Case report

Severe statin-induced autoimmune myopathy successfully treated with intravenous immunoglobulin

Cansu Güngör,1 Udo Carl Wieshmann2

SUMMARY

Statin-induced autoimmune necrotising myopathy causes a severe progressive muscle weakness even when the statins are discontinued. First-line treatment is usually with high dose steroids followed by immunosuppressants, but this is often ineffective and there is a high risk of side effects. We describe a diabetic patient who had a very severe statin-induced autoimmune myopathy. He made a full recovery with regular intravenous immunoglobulin (IVIg) infusion in relatively low dose (55g the first day followed by 50g/day the second and third day, subsequently he was given 50g/day for 3 days every 6 weeks). His symptoms relapsed when the IVIgs were discontinued for 28 weeks but remitted again following recommencement of IVIg infusions (50g/day for 3 days every 7 weeks). Our case suggests IVIgs are an effective and well tolerated alternative to steroids and immunosuppressants.

BACKGROUND

Statins are among the most widely prescribed drugs. Myopathy is a rare but severe side effect of statins. Statin-induced necrotising autoimmune myopathy with anti 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) antibodies is the most severe form of statin-induced myopathy. Unfortunately the weakness progresses even when the statins are discontinued. Treatment with steroids and immunosuppressants is often ineffective. We describe a patient with statin-induced autoimmune myopathy who was successfully treated with intravenous immunoglobulin (IVIg).

CASE PRESENTATION

A 65-year-old man was referred with proximal leg weakness. The weakness started about 2 months ago and got progressively worse. The patient had difficulties climbing stairs, getting up from the squatting position, walking or even putting on his trousers. He had started using a wheelchair outside the house. He denied having any associated pain, and upper limbs as well as speech and swallowing were uninvolved. He had longstanding mild sensory symptoms in both feet but no autonomic symptoms. He had also lost around 6 lb (about 2.72 kg) in weight in the previous month. He had been diagnosed with hypertension and type 2 diabetes mellitus 10 years ago and had taken atorvastatin 10 mg and aspirin 75 mg alongside antihypertensive (ramipril and amlodipine) and anti-diabetic (metformin, dapagliflozin, sitagliptin and gliclazide) medication since then.

On examination fasciculations and wasting were noticed in both quadriceps muscles. The muscular tone was normal. There was proximal weakness in both legs. Trendelenburg’s and Gower’s signs were positive. The knee jerks were brisk. Sensation was intact except loss of vibration sense up to the tibial plateau bilaterally which had been known to the patient for many years. Examination of the cranial nerves and the upper limbs was unremarkable.

INVESTIGATIONS

The creatine kinase (CK) level was raised up to 4292 U/L which was more than 10 times the normal range of 40–320 U/L. The alanine transaminase (ALT) was raised to 234 U/L (normal range: 11–55 U/L). Alkaline phosphatase and bilirubin were normal. Autoantibodies against HMG-CoA reductase were positive. Haemoglobin A1C was 51 mmol/mol. Full blood count, thyroid function tests, renal profile, vitamin D and B12 were all normal.

Electromyography (EMG) showed active denervation (fibrillation and positive sharp waves) and fasciculation potentials along with chronic denervation/re-innervation motor unit patterns (higher amplitude, increased duration and reduced interference patterns) from the cranial muscle, thoracic, paraspinal as well as upper and lower limb muscles.

DIFFERENTIAL DIAGNOSIS

The clinical findings with proximal weakness and preserved reflexes, and the raised ALT and CK were in keeping with an acquired myopathy. The EMG showed polyphasic short muscle action potentials in keeping with a myopathy. Spontaneous activity in the form of fasciculations (both clinically and on EMG) and positive sharp waves (on EMG) is seen in patients with inflammatory myopathies and must not be confused with motor neuron disease.1 A markedly raised CK like in our case is not compatible with motor neuron disease.2 The positive autoantibodies against HMG-CoA reductase confirmed the clinical diagnosis of ‘statin-associated autoimmune myopathy’.3 4

TREATMENT

Atorvastatin was stopped but the patient’s muscle weakness deteriorated in his legs and also spread to his arms making immune-modulating treatment necessary.5 It is normally recommended to start the

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therapy with oral prednisolone (1 mg/kg body weight) and to combine it with an immunosuppressive agent-like methotrexate, azathioprine or mycophenolate mofetil.8

However, due to potential side effects of glucocorticoids on the patient’s diabetes mellitus, an alternative approach was considered in this case. Since a positive outcome with first-line therapy IVIg (gamunex 10%) treatment was already reported for three diabetic patients,6 we decided for monotherapy with IVIg for the patient. A total dose of 155 g IVIg, equivalent to 1.6 g/kg of body weight, was administered over 3 days. He received 55 g on the first day and 50 g on the second and third day. After the first administration, he developed a headache but he tolerated the second and third infusion and all subsequent infusions without side effects. He was ‘feeling stronger’ despite initial headaches and a lack of significant improvement of the CK. Nonetheless, since CK improvement was not expected before two or three treatment courses with IVIg, the treatment was continued every 2–3 weeks while slightly reducing the dose to 150 g because of the headache. After the third course of treatment, the CK in serum dramatically fell to a mildly elevated level. Following the fourth administration, CK was stable around 500 U/L and weakness in his limbs had been greatly improved.

OUTCOME AND FOLLOW-UP

At the first follow-up, 7 weeks after the fourth round of IVIg, the patient nearly regained full strength in his legs and CK did not show any further increase. Taking potential severe adverse effects of IVIg into account, such as anaphylaxis, Stevens-Johnson syndrome, hypotension, myocardial infarction, cytopenia, haemolysis, pulmonary oedema, it was decided to stop IVIg infusions, and to solely monitor the symptoms and serum CK level instead.

However, 28 weeks later the patient’s weakness returned and CK was elevated over 1500 U/L. Treatment with IVIg was hereupon resumed at 50 g/day for 3 days every 7 weeks and resulted in a quick regression of the symptoms and CK which, already after the second course of IVIg, almost returned to the level before the flare-up. Following the next administration, the seventh in total, the patient reported almost full recovery of the muscle strength and his CK level was back in the normal range for the first time since the patient had been admitted to our clinic 1.5 years prior (figure 1).

Figure 1  Creatine kinase level (U/L) in patient serum over time, from the admittance (01/2018) to hospital until the latest round of IVIg infusion (11/2019), red arrowheads: rounds of IVIg infusions, spread over 3 days in each round (50 g/days), CK, creatine kinase; IVIg, intravenous immunoglobulin.

DISCUSSION

Muscle related adverse effects are common side effects of statins ranging from muscle pain and asymptomatic CK elevation to muscle weakness to rhabdomyolysis.9 Statin-induced myopathy affects 1.2 per 10 000 patients and often manifests itself after 6 months of use.9 Although it is of non-immune and self-limiting character, in 2007 Needham et al described eight cases where the myopathy persisted or progressed following the cessation of statins, sarcolemmal major histocompatibility complex (MHC)-I expression was upregulated and strength in proximal muscles was recovered with an immunosuppressive therapy with prednisolone and methotrexate, indicating an immunological nature.10

Statin-induced autoimmune myopathy is a rare side effect of middle to long-term statin use which is seen in approximately 2–3 per 100 000 patients treated with statins.7 The average age of onset is around 65 years of age11 12 and there is no sex predominance.11 It manifests itself as symmetrical proximal weakness which begins after several years of statin taking and may lead to immobility if untreated. The mean duration of statin use prior to symptom onset is approximately 41 months.11 Moreover, it is marked by significantly increased CK in blood, usually more than 10 times the upper limit of the normal range5 and autoantibodies against the enzyme HMG-CoA reductase.4 Our patient also presented with severely decreased strength of hip flexion, CK over 3000 U/L and was anti-HMG-CoA reductase antibodies positive.

Muscle oedema and necrosis can be detected on MRI12 13 while abnormal spontaneous activity in the form of fibrillation potentials, positive sharp waves and myotonic or pseudomyotonic discharges is commonly found on EMG14 along with small-amplitude motor-unit potentials.15 Long-duration, high-amplitude motor unit patterns like in our patient are seen in (sub)acute and chronic stages of autoimmune myositis.15

Muscle biopsies, which are ordinarily considered to be part of the standard workup, demonstrate degeneration, necrosis and regeneration of myofibres14 16 and cellular infiltrates that are mainly constituted by macrophages and focused in endomyial and perivascular areas.17 CD4+ and CD8+ lymphocytes as well as CD123+ plasmacytoid dendritic cells were also present in some samples.15 Besides, it is reported that MHC class I protein expression is up-regulated on the sarcolemma surface of non-necrotic muscle fibres scattered diffusely throughout the endomyium.10 16 We did not proceed to a muscle biopsy, because of the distinctive combination of progressive bilateral proximal weakness with wasting, raised CK level and anti-HMG-CoA antibodies in blood.

Anti-HMG-CoA reductase-associated myositis constitutes a subtype of immune-mediated necrotising myopathies alongside antisignal recognition particle-associated myositis. The anti-HMG-CoA reductase antibodies are associated with prior statin use in two thirds of the cases, especially for those over 50, whereas anti-HMG-CoA reductase positive patients without statin exposure tend to be younger at the disease onset and have higher CK levels.1 A study found that anti-HMG-CoA reductase myopathy is more likely in the patients having type 2 diabetes mellitus or using atorvastatin (comparing with rosuvastatin and simvastatin).18 Both of these risk factors were also present in our case.

The exact pathogenesis of HMG-CoA autoimmunity is yet to be known. Nevertheless, class II HLA allele DRB1*11:01 is associated with a higher risk of developing anti-HMG-CoA reductase antibodies18 and may explain why some people are susceptible to this kind of autoimmune myopathy. Furthermore, the HMG-CoA


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reducease expression is increased in the presence of statins and in the regenerating muscle cells.5 These findings may suggest that overexpression of HMG-CoA reductase induced by statins triggers an autoimmune response in susceptible subjects. As muscles regenerate more due to the damage by autoantibodies, high levels of HMG-CoA reductase in regenerating muscles would maintain the autoimmune reaction in absence of statins as well. It would be interesting to know whether the levels of HMG-CoA antibodies fell along with CK levels with IVlg treatment. Unfortunately, we are unable to report whether the autoimmune titres for this research from any funding agency in the public, commercial or for-profit sectors.

Learning points

- Severe statin-induced autoimmune myopathy can occur after years of treatment with statins.
- Cessation of statins is not sufficient; immune-modulating treatment is required.
- Intravenous immunoglobulin is an alternative to steroids but long-term treatment is required.

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