Ascites Due to Eosinophilic Gastroenteritis: A Case Report

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Abstract

Eosinophilic gastroenteritis is an uncommon disease characterized by eosinophilic infiltration in the gastrointestinal tract. Kajser was probably the first to report a patient with eosinophilic gastroenteritis in 1937. Eosinophilic gastroenteritis may involve more than one layer of the gastrointestinal tract. Clinical features depend on the layer and location to be involved. Mucosal involvement leads to protein-losing enteropathy, fecal blood loss, and malabsorption. Involvement of muscle layer often causes obstruction of gastric outlet or small bowel. Subserosal involvement manifests as eosinophilic ascites. We presented a 34-year-old female with progressive ascites and lower leg edema. Eosinophilic gastroenteritis was diagnosed after serial examinations. The patient was treated with prednisolone. The ascites subsided soon after initation of steroid treatment. (J Intern Med Taiwan 2009; 20: 81-85)

Key Words: Ascites, Eosinophilic gastroenteritis, Eosinophilia, Eosinophilic infiltration

Introduction

Eosinophilic gastroenteritis is an uncommon disease characterized by eosinophilic infiltration in the gastrointestinal tract. Kajser was probably the first to report a patient with eosinophilic gastroenteritis in 1937\textsuperscript{1}.

Eosinophilic gastroenteritis may involve serosa of the gastrointestinal tract and result into the development of ascites\textsuperscript{2-4}. The ascites analysis may reveal eosinophils to account for a high proportion of white blood cells. We here reported a case of eosinophilic gastroenteritis with predominant symptom of ascites, which was resolved with steroid treatment.

Case report

A 34-year-old Chinese female was admitted for evaluation of progressive abdominal distension and lower leg edema.
She was relatively well in the past until three weeks before this admission. Her chief complaints were upper abdominal pain and frequent loose stool passage. The pain was not related to meal and had no radiation. The abdominal pain was not associated with fever. No obvious body weight loss was noted. Two weeks before admission, she suffered from poor appetite, nausea, abdominal fullness, and swelling of bilateral lower legs. She didn’t have past history of liver, renal or gynecological disease. She denied travel history and special food intake before the symptoms happened.

Physical examination disclosed a moderately developed and nourished woman with acute ill-looking. Her consciousness was clear, blood pressure was 145/75 mm-Hg, pulse rate was 94/min, respiratory rate was 20/min, and body temperature was 36.4°C. Abdomen was soft and distended with shifting dullness on percussion.

Laboratory examination disclosed white blood cell count of 13080/mm³ with neutrophils 69%, lymphocytes 5% and eosinophils 25%, hemoglobin of 13.3 g/dL, hematocrit of 38.8%, and platelet 277000/mm³. Total eosinophils count was 5900/mm³. The blood chemistry disclosed albumin 3.2 g/dL, aspartate aminotransferase (AST) 19 U/L, alanine amino-transferase (ALT) 4 U/L, alkaline phosphatase 69 U/L, total bilirubin 0.9 mg/dL, lactate dehydrogenase (LDH) 417 U/L, blood urea nitrogen (BUN) 5 mg/dL, creatinine 0.9 mg/dL, sodium 133 mmol/L, potassium 3.5 mmol/L, fasting blood sugar 91 mg/dL, total cholesterol 147 mg/dL, triglyceride 85 mg/dL. The hepatitis markers and autoantibody test were negative. Stool was negative for occult blood, ova and parasites. Abdominal sonography and computed tomography scan demonstrated markedly increased wall thickening of bowel loops and severe degree of ascites.

Abdominal paracentesis was performed. The gross appearance of ascites was yellowish and turbid. Analysis of ascites disclosed white blood cell count of 6700/mm³ (differential count: neutrophils 3%, lymphocytes 4%, eosinophils 90%, monocytes 3%), red blood cells 15350/mm³, LDH 242 U/L, protein 2.3 g/dL, sugar 93 mg/dL. The culture & cytology of ascites were negative finding.

The patient underwent an esophagogastroduodenoscopy (Figure. 1), which demonstrated several linear erythematous patches at fundus and multiple erythematous patches at the superior duodenal angle, near the second portion of duodenum. The histological examination of these duodenal lesions (Figure. 2) showed marked increase of eosinophilic infiltration in the lamina propria. The
sigmoidoscopy revealed several erythematous patches over the sigmoid colon. The histological examination of these colonic lesions also showed increased eosinophilic infiltration in the lamina propria.

These features were consistent with the diagnosis of eosinophilic gastroenteritis with mucosa and subserosa involvement. The patient was thus treated with prednisolone. The dosage of prednisolone for initial treatment was 40 mg/day. The patient's symptoms subsided after steroid treatment for 3 weeks and her steroid dose was successfully tapered within two months.

**Discussion**

Eosinophilic gastroenteritis is an uncommon disease characterized by eosinophilic infiltration in the gastrointestinal tract. The infiltration may involve one or more layers of the gastrointestinal wall and other abdominal organs.

The pathogenesis of this disease is poorly understood, but speculation has focused on the selective release of eosinophil major proteins leading to the intestinal epithelial damage. Keshavarzian et al. demonstrated that the number of activated degranulated eosinophils in the mucosa correlated with the severity of eosinophilic gastroenteritis. Several reports have pointed to a possible role of allergies to food and other allergens, while other investigators have refuted an allergic reaction as the etiology of this disease.

Eosinophilic gastroenteritis may involve more than one layer of the gastrointestinal tract. Clinical features depend on which layer and location are involved. Mucosal involvement leads to protein-losing enteropathy, fecal blood loss, and malabsorption. Involvement of muscle layer often causes obstruction of gastric outlet or small bowel. Subserosal involvement manifests as eosinophilic ascites.

In a retrospective study of 40 patients, the most common symptoms were abdominal pain, nausea, vomiting, and diarrhea. In that study, the percentage of involving mucosal layer, muscle layer and subserosal disease were 58%, 30% and 12% respectively.

Talley et al. reported that the patients with subserosal disease could be distinguished from the other two groups by clinical symptoms (abdominal bloating, ascites), higher eosinophils counts, and their dramatic responses to steroid therapy. The diagnostic feature of subserosal disease is a marked eosinophilia, up to 88 percent, in the ascitic fluid. Our patient presented with severe ascites. This picture may be confused with cirrhotic ascites or abdominal carcinomatosis. Fortunately, marked eosinophilia in the ascites is an important clue to get the diagnosis of eosinophilic gastroenteritis.

Hypereosinophilia in peripheral blood is present in 20-90% of patients with eosinophilic gastroenteritis. Thus, the absence of peripheral hypereosinophilia should not exclude the diagnosis of eosinophilic gastroenteritis in patients with unexplained gastrointestinal symptoms.

Radiologically, the hallmark of eosinophilic gastroenteritis is the thickening of mucosal folds demonstrated by barium study, computerized tomography, or ultrasound, depending on the severity and layer(s) involved. However, similar thickening may also be seen in Menetrier's disease, lymphoma, Crohn's disease, and granulomatous disease. Thus, the thickening of bowel wall is not specific for eosinophilic gastroenteritis. The endoscopic appearance in eosinophilic gastroenteritis is also nonspecific, including erythematous, friable, nodular, and occasional ulcerative changes.

The definite diagnosis of eosinophilic gastroenteritis must fulfill the following criteria: (1) the presence of gastrointestinal symptoms, (2) demonstration of eosinophilic infiltration of one or more areas of the GI tract on biopsy, (3) no evidence of parasitic or extraintestinal disease. The bowel
involvement is often patchy in distribution, so more than one panendoscopic examination and biopsies sometimes are necessary to establish the diagnosis. Laparotomy is needed for diagnosis only for those cases complicated with perforation, or involving jejunum and ileum.

In general, the clinical response of eosinophilic gastroenteritis to steroid treatment is usually dramatic. The appropriate duration of steroid treatment remains unknown, but improvement usually occurs within two weeks regardless of the layer of bowel involved. But some patients require more prolonged therapy (up to several months) to induce resolution of symptoms. Patients with refractory relapsing disease may require long-term low-dose steroids or immunosuppressive therapy.

In conclusion, eosinophilic gastroenteritis presenting as ascites is a rare disease. Eosinophilic gastroenteritis should be suspected in the presence of unexplained chronic or relapsing gastrointestinal symptoms. Ascites analysis is helpful in the diagnosis of eosinophilic gastroenteritis with subserosal involvement. Steroid is the mainstay of therapy.

References
嗜伊红性胃腸炎引起腹水：病例報告

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摘 要

嗜伊红性胃腸炎是一個少見疾病，是因為嗜伊红性白血球浸潤腸胃道的黏膜層、肌肉層或漿膜層所引起。此疾病最早是由Kaier在西元1937年提出。臨床症狀依侵犯消化道的位置、範圍及深淺而有所不同。若侵犯黏膜層會引起蛋白質流失性腸病變、慢性消化道出血及消化道吸收不良。若侵犯肌肉層會引起胃出口阻塞或腸道阻塞。而侵犯到漿膜層時大部份會以腹水的症狀來表現。本病人是34歲女性，因腹痛、腹瀉、腹脹及雙腳水腫來本院就診。腹部超音波及電腦斷層檢查發現腸壁增厚及腹水。週邊血液中嗜伊紅性白血球增加，腹腔放液檢查亦發現嗜伊紅性白血球增加。上消化道及下消化道內視鏡檢查發現有胃、十二指腸及乙狀結腸有發炎性反應，經切片發現粘膜層有嗜伊紅性白血球浸潤。證實為嗜伊紅性腸胃炎。經給予類固醇治療後，病況改善並於門診繼續追蹤治療。