Wegener’s granulomatosis mimicking inflammatory bowel disease and presenting with chronic enteritis

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Abstract: Wegener’s granulomatosis, also known as anti-neutrophil cytoplasmic antibody (ANCA)-associated vasculitis, is a small vessel vasculitis with primarily pulmonary, renal, and sinus disease manifestations. The prevalence of Wegener’s granulomatosis is three cases per 100,000 patients. Cardiovascular, neurologic, cutaneous, and joint manifestations have been reported in many case reports and case series. Gastrointestinal manifestations are less noted in Wegener’s granulomatosis, although they have been previously reported in the form of intestinal perforation and intestinal ischemia. Additionally, there are characteristic findings of vasculitis that are noted with active Wegener’s granulomatosis of the small bowel. We report a case of an elderly patient who presented with weight loss, diarrhea, and hematochezia. His symptoms were chronic and had lasted for more than 1 year before diagnosis. Inflammatory bowel disease or chronic enteritis due to Salmonella arizonae because of reptile exposure originally were suspected as etiologies of his presentation. The findings of proteinuria, renal failure, and pauci-immune glomerulonephritis on renal biopsy, in conjunction with an elevated c-ANCA titer, confirmed the diagnosis of Wegener’s granulomatosis with associated intestinal vasculitis. This case demonstrates an atypical presentation of chronic duodenitis and jejunitis secondary to Wegener’s granulomatosis, which mimicked inflammatory bowel disease.

Keywords: ANCA-associated vasculitis, Wegener’s syndrome, pauci-immune glomerulonephritis, Salmonella arizonae, inflammatory bowel disease

Introduction

Wegener’s granulomatosis, also known as anti-neutrophil cytoplasmic antibody (ANCA)-associated vasculitis, is a vasculitis of the small arteries of the body. It is a rare disease, with a prevalence rate of three per 100,000 in the United States according to a 1996 epidemiologic study.1 Wegener’s granulomatosis can present with multiple pulmonary and renal lesions, including cavitary lung lesions, fibrosis, and a rapidly progressive crescentic glomerulonephritis of the kidney.2-4 There are many other reported manifestations of Wegener’s granulomatosis including cerebral vasculitis,3 cutaneous lesions,6 and otitis media.7 There are many case reports of cardiac conduction abnormalities, pulmonary hypertension, pericarditis, valvular disease, and other forms of cardiac manifestations of ANCA-associated vasculitis.4 The American College of Rheumatology’s criteria for diagnosing Wegener’s granulomatosis are clinical and do not require positive c-ANCA serology, although this test is generally helpful. The criteria include abnormal urinary sediment, abnormal findings on chest radiography, oral ulcers or nasal discharge, and granulomatous inflammation on biopsy.9
Gastrointestinal manifestations are unusual in Wegener’s granulomatosis; however, several manifestations have been reported in the literature, including esophageal, small bowel, and colonic lesions.10–25 One of the more common manifestations of Wegener’s vasculitis is intestinal perforation, likely due to bowel ischemia from severe vasculitis. There have been four case reports in the medical and surgical literature regarding small bowel perforation secondary to Wegener’s granulomatosis.11,12,20,25 One case of bowel perforation occurred in c-ANCA-negative Wegener’s granulomatosis.14 In one series, the prevalence of bowel perforation in 62 patients with systemic vasculitides including Wegener’s granulomatosis was 15%. Among those patients with bowel perforation, 16% also had bowel infarction and the observed mortality rate was 62%.6

Other manifestations of Wegener’s granulomatosis in the gastrointestinal tract include spontaneous gastrointestinal bleeding,12,19 and esophageal strictures presenting with odynophagia were reported in 2000 by Fallows et al.21 Jejunal stenosis was also reported in 1997 by Rabe et al,17 and intestinal ischemia after a ruptured aneurysm of the middle colic artery was reported in a patient with c-ANCA-positive vasculitis.22 Most of these uncommon gastrointestinal manifestations reported in the literature are acute and often catastrophic. Pagnoux et al reported on surgical emergencies in patients with small-vessel and medium-vessel vasculitides, and noted extremely high mortality rates and depressed 5-year and 10-year survival rates post abdominal surgery in patients with necrotizing vasculitides such as Wegener’s granulomatosis.6 In this report, we examine a more chronic and insidious presentation of gastrointestinal Wegener’s granulomatosis that closely mimicked inflammatory bowel disease. An interesting side issue was the isolation of Salmonella arizonae in the patient’s stool after exposure to a reptile.

Case report
A 57-year-old Ukranian male with a 45-year history of smoking but no other medical history presented in August 2008 complaining of months of intermittent abdominal pain, diarrhea, and a 40 lb weight loss. His symptoms presented over the course of 1 year. His diarrhea was initially evaluated and attributed to gastroenteritis. Routine studies were sent, including a stool culture. S. arizonae (subtype III), which is associated with osteomyelitis and gastroenteritis in reptile handlers,26 was isolated and the patient reported contact with a pet iguana. Computed tomography of the abdomen and pelvis was ordered and revealed a 1.5 × 1.7 cm spiculated pulmonary nodule in the right upper lobe and thickening of the duodenal, jejunal, and colonic wall with scattered surrounding lymphadenopathy.

The patient was placed on antibiotics and continued to suffer from continued diarrhea and weight loss, so inflammatory bowel disease was suspected. In September 2008 he underwent endoscopic studies which revealed a hiatal hernia, old blood in the stomach, mucosal swelling near the gastric antrum, and three areas of circumferential ulceration and inflammation with mucosal pigmentation/erythema (see Figure 1). The areas were biopsied, and the biopsy showed reported duodenitis and jejunitis with acute on chronic inflammation. There were no signs of architectural distortion or cryptitis/crypt abscesses, Helicobacter pylori stains were negative, and no bacteria were apparent on biopsy. Given that

Figure 1 Multiple images from EGD procedures. (A) EGD image from September 2008 showing a normal gastric antrum. (B) EGD image from September 2008 showing mucosal swelling near the gastric antrum. (C) EGD image showing three discrete patches of inflammation in the duodenum. (D) EGD image from September 2008 showing inflammation and ulceration in the jejunum. (E) EGD image from November 2008 showing inflammation, hyperpigmented mucosa, and erythematous mucosa in the duodenum. (F) EGD image from November 2008 again showing duodenitis.

Abbreviation: EGD, esophagogastroduodenoscopy.
the biopsies did not show any findings consistent with the suspected diagnosis of inflammatory bowel disease, his symptoms were attributed to *S. arizonae* gastroenteritis.

Fine needle aspiration of his pulmonary nodule was nondiagnostic, showing only clustered atypical epithelial cells. A positron emission tomography scan was obtained and showed a focus of hypermetabolic activity in the right upper lobe which was read as probable granulomatous disease.

He continued to have recurrent episodes of abdominal pain and vomiting in November 2008, and treatment of the presumptive *S. arizonae* gastritis/enteritis with trimethoprim-sulfamethoxazole and ciprofloxacin failed to improve his symptoms. *S. arizonae* was not grown from stool samples that were resent.

One month later, the patient presented with back pain and decreased urine output. Acute kidney injury was noted, with a serum creatinine of 4.37 mg/dL, while urine studies revealed severe proteinuria and hematuria. Initially the acute kidney injury was thought to be secondary to prerenal azotemia from dehydration or acute interstitial nephritis secondary to treatment with ciprofloxacin and trimethoprim-sulfamethoxazole. A c-ANCA level was drawn for workup of his renal failure and came back positive only in the ascending colon, which showed a focus of hypermetabolic activity in the right upper lobe and unusually pigmented duodenal mucosa (Figure 1). Biopsies of multiple sites were obtained, which later came back positive only in the ascending colon, which showed a tubular adenoma. Slides from a macroscopic jejunal ulcer seen on endoscopy (Figure 2) showed inflammation but no features typical of inflammatory bowel disease. Given the unexplained nature of his systemic symptoms, acute kidney injury, and the positive c-ANCA, a kidney biopsy was performed and showed rapidly progressive glomerulonephritis due to pauci-immune crescentic glomerulonephritis with 25% crescents (Figure 2).

This renal biopsy result was clearly consistent with an ANCA-associated lesion, and the patient received three doses of cyclophosphamide between February and April 2009. Given the finding of a pulmonary nodule, video-assisted thoracoscopic surgery was performed in June 2009. The pathology results later came back positive for adenocarcinoma with clear margins. The lesion was surgically excised and a subsequent positron emission tomography scan was found to be negative for metastasis. He resumed cyclophosphamide therapy in December of 2009 and has had significant improvement of his gastrointestinal symptoms, renal impairment, and systemic symptoms on steroid and immunosuppressive therapy.

**Discussion**

While inflammatory bowel disease commonly occurs in middle-aged patients and can present in a manner very similar to the patient presented in the above report, it was not the underlying pathological process in this case. *S. arizonae*, which was also isolated in this case, is associated with exposure to reptiles and birds, and has been associated with bone infection, as well as joint infection, sepsis, and gastroenteritis. Rattlesnake products, such as rattlesnake venom pills ingested by cancer patients as a homeopathic remedy, have also been associated with infection due to this strain of bacteria. Patients infected with *S. arizonae* are usually immune-compromised secondary to young age, malignancy, or human immunodeficiency syndrome. Ultimately, it turned out to be a very interesting finding that was probably unassociated with the patient’s pathology.

The renal biopsy in particular was helpful because, unlike endoscopic biopsies, it was able to demonstrate the vasculitis adequately on histologic examination. While inflammatory bowel disease can often be diagnosed using endoscopic mucosal biopsies, vasculitides like Wegener’s granulomatosis are often not well demonstrated on biopsy of the gastrointestinal tract since these biopsies are too superficial to visualize the intestinal arterioles.

The pigmentation of the duodenum and jejunum and the chronic inflammation seen were helpful clues that a vasculitis was present. The macroscopic pigmented appearance of the mucosa in the duodenum and jejunum predicted microscopic

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Figure 2 Hematoxylin and eosin stains of mucosal biopsies from esophagogastroduodenoscopy and trichrome stain of renal biopsy showing pauci-immune crescentic glomerulonephritis. (A–C) Multiple slides of jejunal mucosal ulcer with nonspecific inflammatory infiltrate without any bacteria, cryptitis, crypt abscesses, or evidence of vasculitis. (D) Trichrome stain from renal biopsy specimen showing a fibrocellular crescent consistent with rapidly progressive crescentic anti-neutrophil cytoplasmic antibody-positive-associated glomerulonephritis.
vasculitis, as was reported by Singhal et al in 2007. This unusual presentation of Wegener's granulomatosis in the small bowel details the need for increased clinical suspicion to diagnose vasculitis in the appropriate setting. The difficulty in making the diagnosis of Wegener's granulomatosis from mucosal biopsies was highlighted because ultimately a renal biopsy was needed to confirm the vasculitic process.

Disclosure
The authors report no conflicts of interest in this work.

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