68-year old woman with refractory cutaneous dermatomyositis

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ABSTRACT
Dermatomyositis is an idiopathic inflammatory disorder of the muscles associated with characteristic cutaneous findings. Herein we report a 68-year old woman who presented with dermatomyositis associated with painful vasculitic lesions on both hands, refractory to conventional treatment. Steroids, topical tacrolimus, antimalarials and intravenous cyclophosphamide were tried with no beneficial effect. Rituximab was also administered with no initial effect; soon afterwards, intravenous immunoglobulin was administered with good results. Some cases of cutaneous dermatomyositis may require trials of different therapies to identify the treatment regimen that produces satisfactory disease control.

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CASE PRESENTATION
A 68-year old woman with a previous history of hyperthyroidism presented with fever, rash, hair loss, muscular weakness, dysphagia, dry cough and arthralgia. She denied weight loss or loss of appetite. She had undergone total hysterectomy 6 months ago for HPV-related cervical dysplasia without features of malignancy. She was a non-smoker and was treated with thiamazole for hyperthyroidism.

Clinical examination revealed diffuse facial erythema, a heliotrope eyelid rash and poikiloderma in a photosensitive distribution including the upper chest, in a “V-neck” configuration, the upper back (shawl sign) and the lateral thighs (holster sign). Gottron's papules were noted overlying both elbows, metacarpophalangeal, proximal and distal interphalangeal joints bilaterally. The palmar and lateral surfaces of the fingers were rough and cracked. She had painful ulcers of the buccal mucosa; especially the hard palate. Painful vasculitic lesions were noted on the palmar aspects of her fingers on both hands, together with an ulcer. The lesions were so painful that the patient was unwilling to shake hands (Figure 1A). She had diffuse alopecia and arthritis of the wrists and metacarpophalangeal joints bilaterally. Neurological examination disclosed bilateral proximal upper and lower extremity weakness (muscle strength: 3/5) with preserved tendon reflexes and sensation. There were fine inspiratory crackles at both lung bases. There was no lymphadenopathy or organomegaly.

Laboratory work-up showed an elevated CRP (2.82mg/dL, normal < 0.8 mg/dL) and ESR (102 mm/h), leukopenia...
nia (WBC: 3850) and normochromic, normocytic anaemia (Hb=10.2 g/dL). Creatine kinase and thyroid-stimulating hormone were within normal limits. Testing for ANA was positive (1/640); anti-Ro was also positive and anti-Jo1 was negative. Anti-MDA-5, anti-NXP2 or anti-TIF1γ autoantibodies were not tested. Levels of C3 and C4 were normal. A diagnosis of dermatomyositis was made, and the patient was started on high dose steroids (prednisolone 1mg/kg).

One month later, the patient presented at the rheumatology clinic without fever or arthritis, with improved muscle strength. Facial rash and poikiloderma were improved, but she still complained of troublesome vasculitic lesions on her hands. Chest and abdominal imaging, mammography and gastrointestinal endoscopy revealed no features of malignancy, and muscle biopsy was negative.

Skin biopsy disclosed myxoid degeneration of the dermis with mucus deposition and perivascular and intermediate lymphoplasmacytic infiltration together with mucus deposition around the vessels. The inflammatory infiltrate was located on the upper and lower dermis. Moreover, there was focal dropscial degeneration of the stratum basale and few melanocytes on the dermis. These features are not pathognomonic, but can be seen in dermatomyositis. Her pulmonary function testing revealed a restrictive pattern (FVC: 75%, FEV1: 88%, FVC/FEV1: 1.17). Based on the patient’s clinical improvement, steroids were tapered and hydroxychloroquine 200mg OD was added. Ten days later, the patient presented urgently with a rash exacerbation and was admitted for a single methylprednisolone i.v. pulse (1 gr). Hydroxychloroquine dose was doubled, and tacrolimus cream was added. One month later she presented with no rash improvement and another i.v. pulse of methylprednisolone was given with improvement. On subsequent follow-up visits, all other dermatomyositis symptoms kept improving, apart from the vasculitic rash. Steroid tapering resulted in flares. Two monthly intravenous pulses of cyclophosphamide were given, resulting in an initial clinical improvement followed by a subsequent flare. A cycle of rituximab was administered (1 gr, two weeks apart) with initial improvement and then a subsequent flare again.

Based on the refractory nature of cutaneous dermatomyositis, intravenous immunoglobulin (IVIG) was administered at a dose of 2gr/kg divided in 5 daily courses every month. A few days before the third monthly IVIG cycle, our patient’s rash greatly improved (Figure 1B). Gottron’s papules and vasculitic lesions had nearly disappeared, as well as her poikiloderma lesions. Painful ulcers of the buccal mucosa healed, and new hair growth was evident in the scalp. Muscle strength was 5/5. She was given the third cycle of IVIG, steroids were tapered, tacrolimus cream was stopped, and hydroxychloroquine dose was reduced to 200mg once daily. She repeated a 2nd cycle of rituximab at 6 months after the initial course.

Today our patient is well on azathioprine (2mg/kg) and methylprednisolone 4mg once daily.

**DISCUSSION**

Dermatomyositis is an idiopathic inflammatory disease of the muscle, associated with cutaneous involvement. A careful work-up to exclude underlying malignancy is always mandatory. Patients who fail to respond to conventional therapies require more aggressive immunosuppressive or immunomodulatory treatment. Although a wide variety of medications have been used for the treatment of refractory cutaneous dermatomyositis, few formal studies have investigated the safety and efficacy of such agents. Patients who have failed conventional therapy (photoprotection, topical agents, hydroxychloro-
The authors declare no conflict of interest.

REFERENCES


