Salvatore Vadala di Prampero, Marco Marino, Francesco Toso, Claudio Avellini, Vu Nguyen, Dario Sorrentino

A Department of Pathology, University Hospital of Udine, Udine, Italy; B Department of Internal Medicine, Virginia Tech Carilion School of Medicine, Roanoke, VA, USA; C Department of Clinical and Experimental Medical Sciences, University of Udine School of Medicine, Udine, Italy

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Crohn disease · Inflammatory bowel disease · Myositis · Adalimumab · Anti-tumor necrosis factor · Extraintestinal manifestations

Abstract
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Single Case

Isolated Bilateral Gastrocnemius Myositis in Crohn Disease Successfully Treated with Adalimumab

Salvatore Vadala di Prampero, Marco Marino, Francesco Toso, Claudio Avellini, Vu Nguyen, Dario Sorrentino

A Department of Pathology, University Hospital of Udine, Udine, Italy; B Department of Internal Medicine, Virginia Tech Carilion School of Medicine, Roanoke, VA, USA; C Department of Clinical and Experimental Medical Sciences, University of Udine School of Medicine, Udine, Italy

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60 mg/day was initiated with rapid resolution of calf tenderness; however, tenderness soon returned when the dose was tapered to 10 mg/day. Subsequently, prednisone and azathioprine were discontinued, and adalimumab was started at standard induction and maintenance doses. The patient’s symptoms resolved shortly after the first induction dose. A repeat magnetic resonance imaging of the calves – 3 months after starting adalimumab – showed complete resolution of muscle inflammation. To our knowledge, this is the first case of gastrocnemius myositis – a rare extraintestinal manifestation of Crohn disease – successfully treated with anti-tumor necrosis factor agents.

Introduction

Inflammatory bowel diseases (IBD), which include Crohn disease (CD) and ulcerative colitis, may be associated with a variety of extraintestinal manifestations (EIMs) that can cause greater morbidity than the underlying intestinal disease and may represent the first sign of IBD. Since EIMs can involve multiple organ systems, some authors believe that IBD is a systemic disorder with predominant bowel manifestations. As many as 36% of the IBD patients have at least one EIM: some correlate to the activity of bowel inflammation (joint, skin, and ocular manifestations), some to small bowel dysfunction (cholelithiasis or nephrolithiasis), while others are nonspecific disorders (osteoarthritis or hepatobiliary disease) [1]. The most frequent EIMs in IBD are arthropathies (4–23%), followed by cutaneous manifestations (2–34%), ocular manifestations (4–12%), hepatobiliary diseases (5–15%), and metabolic bone diseases [2]. Muscular involvement in IBD is rarely reported in the literature [3].

Case Presentation

A 26-year-old male with a 6-year history of ileocolonic CD presented with mild diarrhea and sudden tenderness in both calves and no other symptoms. His CD at diagnosis involved the terminal ileum and cecum and was initially treated with a cycle of prednisone 60 mg daily for 4 weeks and azathioprine 2.5 mg/kg daily. Azathioprine was then continued in monotherapy, and the patient remained in clinical remission with normal inflammatory markers (fecal calprotectin and C-reactive protein) for the following 5 years. His past medical history included only allergic rhinitis.

On physical examination, calves bilaterally were tender on palpation; no rash, skin lesion, or signs of systemic vasculitis were observed. Hematologic investigations showed elevated C-reactive protein at 14.30 mg/L (normal 0–5), erythrocyte sedimentation rate at 31 mm/h (normal 1–10), and positive anti-Saccharomyces cerevisiae Ig G [ASCA IgG] titer at 1:160. Other laboratory tests including antineutrophil cytoplasmic antibodies [c-ANCA], creatine kinase [CK], antinuclear antibodies, anti-smooth-muscle antibodies, antimitochondrial antibodies, and anti-Jo-1 antibodies were within normal limits.

Ultrasound Doppler studies of the lower extremities excluded deep vein thrombosis. Magnetic resonance imaging [MRI] of the legs showed bilateral high-intensity signal in the gastrocnemius muscle fibers beneath the muscle membrane spreading to the right flexor
muscle of the hallux and the tibialis anterior – a picture consistent with myositis (Fig. 1). Electromyography showed neither nerve nor muscle abnormalities. Subsequently, a gastrocnemius muscle biopsy was performed, which showed nonspecific lymphocytic myositis (Fig. 2). A colonoscopy was also performed to assess intestinal disease activity, which showed few aphthous ulcers in the ileum (simple endoscopic CD score of 3).

The patient was started on prednisone 60 mg daily, resulting in complete clinical remission of his myositis within a few days. After 1 month, prednisone was tapered at 10 mg every other week. However, at the dose of 10 mg daily, the patient experienced recurrence of symptoms with calf pain. Prednisone was increased to 60 mg daily, and after 1 month it was slowly tapered by 5 mg every other week. Azathioprine was maintained all along at the dose of 2.5 mg/kg daily. Once again, tapering prednisone below 10 mg daily resulted in symptom relapse. At this point, prednisone and azathioprine were discontinued and adalimumab was initiated at regular induction (160/80 mg) and maintenance (40 mg every other week) doses. The patient became completely asymptomatic few days after the first induction dose. Repeat MRI 3 months later did not show any significant inflammation of the gastrocnemius muscles (Fig. 3). After 9 months of adalimumab therapy with the subject still asymptomatic, azathioprine was reintroduced at a dose of 2.5 mg/kg daily, and after 3 additional months, only azathioprine was continued. 1 year after adalimumab discontinuation, the patient remained in full clinical remission from myositis while on azathioprine monotherapy.

Discussion

We report here the case of a CD patient who developed gastrocnemius myositis – apparently as an EIM – with normal CK and an elevated ASCA IgG titer. Published case reports of gastrocnemius myalgia linked to CD have shown similar prevalence in both genders, with patient age ranging from 19 to 50 years. Muscle histology showed granulomatous myositis in 2 cases [4, 5], nonspecific myositis in 4 cases [6–8, 11], and vasculitis in 3 other cases [9, 10].

Most cases, as well as our own, had normal serum levels of CK and c-ANCA. Only 1 case presented with high c-ANCA titers [11]. The main features of reported cases of gastrocnemius myalgia associated with CD are summarized in Table 1. All cases of calf-limited myalgia reported in the literature responded to prednisolone [4, 6–12] except 1 [5]. In 2 young females, a combination treatment of prednisolone plus azathioprine or cyclophosphamide [10] seemed to be effective.

Our patient initially responded to prednisone, but his symptoms quickly relapsed after the dose was lowered below 10 mg/day. This case appears to differ from the others since this is the first case in which ASCA positivity (frequently reported in CD) was detected. It is also the first case of gastrocnemius myositis successfully treated with an anti-tumor necrosis factor (TNF) agent.

Anecdotal reports suggest that TNF inhibitors can be helpful in the treatment of dermatomyositis and polymyositis, although other reports do not show a benefit or even show that these disorders could actually be the result of such treatment (e.g., the onset of myositis has been reported in rheumatoid arthritis patients during anti-TNF therapy) [13–15]. The myositis in IBD patients differs from the myositis resulting from other causes since it does not
appear to be associated with an increased CK or with specific autoantibodies. It also does not show typical electromyography or histology features. Its relationship to IBD may explain why this type of myositis, but not others, might respond to anti-TNF therapy.

In conclusion, gastrocnemius myositis can be a rare EIM of CD, which may be particularly troublesome for the patient and refractory to conventional therapies. Our case report is the first in the literature that describes a case successfully managed with anti-TNF agents.

**Statement of Ethics**

The authors have no ethical conflicts to declare.

**Disclosure Statement**

The authors have no conflicts of interest to disclose.

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Fig. 1. Axial MRI of the calves. T2-weighted spin-echo MR images show diffuse myositis pattern involving several different muscle groups, most marked in the medial and lateral head of the gastrocnemius muscles (arrows).

Fig. 2. Histological findings of calf muscle biopsy (HE × 20). Lymphocytic infiltration in striated muscle cells with typical aspects of cellular regression.
Fig. 3. Axial MRI of the calves during treatment with adalimumab. T2-weighted spin-echo MR images demonstrate the absence of inflammation in the previously involved muscle groups (for comparison, see Fig. 1).
### Table 1. Features of reported cases of gastrocnemius myalgia in CD

| First author [ref.] | Age, years | Sex | Laboratory findings | Muscle histology | Treatment |
|---------------------|------------|-----|---------------------|------------------|-----------|
| Ménard [4]          | 44         | M   | normal CK           | granulomatous myositis | prednisolone |
| Gilliam [9]         | 19         | M   | normal CK           | vasculitis        | prednisolone |
| Hall [11]           | 32         | F   | normal CK           | myositis          | prednisolone |
| Drabble [12]        | 50         | M   | normal CK           | not performed     | prednisolone |
| Diószeghy [5]       | 41         | M   | normal CK           | granulomatous myositis | no response to steroids or nonsteroidal anti-inflammatory drugs |
| Disdier [10]        | 21         | F   | normal CK           | vasculitis        | prednisolone |
|                     | 26         | F   | normal CK           | vasculitis        | prednisolone and cyclophosphamide |
| Christopoulos [6]   | 19         | F   | normal CK           | myositis          | prednisolone |
| Shimoyama [7]       | 33         | F   | elevated CK         | myositis          | mesalamine, colon surgery |
| Mogul [8]           | 15         | M   | normal CK           | myositis          | methylprednisolone, methotrexate |
| Current report      | 26         | M   | normal CK           | myositis          | adalimumab, then azathioprine |

CD68, IgG, ASCA, c-ANCA, ANCA.