ORIGINAL ARTICLE

INVASIVE ENTERITIS BY STRONGYLOIDES STERCORALIS PRESENTING AS ACUTE ABDOMINAL DISTRESS UNDER CORTICOSTEROID THERAPY

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Overwhelming helminthiasis is still a problem in endemic areas, especially in immunocompromised patients. We report a case of invasive intestinal strongyloidiasis that was clinically expressed as acute abdominal distress in a 73-year-old man from São Paulo who had been receiving methylprednisone, 20 mg/day, for one year for osteoarthritis. A surgical specimen from the ileum revealed invasive enteritis with severe infestation by Strongyloides stercoralis. The patient died of sepsis 6 days after surgery. The possibility of invasive strongyloidiasis should be considered in the differential diagnosis of acute abdominal distress in patients undergoing immunosuppressive therapy.


Strongyloidiasis is a worldwide parasitic disease, endemic in tropical countries like Brazil. The great majority of cases are asymptomatic, but when symptoms develop, patients may present mild abdominal pain, nausea, vomiting, and watery diarrhea. Poor personal hygiene, cutaneous rashes, and eosinophilia are important clues for the diagnosis. However negative stool cultures or eosinopenia do not rule out the diagnosis and under severe conditions may point to the disseminated form of strongyloidiasis.

This report presents a patient who was undergoing corticosteroid therapy for osteoarthritis and who developed a small bowel obstruction due to Strongyloides stercoralis hyperinfection.

CASE REPORT

A 73-year-old man from São Paulo came to the emergency room with a 4-day history of cramping abdominal pain, nausea, vomiting, and watery diarrhea. On examination, he was afebrile with a markedly tender, distended abdomen with rebound tenderness and guarding. His white blood cell count was 8 200 / mm³, and no eosinophils were detected on peripheral smear. An abdominal roentgenogram revealed air-filled dilated loops of small bowel consistent with small bowel obstruction. He underwent exploratory laparotomy that showed an inflamed section of distal ileum with a narrowed lumen adhered to the retroperitoneum. Resection of the affected segment was performed with construction of an ileocolic anastomosis. The patient subsequently developed sepsis with blood cultures positive for anaerobic bacteria and died on the sixth postoperative day. The histopathologic examination of the surgical specimen revealed invasive enteritis with severe infestation by...
Strongyloides stercoralis. Many larval forms were found within the crypts and invading the deeper layers of the intestinal wall with inflammation and focal granulomas (Fig. 1).

Figure 1 - Enteritis by Strongyloides stercoralis. Larval forms invading the crypts of the mucosa.

Retrospective analysis of the patient’s records, which were not available at the emergency room, revealed that he had been receiving treatment with methylprednisone, 20 mg/day, for one year for osteoarthritis.

DISCUSSION

Even though Strongyloides stercoralis is an intestinal nematode, infected patients rarely present with symptoms from the gastrointestinal tract, such as intestinal obstruction or acute abdominal distress. Severe gastrointestinal symptoms due to increased worm burden occur in the hyperinfection syndrome. This pathologic condition results from larvae that become infective within the intestine. Predisposing factors for this overwhelming autoinfection include immunosuppression, particularly impaired cell mediated immunity, steroid therapy, severe malnutrition or alcoholism, leukemia, lymphoma, and the expanding population of HIV-infected patients. In this condition, filariform larvae invade the intestinal walls and may spread through lymphatic and hematogenic pathways to distant organs such as the liver and biliary tract, pancreas, lungs, and the central nervous system. This hyperinfection is usually accompanied by bacteremia with enteric organisms, leading to fatal septicemia.

The reported case is from an elderly patient who was undergoing chronic corticosteroid therapy for osteoarthritis who developed signs and symptoms of small bowel obstruction. The information about his treatment with corticosteroid was not available when he was admitted at the emergency room. Retrospective analysis of the patient’s records revealed that he was under investigation for anemia and recurrent diarrhea. The negative results for the parasitologic stool examinations did not rule out the possibility of parasitic infection, and the chronic use of corticosteroid should have prompted prophylactic treatment with thiabendazole. With this panel of clinical data and the development of signs and symptoms of acute abdominal distress, the hypothesis of hyperinfection syndrome by Strongyloides stercoralis would have led to a course of treatment that did not include surgery.

Boken et al. reported a case of Strongyloides stercoralis hyperinfection in a patient who was treated successfully with rectal administration of thiabendazole. Undiluted thiabendazole suspension (500 mg/5 mL) was given by retention enema for 14 days (1.5 g every 12 hours) because severe upper intestinal obstruction precluded oral administration.

In the present case, the diagnosis of Strongyloides stercoralis hyperinfection was made only after surgery by pathological examination of the surgical specimen and did not prevent subsequent complications that led to the patient’s death. Thus, this report highlights the importance of the preoperative evaluation of co-morbid states, even under emergency conditions, since they may lead to life-threatening complications.

In conclusion, the possibility of Strongyloides stercoralis hyperinfection should be considered in the differential diagnosis of acute abdominal distress in patients receiving cortisone analogs and other immunosuppressive drugs. Timely appropriate therapy before the development of significant abdominal complications such as perforation and peritonitis would be life-saving.
RESUMO


A síndrome de hiperinfeccão por Strongyloides stercoralis ainda constitui problema em áreas endêmicas, especialmente em pacientes portadores de imunossupressão. Os autores relatam um caso de estrongiloidíase intestinal invasiva, que se apresentou clinicamente com quadro de abdome agudo obstrutivo, em um paciente masculino de 73 anos, que estava recebendo metilprednisona 20 mg/dia há um ano devido à osteoartrite. O exame anatomo-patológico de segmento do íleo demonstrou enterite invasiva com intensa infestação por Strongyloides stercoralis. O paciente evoluiu para óbito com quadro de septicemia, 6 dias após a intervenção cirúrgica. O presente caso destaca a importância da avaliação pré-operatória de estados comorbidos, mesmo em condições de urgência, visto que podem comprometer o êxito do tratamento cirúrgico e a vida do paciente. A possibilidade de estrongiloidíase intestinal invasiva deve ser considerada no diagnóstico diferencial dos quadros de abdome agudo em pacientes submetidos à terapêutica imunossupressora.


REFERENCES


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