Pneumatosis intestinalis and hepatic portal venous gas associated with gas-forming bacterial translocation due to postoperative paralytic ileus

A case report

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Abstract

Rationale: Pneumatosis intestinalis (PI) and hepatic portal venous gas (HPVG) are rare but potentially lethal conditions in which gas pathologically accumulates in the portal vein and intestinal wall, respectively. Proposed mechanisms include flatus escaping through an injured intestinal mucosa into the submucosa and thence into the portal venous system, or bacterial translocation (BT) of gas-forming enteric microorganisms from the gut into and through the intestinal wall to other organs. However, there has been no clear histopathological evidence to support these hypotheses.

Patient concerns: A 61-year-old man underwent sigmoidectomy for colonic adenocarcinoma. Postoperatively, he developed paralytic ileus and then had a sudden cardiopulmonary arrest.

Diagnoses: PI and HPVG were found at autopsy, presumably caused by the postoperative paralytic ileus and associated with BT of gas-forming organisms.

Interventions: Cardiopulmonary resuscitation was unsuccessful.

Outcomes: Postmortem imaging indicated the presence of massive PI and HPVG. At autopsy, there was marked intestinal emphysema with diffuse ischemic mucosal necrosis and severe pneumatosis in the stomach and intestine and marked gaseous dilation of the intrahepatic portal veins. Postmortem bacterial cultures revealed enteric bacteria in the peripheral blood and liver tissue.

Lessons: Postoperative ileus leading to intestinal mucosal damage may be associated with BT of gas-forming enteric bacteria and the rapid onset of PI and HPVG with a lethal outcome.

Abbreviations: BT=bacterial translocation, HPVG=hepatic portal venous gas, PI=pneumatosis intestinalis.

Keywords: bacterial translocation, gas-forming bacteria, hepatic portal venous gas, paralytic ileus, pneumatosis intestinalis

1. Introduction

Pneumatosis intestinalis (PI) and hepatic portal venous gas (HPVG) are abnormal accumulations of gas in the intestinal wall and portal vein, respectively. Bacterial translocation (BT) is defined as the passage of bacteria and bacterial endotoxins from the gut to other organs.1 Experimental studies have suggested that PI, HPVG, and BT share a similar pathogenetic mechanism in which mucosal disruption allows for the invasion of flatus and infectious agents into tissues where they are not normally present.[2,3] However, these mechanisms have not been fully understood because of insufficient histopathologic and postmortem evidence in humans. Moreover, little has been reported on the clinical and pathologic relationship between BT and pneumatoses such as PI and HPVG.

We present the case of a 61-year-old Japanese man who died after a sigmoidectomy complicated by postoperative paralytic ileus, with the subsequent rapid onset of BT, PI, and HPVG. We review the ante- and postmortem clinical, radiological, and histopathological findings.

2. Case report

A 61-year-old Japanese man, with no relevant previous history of disease, was admitted with suspected sigmoid colon cancer. He had been evaluated at another hospital with computed tomography after a positive fecal occult blood test. Physical examination of the patient on admission revealed obesity (body mass index: 34.2 kg/m²) with normal vital signs, a blood pressure of 125/73 mmHg, and a pulse rate of 67 with a regular rhythm. Laboratory data on admission were within reference ranges, except for slightly hyperglycemia (hemoglobin A1c 6.3%, reference range: 4.6–6.2%) and mild renal dysfunction (creatinine 1.14 mg/dL,
Computed tomography revealed a well-demarcated solid mass in the sigmoid colon. Endoscopic biopsy of the mass demonstrated a well to moderately differentiated adenocarcinoma.

Sigmoidectomy and regional lymph node dissection were performed. After the resection of the sigmoid, an initial attempt to perform colorectal anastomosis using a 29-mm diameter intraluminal stapler failed, as the intestinal wall was partially torn because of excessive tensile strength. Eventually, anastomosis was successfully completed using a 25-mm diameter stapler with no air leak. Pathological examination of the resected specimen revealed a moderately differentiated adenocarcinoma of the sigmoid colon, pT1, pN0, M0, stage I (according to the 8th edition of the TNM classification), which had been completely resected (R0).

The postoperative course was uneventful until 5 days after the operation. The patient had begun drinking water on the 6th postoperative day, but he vomited twice and complained of abdominal discomfort at that day. An upright abdominal radiograph showed marked gaseous dilatation of the stomach, small intestine, and colon with air-fluid levels, indicating paralytic ileus (Fig. 1). Therefore, the patient again resumed fasting that evening. The nursing staff noted that he was in bed and conscious at 10:30 pm. However, at 1:50 am, he was found in the toilet in cardiopulmonary arrest. Resuscitation was started immediately but was unsuccessful, and he was pronounced dead.

An autopsy was performed. Computed tomography revealed marked PI, gastric pneumatosis, and HPVG (Fig. 2), but it did not indicate any clear cause of these pneumatosis. There was abdominal distention, but the surgical wound appeared normal for that stage postoperatively. There was sanguineous ascites in the abdominal cavity. However, there was no leakage from the colorectal anastomosis, nor was any residual or metastatic colon cancer found, indicating that the sigmoidectomy itself did not appear to have contributed to the outcome. However, there was marked gastric and intestinal pneumatosis, especially from the jejunum to the ileum, with the intestines spilling out as soon as the abdomen was opened (Fig. 3A and B). Histologically, there were ischemic changes indicated by mucosal thinning and necrosis of the gastrointestinal wall, as well as marked pneumatisos in the mucosal lamina propria and submucosa (Fig. 3C–F). The intrahepatic portal veins were markedly dilated due to gas accumulation, but the portal vein walls themselves were structurally normal (Fig. 4). There was centrilobular liver cell congestion and necrosis. Enterococcus faecalis and Klebsiella oxytoca were detected in blood drawn from the heart and also in tissue harvested from the liver, indicating BT and subsequent bacteremia. Informed written consent was obtained from the patient for publication of this case report and accompanying images.

3. Discussion

This is a rare autopsy case of PI and HPVG along with BT, which may itself have contributed to the pneumatisos. These were the findings in a patient who underwent an apparently uneventful sigmoidectomy but then developed paralytic ileus, with subsequent intestinal necrosis and bacteremia. The patient’s only known risk for postoperative complications was obesity. His sudden deterioration was completely unexpected. Although there have been a few reports of PI and HPVG associated with BT, they were based only on clinical and radiological observations rather than histopathological findings.\(^4\)\(^–\)\(^7\)

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**Figure 1.** Abdominal upright radiograph 5 days after the operation. The stomach and intestine are markedly dilated with gas, and air-fluid levels are seen, indicating paralytic ileus.

**Figure 2.** Postmortem computed tomography reveals extensive dilatation of intrahepatic portal veins in left and right branches and central and peripheral vessels and marked gastric and intestinal dilatation with fine, bubble-like aerocysts in the walls. These radiological findings indicate the presence of hepatic portal venous gas, pneumatosis intestinals, and gastric pneumatosis.
BT is defined as passage of bacteria and bacterial endotoxins from the gut through the mucosa to other organs. Intraluminal microorganisms are able to breach the intestinal mucosal barrier in an event that can be seen in cases of intestinal obstruction, inflammatory bowel disease, jaundice, acute pancreatitis, organ transplantation, Stevens–Johnson syndrome, and malignant neoplasms. In the present case, the postmortem morphologic analysis revealed extensive necrosis of the mucosal epithelia in the small and large intestine. The intestinal crypts overlying the gas-distended submucosa were partially desquamated. There was no increase in the numbers of lymphocytes and plasma cells over what are usually seen in normal, uninflamed mucosa. These histological changes were compatible with a diagnosis of ischemic enterocolitis. In the presence of ileus, as seen in our presenting patient, a rapid increase in intestinal luminal and intra-abdominal pressure can lead to multisystem organ failure, including intestinal ischemia and hepatic fragility. The autopsy findings of the patient suggest that postoperative
paralytic ileus may have led to widespread ischemic intestinal necrosis and subsequent BT.

Generally, BT is clinically confirmed by blood cultures positive for intestinal bacterial flora.\textsuperscript{[18]} The autopsy in this patient revealed that \textit{E. faecalis} was isolated not only from the blood but also from the liver tissue, suggesting that BT may have led to lethal bacteremia. \textit{Enterococcus} species, including \textit{E. faecalis}, are part of the normal flora of the human gut.\textsuperscript{[19]} However, life-threatening \textit{E. faecalis} infections, including urinary tract infections, intra-abdominal and pelvic infections, meningitis, and endocarditis, are sometimes seen in hospitalized patients with weakened immune systems as a result of surgery, cancer treatment, dialysis, organ transplantation, and immunodeficiency.\textsuperscript{[20–23]} As discussed below, PI and HPVG may be also induced in a patient postoperatively by BT by gas-forming \textit{E. faecalis}. The aggressiveness of \textit{E. faecalis} depends on a patient’s clinical status independent of the virulence of the bacterial strains.\textsuperscript{[24]} Thus, postoperative patients are unusually vulnerable to \textit{E. faecalis} infection, which worsens disease severity and clinical outcome.\textsuperscript{[25,26]}

PI and HPVG are rare but potentially lethal conditions mostly associated with several acute abdominal conditions such as inflammatory bowel disease, infections, graft-versus-host disease, diverticulitis, bowel obstruction, and ischemic enterocolitis.\textsuperscript{[27–29]} Although the mechanism is not fully understood, many researchers hold to a mechanical theory, in which PI and HPVG stem from flatus escaping through a damaged intestinal mucosa into the submucosal stroma and vessels and, from there, into the portal venous system. An association between BT and PI and HPVG has rarely been described in the literature. However, in an experimental study, PI was induced artificially by injecting gas-forming enteric bacilli into the wall of rats.\textsuperscript{[31]} The present autopsy case may support the theory that this same mechanism may occur in humans, namely, that PI and HPVG may arise from BT of gas-forming microorganisms such as \textit{Enterobacter} and \textit{Klebsiella} species. Normally, the intestinal mucosal barrier physiologically resists such invasion, but mucosal necrosis disrupts the barrier’s integrity, which can predispose to BT. Thus, both the mechanical pressure from the distended gut and BT by gas-forming bacteria may act synergistically to produce PI and HPVG, a potentially lethal condition even in patients with no known risks. In patients with postoperative ileus, close attention to possible bacterial infections are warranted to avoid clinical deterioration.

Author contributions
Sayumi Tahara wrote the manuscript; Yasuhiro Sakai wrote and organized the manuscript; and Hidetoshi Katsuno, Makoto Urano, Makoto Kuroda, and Tetsuya Tsukamoto critically reviewed the manuscript and had useful professional suggestions. All authors reviewed the final version of the manuscript.

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Figure 4. Autopsy findings in the liver. (A, B) Central and peripheral (arrow) portal veins are markedly dilated. (C) Histologically, the portal veins in Glisson’s sheaths are dilated, but the venous walls are structurally normal. Hematoxylin and eosin staining. Bar = 200 μm.
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