Case Report

Pneumatosis cystoides intestinalis – an incidental finding with unpredictable evolution

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ABSTRACT

Pneumatosis cystoides intestinalis is an uncommon disease with unknown etiology characterized by the presence of multiple gas-filled cysts within the submucosa or subserosa of the intestinal wall. Pneumoperitoneum and/or intestinal perforation are complications that may be associated with pneumatisos cystoides intestinalis. The patients are often prone to misdiagnosis or mistreatment.

We are presenting a case of pneumatisos cystoides intestinalis in a 42 year-old woman affected by peritoneal free air and numerous, diffuse, bubble-like intramural gas collections into the jejunum and ileum, showed in CT-enterography images. The woman had a carcinoid tumor located in jejunum two years ago, treated with enterectomy. Recent complaints of nonspecific symptoms of abdominal discomfort and diarrhea motivated the realization of CT scan, serum chromogranin and urine 5-hidroxindolacetic acid for hypothesis of tumor carcinoid recurrence withdraw. The only change found was the presence of pneumatisos cystoides intestinalis in CT-enterography images without intestinal necrosis, bleeding or evident obstruction. For that reason no surgical procedure was realized and the patient stayed on surveillance. Actually, the patient complaints are sporadic abdominal discomfort, without pneumatisos cystoides intestinalis clinical evidence.

Conclusion: The treatment plan of patient with PCI depends on underlying cause and clinical condition severity. When conservative treatment is adopted the clinical evolution of pneumatisos cystoides intestinalis is unpredictable and can even disappear in an indeterminate number of patients.

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Pneumatose cistoide intestinal – um achado incidental com evolução imprevisível

R E S U M O

A pneumatose cistoide intestinal é uma doença incomum, de etiologia desconhecida, caracterizada pela presença de múltiplos cistos preenchidos com gás na submucosa ou subserosa da parede intestinal. O pneumoperitoneu e/ou a perfuração intestinal são complicações que podem estar associadas à pneumatose cistoide intestinal. Os pacientes geralmente estão sujeitos a erros de diagnóstico ou de tratamento.

Apresentamos um caso de pneumatose cistoide intestinal em paciente do sexo feminino, 42 anos de idade, com ar livre peritoneal e numerosas coleções gassosas intramurais, difusas e semelhantes a bolhas no jejuno e íleo, visualizados em imagens de enterografia por tomografia computadorizada (TC). Há dois anos, a paciente teve um tumor carcinoide localizado no jejuno que foi tratado com enterectomia. As queixas recentes de sintomas inespecíficos, desconforto abdominal e diarreia motivaram a realização da TC e exame de cromogranina sérica e ácido 5-hidroxindolacético na urina para excluir a hipótese de recorrência do tumor carcinoide. A única alteração encontrada foi a presença de pneumatose cistoide intestinal em imagens de enterografia por TC sem necrose intestinal, sangramento ou obstrução evidente. Por esse motivo, nenhum procedimento cirúrgico foi realizado, e a paciente permaneceu em observação. Atualmente, a queixa da paciente é de desconforto abdominal esporádico, sem evidência clínica de pneumatose cistoide intestinal.

Conclusão: O plano de tratamento de pacientes com PCI depende da causa subjacente e da gravidade da condição clínica. Quando o tratamento conservador é adotado, a evolução clínica da pneumatose cistoide intestinal é imprevisível e pode até desaparecer em alguns pacientes.

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Introduction

Pneumatosis cystoides intestinalis is a clinical entity and may be very heterogeneous representing a challenge for the clinician. It is characterized by polycystic air in the submucosa or subserosa of the intestine developing acute abdomen with intestinal obstruction, intestinal necrosis, perforation, pneumoperitoneum and/or peritonitis, or being present with nonspecific abdominal complain.1 The CT scan may confirm the diagnosis, give some additional information, providing differential diagnosis, to help determining the primary cause or some possible coexistent complications.2 It usually related to respiratory infections, tumor or collagen disease, traumas, immunosuppression.3-5 Treatment plan to PCI patients depend on underlying cause, and clinical condition severity.

Case report

A 42 year-old woman with recent nonspecific symptoms of abdominal discomfort, nausea and diarrhea was observed in a routine surgical outpatient appointment. The patient was submitted to an enterectomy two years ago for a jejunum carcinoid tumor. The physical examination detected a median incision scar, and only a slight tenderness at palpation in the epigastrium with no acute abdomen. CT scan, serum chromogranin and urine 5-hydroxindolacetic acid were performed to withdraw tumor carcinoid recurrence hypothesis. Patient study was completed with complete blood count, blood biochemistry with amylase, lipase, creatinine phosphokinase, alkaline phosphatase and lactic dehydrogenase among others, and CT enterography. The only unexpected change found was the presence of PCI in CT enterography images with pneumoperitoneum but without intestinal necrosis, bleeding or evident obstruction. The CT enterography images showed the presence of numerous gas collections within the wall of the small bowel, consisting mainly of isolated bubbles, free intraperitoneal airs, multiple air-fluid levels, and distention in the small bowel, proximal to ileo-ileal anastomosis (Fig. 1).

A conservative approach with antibiotic therapy was adopted with patient symptoms improvement. She was medicated with proton pump inhibitor (omeprazole 20mg/day), metoclopramide (10mg 15 min before breakfast, lunch and dinner) during two months and antibiotic (rifaximin 200mg 12/12h during 12 days).

Actually, sporadic abdominal pain and discomfort, but without PCI, are the only patient complaints (study with CT enterography and capsule enteroscopy) (Fig. 2) or tumor recurrence.

Discussion

PCI is a rare condition characterized by multilocular gas-filled cysts localized in the submucosa and subserosa, most frequently observed in the terminal ileum of gas-trointestinal tract.6 The incidence of PCI is unknown and the etiology is
unclear. Pneumatosis cystoides intestinalis can be classified as primary disease, (15%) of unknown cause, and secondary (85%) associated to gastrointestinal diseases, such as intestinal obstruction, cystic fibrosis, peptic ulcer, diverticula, inflammatory bowel disease, mesenteric infarction, chronic intestinal pseudo-obstruction, respiratory infections, tumor or collagen disease, traumas, immunosuppression and the use of steroids.

The underlying diagnosis, when the cause is secondary, is not always easy.

There are several theories to explain the secondary PCI pathogenesis: mechanical theory, bacterial theory, pulmonary theory, and chemical theory. The mechanical theory suggests existence of air inflow through damaged mucosa, caused by intestinal obstruction, severe constipation, gastrointestinal ulcers, or intestinal necrosis, resulting in increased intestinal pressure. The bacterial theory suggests the invasion of gas-producing bacteria into intestinal submucosa, producing gas at the intestinal walls. The pulmonary

Fig. 1 – CT enterography (A, axial image; B and C, axial and coronal images, lung window) demonstrates free extra-peritoneal gas (arrows) and multiple air-filled cysts in the small bowel walls. There were no other abnormalities in the small bowel or in the colon.

Fig. 2 – Capsule endoscopy and CT enterography images, 6 months after the acute event. Signs of pneumatosis are absent with few enteric stasis near the enteric anastomosis in CT enterography.
Pneumatosis cystoides intestinalis refers to intramural gas often presenting as one incidental finding in benign cases. Patients treatment with PCI, depend on underlying cause and clinical condition severity. Benign PCI should always be treated conservatively. When conservative treatment is adopted the clinical evolution of PCI is unpredictable and can even disappear in an indeterminate number of patients.

Conflicts of interest

The authors declare no conflicts of interest.

REFERENCES