Case report

Antisynthetase Syndrome: two case reports and literature review

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A B S T R A C T

Antisynthetase Syndrome (ASS) is characterized by myositis, Raynaud’s phenomenon, fever, interstitial lung disease, mechanic’s hands and arthropathy associated with the presence of antibodies against tRNA synthetase, especially anti-Jo-1. This article aims to review the literature on ASS and report two cases where the first is a patient with polymyositis who developed subluxation on the proximal interphalangeal joint of bilateral first right finger after a few years of the disease, associated with pulmonary manifestations and positive anti-JO-1. In the second case, we present a patient with dermatomyositis, who developed a subluxation of the two first fingers, anti-Jo1 positive and chest CT changes, but without clinical evidence of pulmonary involvement. These cases reveal the importance of performing early diagnosis. The authors describe two cases of this rare syndrome, emphasizing the severity of interstitial lung disease and arthritis.

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Síndrome Antissintetase: relato de dois casos e revisão da literatura

R E S U M O

A Síndrome Antissintetase (SAS) é caracterizada por miosite, fenômeno de Raynaud, febre, doença pulmonar intersticial, artropatia e mãos de mecânico associados à presença de anticorpos contra a sintetase do tRNA especialmente anti-Jo-1. Este artigo tem como objetivo revisar a literatura sobre SAS e relatar dois casos, sendo o caso 1 de uma paciente com Polimiosite que desenvolveu, após alguns anos de doença, subluxação da articulação interfalangeana proximal do primeiro quirododáctilo direito, associada a manifestações pulmonares e anti-Jo-1 positivo. O caso 2 é de uma paciente com Dermatomiosite que evoluiu com subluxação dos dois primeiros quirodáctilos, anti-Jo-1 positivo e alterações pulmonares intersticiais na

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Introduction

Idiopathic inflammatory myopathies are a heterogeneous group of acquired muscular diseases characterized by various types and degrees of skeletal muscle inflammation. Polymyositis (PM) and dermatomyositis (DM) are the two major conditions making up this group of diseases. They are associated with serum antibodies related to particular clinical manifestations, therefore marking important clinical subgroups. The largest subgroup in muscular inflammatory diseases comprises the Antisynthetase Syndrome (ASS), firstly described by Marguerie et al. in 1990 as the triad polymyositis, diffuse interstitial lung disease, and self-antibodies to aminoacyl-tRNA synthetase (anti-ARS). It is currently characterized by the production of antibodies to tRNA synthetase, with anti-Jo-1 being the best known of these antibodies associated with myositis, interstitial lung disease, joint disease, fever, Raynaud’s phenomenon and “mechanic’s hands". This is a rare disease with an incidence in general population that is still unknown and a twofold higher prevalence in females. The interstitial lung disease (ILD) is the main morbidity and mortality determinant in the syndrome.

Case 1

A white craftswoman aged 61 with Hashimoto’s thyroiditis was seen in the rheumatology clinic in January 2001 reporting some manifestations that were consistent with Raynaud’s phenomenon, but she did not have muscle weakness or joint pain. A number of laboratory tests were obtained and the results are as follows: LDH: 628 (RV: 115–225); CK: 1041 (RV: 96–140); aldolase: 20.6 (RV: 3–7); ALT: 20 (RV: 7–56); AST: 36 (RV: 5–40); rheumatoid factor (RF): 12 (RV = up to 20), antinuclear antibody (ANA): negative; anti-Jo-1: positive; chest X-ray: no abnormalities were seen. Corticosteroids were initiated and muscle enzymes were normalized 4 months later. In January 2002, the patient complained of dyspnea and the chest X-ray showed a bilateral lung base infiltrate, in addition to elevated muscle enzymes. In May 2002, a chest computed tomography (CT) was obtained and diffuse “ground glass” imaging was seen, in addition to a pulmonary function test exhibiting a restrictive pattern. At that time, the patient complained of fever. In 2004, she had an interstitial thickening of the right upper lobe and both bases on chest X-ray, with dense bands showing a residual fibrous appearance in the right lower lobe.

In 2007, due to a respiratory and laboratory worsening, a monthly pulse therapy with cyclophosphamide was initiated. In that very year, she had a synovitis at the first finger of the right hand, progressing to a bilateral proximal interphalangeal subluxation of the first finger. Hand X-ray: slightly reduced carpometacarpal joint space.

Case 2

A white female patient aged 78, born in Portugal and who retired as an embroiderer, presented with a classic dermatomyositis since 1980. She had used corticosteroids for 11 years, with symptom remission (proximal muscle weakness, heliotrope rash, Gottron’s papules, Raynaud’s phenomenon, joint pain, and skin rash). Remission was maintained. Five years earlier, she had presented a deformity with distal interphalangeal joint subluxation of the first fingers (Fig. 1). A hand X-ray in 2009 showed bilateral interphalangeal (IP) arthrosis signs from the 2nd to the 5th finger and at the thumb IP, unilateral rizarthrosis and right pyramidal osteophytes. Chest CT: slight reticular pattern, subpleural lines, and ground glass opacities in the lung periphery. Positive anti-Jo-1.

Discussion

The Antisynthetase Syndrome is characterized by myositis, interstitial lung disease, joint pain, arthritis, “mechanic’s hands”, fever, and Raynaud’s phenomenon. It is associated with self-antibodies to tRNA synthetase, mainly anti-Jo-1. In general, the myositis either precedes or is concomitant with the lung involvement, as in the first reported case, in which the patient only had Raynaud’s phenomenon, elevated enzymes (LDH, CK, and aldolase) and positive anti-Jo-1, manifesting dyspnea and chest CT changes with diffuse ground glass imaging and restrictive pattern on the PFT. The patient in case 2,
Raynaud’s phenomenon is early in 2/3 of patients and can precede the myositis in years. It is more frequently found in anti-Jo-1-positive patients.  

“Mechanic’s hands” are a hyperkeratosis, scaling, and fissure in fingertips and later aspects of the fingers, being more frequently found in the Antisynthetase Syndrome and interstitial lung disease.

A skin vasculitis has also been described. The heart involvement does not have a different prevalence from that in polymyositis/dermatomyositis. A mesangial proliferative glomerulonephritis has been observed, but it is rare and has a good prognosis. Regarding idiopathic inflammatory myopathies associated with a interstitial lung lesion, recent reports show a favorable response to prednisone: 30–40% of the patients had an improvement and 20–40% became stable for symptoms and pulmonary function. A small group of patients is resistant to corticosteroids. There are remission reports with azathioprine, methotrexate, cyclophosphamide, cyclosporine, tacrolimus or mycophenolate mofetil. A late relapse following an initial remission is more common in the Antisynthetase Syndrome.

Bruhlart et al. described a patient aged 57 with an Antisynthetase Syndrome treated by prednisone (high doses) and methotrexate 15 mg/week who had a good response. A severe clinical and laboratory relapse was seen three months later. Rituximab (2 x 1-g infusions) was introduced, with myositis and ILD improvement. Studies suggest that CD20 depletion therapy might be valuable in refractory Antisynthetase Syndrome.

Anti-Jo-1-positive myopathies have a worse prognosis than other inflammatory myopathies.

Tests for anti-Jo-1 should be obtained in patients with inflammatory muscle disease, as well as a screening for interstitial lung disease.

**Conflicts of interest**

The authors declare no conflicts of interest.

**References**


