Statin-Associated Polymyositis Following Omeprazole Treatment

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Abstract

Statins are an extensively used class of drugs, and myopathy is an uncommon, but well-described side effect of statin therapy. Inflammatory myopathies, including polymyositis, dermatomyositis, and necrotizing autoimmune myopathy, are even rarer, but debilitating, side effects of statin therapy that are characterized by the persistence of symptoms even after discontinuation of the drug. It is important to differentiate statin-associated inflammatory myopathies from other self-limited myopathies, as the disease often requires multiple immunosuppressive therapies. Drug interactions increase the risk of statin-associated toxic myopathy, but no risk factors for statin-associated inflammatory myopathies have been established. Here we describe a male, aged 59-years, who had been treated with a combination of atorvastatin and gemfibrozil for approximately 5 years and developed polymyositis after treatment with omeprazole for 7 months. Symptoms did not resolve after discontinuation of the atorvastatin, gemfibrozil, and omeprazole. The patient was treated with prednisone and methotrexate followed by intravenous immunoglobulin, which resulted in normalization of creatinine kinase levels and resolution of symptoms after 14 weeks. It is unclear if polymyositis was triggered by interaction of the statin with omeprazole and/or gemfibrozil, or if it developed secondary to long-term use of atorvastatin only.

Keywords: Autoimmune; Myopathy; Statins
Statins are an extensively used class of drugs. According to a report from the IMS Institute for Healthcare Informatics, close to 94 million individual prescriptions for generic simvastatin were issued in 2010. Myopathy is an uncommon, but well-described side effect of statin therapy. Drug interactions increase the risk of statin-associated myopathy by as much as ten-fold. Statin-associated myopathies generally resolve after discontinuation of the drug; however, in rare cases of statin-associated autoimmune myopathy, symptoms may persist or worsen after the drug is discontinued, requiring immunosuppressive therapy. Polymyositis is a rare side effect of statin therapy that is characterized by symmetric proximal muscle weakness and the persistence of symptoms even after discontinuation of the drug. Here we describe a patient with probable polymyositis in whom drug interactions may have played a role.

**Case Presentation**

A male patient, aged 59-years, presented in September 2011 with complaints of insidious onset of weakness, muscle soreness, and trouble climbing stairs for 5 months. The patient had symptoms in both upper and lower extremities, but his symptoms were more pronounced in the hip flexors. He described his symptoms as gradually worsening and denied any history of viral illness or fever.

The patient’s medical history was significant for ST-elevated myocardial infarction (STEMI) in April 2006, for which he had been treated with percutaneous coronary intervention (PCI) with drug-eluting stent placement and had been started on atorvastatin 20 mg daily; 3 months later gemfibrozil 600 mg twice daily was added. In September 2010, the patient had symptoms of
gastroesophageal reflux disease (GERD) and dysphagia, for which he underwent esophagogastrroduodenoscopy with findings of mild distal peptic esophagitis and relatively patent Schatzki’s ring. Balloon dilation of the esophagus was performed, and he was started on omeprazole 20 mg daily. He denied use of tobacco or alcohol. Other home medications included amlodipine, aspirin, nitrate, and metoprolol. Atorvastatin, gemfibrozil, and omeprazole were stopped in August 2011 secondary to muscle soreness. The patient had no known history of endocrinopathy, neurogenic disease, exposure to other myotoxic drugs, any symptoms suggestive of underlying malignancy, or family history of neuromuscular disease.

Upon physical examination, the patient scored 5/5 on strength of the upper extremities and 4/5 on strength of lower extremities (hip flexors, ileopsoas), indicating he was able to raise the leg only against slight resistance, but not against full resistance. The issue appeared to be localized to the ileopsoas, as strength in the quadriceps was 5/5. Sensation was intact for pin and vibration, and reflexes were intact. Physical exam was otherwise normal, including extraocular and facial muscles, and there was no evidence of rash.

The patient had been on atorvastatin and gemfibrozil for approximately 5 years without any muscle symptoms. Seven months after starting omeprazole, he began experiencing muscle soreness and weakness. The patient’s serum creatinine kinase (CK) level at the time of presentation was highly elevated at 10,554 U/L (normal 50–235 U/L) compared to a level of 700 U/L in April of 2006. Table 1 details CK levels over time. Atorvastatin, gemfibrozil, and omeprazole were stopped, but the patient remained symptomatic with persistent CK levels >10,000 U/L that reached as high as 11,831 U/L in September, one month after presentation.
Magnetic resonance imaging (MRI) of the bilateral thighs was performed with intravenous administration of 20 ml gadolinium contrast agent. MRI with axial and coronal T1-weighted and short tau inversion recovery (STIR) sequences showed an abnormal mild increase in STIR signal and enhancement of the bilateral posterior thigh musculature, especially involving the bilateral hamstring and adductor muscles (figure 1), compatible with polymyositis, rhabdomyolysis, or an inflammatory neuropathy. A biopsy was obtained from the left posterior thigh muscle and revealed findings consistent with acute necrotizing myopathy, including countless necrotic and regenerating fibers (figures 2A and B) with focally prominent perivascular perimysial and endomysial inflammation composed of mature lymphocytes and macrophages (figure 2C). Immunohistochemical studies revealed that the lymphocytic infiltrates were composed of CD3- and CD8-positive cytotoxic T-cells (figure 2D and E). A major histocompatibility complex (MHC)-1 immunostain showed patchy membranous reactivity with accentuation around necrotic fibers and foci of perivascular inflammation (figure 2F). Laboratory results were negative for all autoantibodies tested including antinuclear antibody (ANA), ds-DNA, chromatin, ribonucleoprotein P, Sjogrens antibodies SSA and SSAB, centromere B, SM, SnRNP, SCL-70, and Jo1. Antibodies to hepatitis A, B, and C were also negative. Testing for anti-SRP and other non-Jo1 anti-synthetase autoantibodies was not performed.

Based on the muscle biopsy report and clinical presentation, the patient was suspected of having polymyositis and was started on prednisone and methotrexate. Oral prednisone was started at 80 mg/day for 7 weeks, then gradually tapered by 10 mg/month for 2 months followed by 5 mg/month for 3 months, which still continues. Oral methotrexate was started at 15–20 mg/week for 12 weeks, and the patient continues to receive 20 mg/week, subcutaneously. Prednisone and
methotrexate administration resulted in a decrease in serum CK levels from 11,831 U/L in September 2011 to 4772 U/L in October 2011, but muscle weakness did not subside. Three doses of intravenous immunoglobulin of 40 g, 60 g, and 60 g were administered over a one-month period in November and December 2011, resulting in a further decrease in serum CK levels to 778 U/L. The patient tolerated the medications well without side effects. By 14 weeks of treatment, CK levels had fallen to within the normal range (180 U/L), and the patient’s muscle weakness had disappeared such that strength in the hip flexors (ileopsoas) had returned to 5/5. At last follow-up after 24 weeks of treatment, his serum CK levels were within normal range at 65 U/L, and he was continuing on prednisone 20 mg/day orally and methotrexate 20 mg/week subcutaneously.

**Discussion**

Statins are used extensively to lower serum cholesterol as well as for primary and secondary prevention of coronary events and stroke.\(^4\) Myopathy is a rare but well-documented side effect associated with statin use and ranges from myalgia to myositis to overt rhabdomyolysis.\(^5,6\) Onset of muscle symptoms usually occurs within weeks to months of statin therapy initiation, and symptoms resolve within months of drug discontinuation.\(^7\) Several possible mechanisms of statin-associated muscle toxicity have been proposed, but the answer remains unclear. Theories suggested are primarily downstream of HMG-CoA reductase inhibition and include instability of skeletal muscle cell membranes by blocking of cholesterol synthesis,\(^8\) decreased synthesis of coenzyme Q10 and mitochondrial dysfunction,\(^9,10\) inhibition of GTP-binding regulatory proteins, and impairment of muscle’s ability to appropriately recover from physical exertion.\(^11,12\) While statins can cause myopathy during monotherapy, many cases are associated with drug
interactions. Drugs of particular concern include CYP3A4 isoenzyme-dependent drugs such as macrolide antibiotics, azole antifungals, and cyclosporine,\textsuperscript{12-14} drugs that interact with the glucuronidation pathway such as gemfibrozil,\textsuperscript{12} and drugs that interact with the p-glycoprotein efflux system such as proton pump inhibitors (PPIs).\textsuperscript{2,15-17} There are many reports in the literature of an increased incidence of myopathy with use of statins combined with gemfibrozil,\textsuperscript{18} as well as case reports of rhabdomyolysis and polymyositis associated with the use of omeprazole and other PPIs in combination with statin therapy.\textsuperscript{2,19,20}

In addition to the self-limited, toxic myopathy associated with statins, there are reports of statin-associated inflammatory myopathies including polymyositis, dermatomyositis, and necrotizing autoimmune myopathy that are characterized by elevated CK levels and proximal muscle weakness during or after statin use that persists even after discontinuation of the drug. Symptoms improve with immunosuppressive therapy, but many patients require treatment with multiple agents including steroids, steroid-sparing agents, and intravenous immunoglobulin,\textsuperscript{21} as in the case reported here. Diagnostic criteria for polymyositis include: symmetric proximal muscle weakness progressing over weeks to months; muscle biopsy demonstrating myofiber necrosis, phagocytosis, regeneration, variation in fiber diameter, and an inflammatory exudate; elevation of serum skeletal muscle enzymes; and electromyography showing low-amplitude, small, polyphasic motor units.\textsuperscript{22} The patient presented here meets three of the four diagnostic criteria, making this a probable case of polymyositis.

The histopathological findings described in the muscle biopsies of patients with polymyositis show necrotic and regenerating muscle fibers with a characteristic endomysial inflammatory
infiltrate rich in CD8 T cells and increased MHC-1 expression on non-necrotic muscle fibers.\textsuperscript{3} Although the clinical presentation of several inflammatory myopathies is similar, histology allows for differentiation of the diagnoses. Histologically, dermatomyositis is similar to polymyositis, except that the infiltrate is composed primarily of CD4 T cells.\textsuperscript{3} The characteristic rash is a further distinguishing characteristic. In contrast, necrotizing autoimmune myopathy is notable for the relative absence of inflammatory cells on histology.\textsuperscript{3}

Autoantibodies are frequently detected in patients with autoimmune myopathies. The patient presented here was negative for autoantibodies against ANA and Jo1, but he was not tested for antibodies against HMG-CoA, anti-SRP, and non-Jo1 anti-synthetase antibodies. Antibodies against HMG-CoA are more typical of necrotizing autoimmune myositis than polymyositis.\textsuperscript{22} Patients with anti-SRP antibodies are very rarely found to have collections of inflammatory cells on biopsy, and those with anti-synthetase antibodies tend to present with additional anti-synthetase syndrome-specific clinical features in addition to myopathy;\textsuperscript{22} however, due to lack of testing, these autoantibodies cannot be ruled out.

The presence of inflammatory CD8 T cells upon biopsy and the overall clinical picture of widespread necrosis and regeneration is most consistent with polymyositis. In the case of statin-associated inflammatory myopathies, symptoms may occur years after the start of statin treatment or even after discontinuation of statin use.\textsuperscript{24} In the case presented here, we are not sure if polymyositis was triggered by interaction with omeprazole and/or gemfibrozil, or if it developed secondary to long-term use of atorvastatin only. Of note, the patient’s CK levels were relatively high (700 U/L) 5 years before presentation, which may have indicated an underlying
polymyositis that was later exacerbated or unmasked by use of a statin, gemfibrozil, and omeprazole. As previously mentioned, omeprazole and gemfibrozil have been reported to cause toxic myopathy with statins, but the incidence of polymyositis is rare.

**Conclusion**

Polymyositis is a rare but serious statin side effect that can occur years after initiation of therapy and persists after discontinuation of the drug, in contrast to the self-limiting toxic myopathy more frequently associated with short-term statin use. It is important to differentiate statin-associated inflammatory myopathies from other self-limited myopathies, because the disease will not subside following discontinuation of the drug, and treatment often requires multiple immunosuppressive agents. Drug interactions increase the risk of statin-associated toxic myopathy, and while no risk factors for statin-associated polymyositis have been established, drug interactions may play a role and deserve further investigation.

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References


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Table 1. Creatinine Kinase (CK) Levels Over Time

<table>
<thead>
<tr>
<th>Date</th>
<th>CK (U/L)</th>
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<tbody>
<tr>
<td>April 2006</td>
<td>700</td>
</tr>
<tr>
<td>August 2011</td>
<td>10,554</td>
</tr>
<tr>
<td>September 2011</td>
<td>11,831</td>
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<tr>
<td>October 2011</td>
<td>4,771</td>
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<tr>
<td>November 2011</td>
<td>3,137</td>
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<td>December 2011</td>
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<td>January 2012</td>
<td>118</td>
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<td>March 2012</td>
<td>65</td>
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**Figure Legends**

**Figure 1.** MRI of lower extremities showing T1 flat saturation sequences. (A) Coronal section and (B) axial section showing inflammation in the posterior (bottom of image) thigh muscles with relative sparing of the anterior and medial compartments. Arrows indicate areas of inflammation.
Figure 2. Histopathological analysis of left posterior thigh biopsy with evidence of acute necrotizing myopathy. (A) Active necrotizing myopathy with concurrent areas of regeneration, H&E, 100x. (B) Higher power detail of areas of myofiber regeneration, H&E, 400x. (C) Perimysial vascular lymphocytic infiltrates, H&E, 400x. (D) CD3 immunohistochemical stain highlights T-cells in perivascular infiltrates, DAB chromogen with hematoxylin counterstain, 100x. (E) CD8 immunohistochemical stain highlights cytotoxic T-cells. DAB chromogen with hematoxylin counterstain, 100x. (F) MHC-1 immunohistochemical stain highlighting diffuse positive membranous reactivity, with accentuation in necrotic and regenerating fibers. DAB chromogen, 200x.