CASE REPORTS

JUVENILE DERMATOMYOSITIS: COMMON MANIFESTATIONS OF A RARE DISEASE

DERMATOMIOSITE JUVENIL: MANIFESTAÇÕES COMUNS DE UMA DOENÇA RARA

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ABSTRACT

Juvenile dermatomyositis is an autoimmune systemic vasculopathy, mainly characterized by chronic inflammation of the striated muscle and skin. The authors report the case of a 15-year-old boy presenting with a four-month history of myalgia and proximal muscle weakness on the upper and lower limbs. These symptoms were associated with heliotrope palpebral exanthema, erythema in the dorsum of the hands, Gottron’s papules, erythematous and petechial rash on the extensor face of the thighs, and mild dysphagia for solids. Blood tests revealed an increase in muscle enzymes and electromyography showed changes suggestive of severe acute myopathy. Intravenous methylprednisolone was initiated, followed by a combination regimen of prednisolone and methotrexate. Progressive dysphagia, cutaneous abnormality, and muscular strength improvement were noted. With this case, the authors intend to raise awareness of a rare disease with an essentially clinical diagnosis, presenting in most cases with characteristic manifestations that should be recognised.

Keywords: exanthema; juvenile dermatomyositis; myositis

RESUMO

A dermatomiosite juvenil é uma vasculopatia sistémica de natureza autoimune, caracterizada por inflamação crónica do músculo estriado e da pele. É apresentado o caso de um adolescente do sexo masculino com um quadro de mialgia e fraqueza muscular proximal dos membros superiores e inferiores com quatro meses de evolução, associado a exantema palpebral em heliotropo, eritema purpúrico no dorso das mãos, papulas de Gottron com atingimento peri-ungueal, exantema eritematoso nas coxas e disfagia ligeira para sólidos. O estudo analítico revelou aumento das enzimas musculares e a eletromiografia evidenciou alterações sugestivas de miopatia grave em fase aguda. O doente iniciou metilprednisolona endovenosa em bólus, seguida de um esquema de prednisolona e metotrexato, com melhoria progressiva da disfagia, força muscular e alterações cutâneas. Com este caso clínico, os autores pretendem alertar para uma doença rara com diagnóstico essencialmente clínico, que se apresenta na maioria dos casos com manifestações características que devem ser reconhecidas.

Palavras-chave: dermatomiosite juvenil; exantema; miosite

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INTRODUCTION

Juvenile idiopathic inflammatory myopathies are a heterogeneous group of immune-mediated diseases. Juvenile dermatomyositis (JDM) is the most common clinical phenotype of myositis in children and adolescents, but remains a rare disease. Most studies suggest that JDM is an autoimmune angiopathy, resulting from exposure of genetically susceptible individuals to certain environmental triggers. JDM has an incidence of 0.55 cases per year and a prevalence of 6 in 100,000 inhabitants in Europe. Mean age of onset is 7.5 years and girls are mostly affected. Diagnostic criteria for JDM were established by Bohan and Peter in 1975 and include pathognomonic exanthema, proximal muscle weakness, raised muscle enzymes, myopathic changes on electromyography, and typical muscle biopsy. Current practice revealed the need of broadening classification criteria by incorporating new exams, such as magnetic resonance imaging (MRI), ultrasound, capillaroscopy, and myositis autoantibodies. These criteria are included in more comprehensive recommendations recently established by the European League Against Rheumatism/American College of Rheumatology (EULAR/ACR). Corticosteroids, as prednisolone, are first-line agents in JDM treatment. However, recent studies suggest superior efficacy of a combined regimen of prednisolone plus methotrexate (MTX). Cyclosporine A, intravenous human immunoglobulin, cyclophosphamide, azathioprine, mycophenolate mofetil, hydroxychloroquine, tacrolimus, rituximab, infliximab, and autologous stem cell transplantation are possible alternatives used in the treatment of refractory disease.

JDM prognosis is variable, with an overall mortality rate of 2–3%. Approximately 24–40% of patients experience a monocyclic course, recovering and successfully discontinuing medication within two years. Around 25% have a polyphasic course, with complete remission and recurrence periods. The remaining 50% of patients have chronic disease.

CASE REPORT

A 15-year-old healthy boy was referred to the Pediatric Rheumatology Clinic due to progressive asthenia, myalgia, and muscle weakness on the upper and lower limbs, with proximal predominance, during the previous four months. Muscular symptoms worsened in the previous two weeks, with gait difficulty and muscle weakness in daily life activities. The patient additionally presented erythema of the dorsal surface of the hands together with malar rash, eyelid exanthema, and periorbital edema with one month of evolution. Two weeks later, he noticed a rash in the anterior surface of lower limbs. Importantly, the rash worsened with sun exposure. The patient also mentioned mild dysphagia for solids with a week of evolution. He denied fever, arthralgia, anorexia, weight loss, hematemesis or melena, abdominal pain, respiratory symptoms, or Raynaud’s syndrome. On physical examination, the boy had good general condition, with stable vital signs. He presented palpebral exanthema in heliotrope associated with periorbital edema (Figure 1) and purpuric erythema in the dorsum of the hands, periungual telangiectasias and Gottron’s papules on metacarpophalangeal and interphalangeal joints (Figure 2), and erythematous and petechial rash on the extensor face of the thighs. Additionally, he presented symmetrical reduction of proximal muscle strength in upper and lower limbs and positive Gower’s signal, with Childhood Myositis Assessment Scale (CMAS) score of 28.

Laboratory tests revealed aspartate aminotransferase 267 U/L (Normal values 4–35), alanine aminotransferase 175 U/L (4–26), lactate dehydrogenase 777 U/L (135–225), creatine kinase 6,390 U/L (26–174), myoglobin 759 ng/mL (28–72), and aldolase 55.7 U/L (0–7.6). Blood count and sedimentation rate were normal, as were C-reactive protein and renal function. Serological tests (Cytomegalovirus, Rubella, Toxoplasmosis, Herpes Simplex I and II, Mycoplasma pn, Chlamydia pn, Legionella pn, Parvovirus, Epstein-Barr virus, Borrelia burgdorferi, and Hepatitis B and C) were all negative. Autoimmunity study revealed normal C3 and C4, 1/80 title of antinuclear antibodies (ANA) with mottled pattern. Anti-neutrophil cytoplasmic antibodies (ANCA), myositis-specific antibodies, anti-
Mi-2, anti-SRP, anti-Jo-1, anti-PL-12, anti-PL-7, anti-OJ, anti-EJ, as well as myositis-associated antibodies, anti-Scl, anti-Ro-52, and anti-Ku were all negative. Electromyography (EMG) revealed spontaneous denervation activity on examined muscles, with typical features of severe acute myopathy. Electrocardiogram, echocardiogram, chest radiograph, and functional respiratory tests were normal. Capillaroscopy showed architectural disorganization, giant capillaries, capillary loss, and angiogenesis.

Juvenile dermatomyositis diagnosis was established and the patient was treated with intravenous methylprednisolone for five days (30 mg/kg/day) during hospitalization. Muscle strength improvement was observed, together with slight improvement of cutaneous manifestations and dysphagia. The patient was discharged with oral prednisolone 1 mg/kg/day, calcium, and vitamin D. Sun protection was recommended and physiotherapy was started. Oral methotrexate 12.5 mg/m² (20 mg) was added two weeks after. One month after starting the combined regimen, almost complete muscle strength recovery (CMAS score of 48) as well as dysphagia and skin rash improvement were observed, with no evidence of calcinosis or other complications. Six months later, methotrexate was changed to subcutaneous administration for therapeutical optimization. Regimen was progressively reduced and after eight months of treatment symptoms completely resolved (CMAS score of 52).

The patient completed two years of combined treatment, without disease relapse. After three years of follow-up, he remains asymptomatic, without treatment, and is followed as an outpatient in the Pediatric Rheumatology Clinic.

**DISCUSSION**

The authors presented a clinical case of JDM diagnosed at 15 years of age in a male patient. In JDM, symptom onset is more frequent at earlier ages and prevalence is higher in females. In this case, no putative triggering factors were identified.

Diagnostic criteria for JDM are essentially based on clinical manifestations, according to the EULAR/ACR Classification Criteria for Adult and Juvenile Idiopathic Autoinflammatory Myopathies. This patient presented the characteristic heliotrope eyelid cutaneous rash associated with periorbital edema, as well as Gottron’s papules on metacarpophalangeal and interphalangeal joints. He also displayed proximal and symmetrical muscular weakness at shoulder and hip level, with difficulty in standing up, as confirmed by CMAS score.

Furthermore, complementary diagnostic tests revealed an increase in muscle enzymes and EMG changes suggestive of myopathy and denervation. Presently, EMG is not mandatory to establish diagnosis, and other exams, as total-body MRI, can be used to corroborate JDM diagnosis and help monitor disease activity. Muscle biopsy is also not mandatory to establish diagnosis, being mainly recommended when presentation is atypical or diagnosis is challenging, and was not performed in the present case. Capillaroscopy is increasingly used when severe capillary abnormalities, such as giant capillaries with intense neoangiogenesis, are common.

Although the role of antibodies in JDM pathogenesis has not been clarified, its value is increasingly acknowledged due to association with specific phenotypes, complications, and prognosis. Around 70% of patients are estimated to present myositis-specific or myositis-associated antibodies. In this patient, all such autoantibodies were screened and negative. Regarding other autoantibodies analyzed, 1/80 titer for ANA should be highlighted, although ANA values are variable in JDM.

Moderate-to-severe striated muscle damage, with limitation of daily activities and dysphagia prompt hospitalization for intravenous corticosteroid therapy. In the present case, combined prednisolone and oral MTX regimen was started early in treatment course, with muscle weakness improvement and cutaneous attainment partial improvement. Due to dysphagia persistence, MTX was changed to subcutaneous administration, with complete symptom resolution. This would have been the best option from the beginning, and the patient should have been initially treated with prednisolone and subcutaneous MTX.

**CONCLUSION**

This clinical case illustrates the challenging JDM diagnosis. Despite the condition’s rarity, JDM diagnosis remains essentially clinical, often presenting with typical clinical disease manifestations. It is therefore crucial to consider JDM in the differential diagnosis of pediatric myopathies and cutaneous rashes. Early recognition and treatment improve disease prognosis, preventing onset of calcinosis and other major complications.

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