CASE REPORT

An Anti-PM/Scl-75 Antibody-positive Japanese Woman Who Developed Inflammatory Myopathy

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Abstract:
A 69-year-old Japanese woman presented with mild muscle weakness of the neck and symmetrical proximal parts of the upper and lower limbs. Laboratory tests, needle electromyography, and a muscle biopsy revealed inflammatory myopathy with an apparent clinical classification of polymyositis and positive findings for anti-PM/Scl-75 antibody. This antibody is rare among Japanese populations, and most Japanese patients with the antibody are not classified with the inflammatory myopathy seen in polymyositis. The muscle biopsy also showed marked necrotic and regenerative fibers. We need to collectively investigate patients with the potential to develop this disease, and to identify any unique characteristics for Asian populations, including Japanese.

Key words: anti-PM/Scl antibody, inflammatory myopathy, Japanese, muscle biopsy, neurology

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Introduction

Anti-PM/Scl (polymyositis/scleroderma) antibodies (PM/Scl-75 and PM/Scl-100) have been recognized as a cause of myositis. These antibodies were first detected in patients with an overlapping syndrome of PM and systemic sclerosis (SSc) (1) and have also been recognized in cases of other autoimmune diseases, such as dermatomyositis (DM) (2).

The detection frequency of these antibodies is influenced by the ethnic background, and studies from Europe and North America account for the majority of published data, while studies from Asia remain rare. Some papers have reported anti-PM/Scl antibody-positive DM in Japanese patients (3) and anti-PM/Scl antibody-positive Japanese patients with collagen disease centered on SSc (4). However, a small number of reports have described cases that appear to represent with PM only. No consensus has been reached regarding the clinical characteristics of anti-PM/Scl antibody-positive Japanese patients.

We herein report the case of a Japanese patient with anti-PM/Scl-75 antibody-positive inflammatory myopathy in which the clinical classification appeared to be PM and also describe the myopathological findings.

Case Report

A 69-year-old Japanese woman was referred and admitted to our department with suspected myositis. She had histories of unexplained weight loss over three years and mild muscle weakness from several months before this presentation. She had no history of statin treatment and did not regularly consume alcoholic beverages. No history suggestive of myositis or congenital myopathy was elicited. She had a history of gastroesophageal reflux disease, atrophic gastritis, and glaucoma. A physical examination revealed mild muscle weakness confined to the neck and symmetrical, proximal parts of the upper and lower limbs. The muscles in those areas showed mild atrophy. No skin lesions, such as sclerosis, Gottron’s lesion, heliotrope rash, or mechanic’s hands, were evident. These skin findings were confirmed by a dermatologist (HT). No myalgia, arthralgia, or dyspnea was observed.

Laboratory tests showed the following: aldolase, 12.7 IU/L; erythrocyte sedimentation rate, 50.9 mm/h; C-reactive protein, 4.76 mg/dL; creatine kinase (CK), 1,081 IU/L; and Krebs von den Lungen (KL)-6, 589 IU/mL (normal range, <
Results for anti-aminoacyl transfer RNA synthetase, anti-DNA, anti-Jo-1, anti-Scl 70, anti-Sjögren's syndrome-A/B, anti-Sm, and anti-U1-ribonucleoprotein were all negative. Antibodies suggesting immune-mediated necrotizing myopathy (anti-signal recognition particle/anti-3-hydroxy-3-methylglutaryl coenzyme A reductase) were also negative. Anti-centromere antibody was positive. Anti-PM/Scl-75 antibody was qualitatively detected using a line blot assay. Negative results were obtained for anti-PM/Scl-100 antibody. Results for molecular human leukocyte antigen (HLA) typing on DNA showed DRB1*01 and DRB1*15.

No abnormalities were evident on magnetic resonance imaging of the upper and lower limbs (Fig. 1). However, needle electromyography of the right upper and lower limbs showed normal spontaneous activity and an abnormal myogenic pattern (early recruitment in voluntary contraction and motor unit potential with small amplitude). A muscle biopsy from the left rectus femoris revealed histological findings reflecting active muscle fiber necrosis and regenerative changes (Fig. 2). Reactive mononuclear cell infiltration was seen surrounding necrotic and regenerating fibers. Immunostaining showed the expression of human leukocyte antigen (HLA)-ABC and deposition of membrane attack complex (MAC) on the muscle fiber membranes. Perifascicular atrophy, fibers with internal nuclei, and endomysial fibrosis were not seen. No perivascular collections of inflammatory cells were apparent. The pathological findings were suggestive of immune-mediated necrotizing myopathy (IMNM), as confirmed by a neurologist specializing in muscle disease (IN). A whole-body search including upper gastrointestinal endoscopy and lower gastrointestinal endoscopy was performed, but no malignant tumors were observed. Chest computed tomography showed mild interstitial changes at the base of the lower lung fields bilaterally.

We initially considered a diagnosis of PM (the PM subset includes IMNM) based on the criteria for idiopathic inflammatory myopathy recently described by Lundberg et al. (5). However, our final diagnosis was inflammatory myopathy with an apparent clinical classification of PM, because the pathological features of anti-PM/Scl antibody-positive myositis have not yet been definitively confirmed, and the pathological features of anti-PM/Scl antibody-positive patients and typical IMNM are not identical (6). Oral prednisolone was initiated at 40 mg/day (1 mg/kg/day), and her muscle weakness was gradually improved. After tapering the dose of oral prednisolone, the patient was discharged from the hospital. The serum CK level just before discharge was 211 U/L. As of six months since her discharge, no recurrence has been noted.
Figure 2. A Hematoxylin and Eosin staining section of the rectus femoris shows mild to moderate variation in the fiber size. A few necrotic fibers and some regenerating fibers are apparent (A). Modified Gomori trichrome-stained sections do not show ragged-red fibers, fibers with rimmed vacuoles, or nemaline rods (B). Immuno-stained sections show the expression of human leukocyte antigen-ABC and deposition of membrane attack complex on the muscle fiber membrane (C, D). Some freezing artifacts are evident.

Discussion

We encountered a rare case of anti-PM/Scl-75 antibody-positive inflammatory myopathy in a Japanese patient in which the clinical classification appeared closest to be PM. Because the clinical symptoms, such as muscle weakness, were minor and no skin lesions were evident, the clinical findings alone did not indicate such a diagnosis. Laboratory findings, needle electromyography, and a muscle biopsy helped in the diagnosis. The prognosis in this case was favorable.

Previous studies of Japanese individuals have not reported patients diagnosed with inflammatory myopathy, IMNM, and PM in which anti-PM/Scl antibodies were detected (Table) (3, 4). Another two studies to detect anti-PM/Scl antibodies in Japanese patients with systemic autoimmune conditions found no positive cases of anti-PM/Scl antibodies (7, 8). For example, in Muro et al., only 9 of 600 patients were ultimately deemed positive for anti-PM/Scl antibodies, including 4 of 16 (25%) patients with undifferentiated connective tissue disease (UCTD), 3 of 126 (2.4%) patients with DM, 1 of 223 (0.4%) patients with SSC, and 1 of 88 (1.1%) patients with Sjögren’s syndrome (4). Two possible reasons for these results were considered. One reason is that anti-PM/Scl antibodies appear to be related to HLA-DRB1*0301 (9). This HLA type is uncommon in Japanese populations, with an estimated frequency of 0.14% (4), making larger-scale research difficult. Another reason is that, in order to investigate anti-PM/Scl antibodies, we generally must ask a commercial center in Japan (BML, Tokyo, Japan) to perform testing. Because of the costs incurred, some patients with inflammatory myopathy may not be examined.

Despite the fact that weight loss started over three years earlier, our case was not able to be diagnosed until our department was consulted because of the absence of skin findings and her relatively mild muscle weakness. A small portion of anti-PM/Scl antibody-positive patients develop PM only (10-15), and determining the clinical features in those cases can be difficult. Another study of 41 anti-PM/Scl-positive patients stated that muscle weakness was present in 37% initially but was ultimately confirmed in 93% of patients during follow-up (mean follow-up, 6.5±4.7 years) (6). Long periods from the onset to the recognition of muscle symptoms may be characteristic of anti-PM/Scl-positive patients. Previous series have warned of lung and esophageal involvement (16). We need to perform continuous, long-term observation concerning the existence of new complications such as dysphagia. However, patients with anti-PM/Scl antibody have been considered to show a good prognosis, with
a good response to steroids and relatively low rate of malignancy (6, 15). Our case also responded well to oral prednisolone, and no cancer was found.

The findings from the muscle biopsy were a point of interest in the present study. To our knowledge, there have been no reports of the muscle biopsy findings for anti-PM/Scl antibody-positive patients in Asian populations. Previous studies concerning the clinicoserologic and pathologic classifications of inflammatory myopathies have shown that 10 of 178 patients had anti-PM/Scl antibodies, with 9 of those 10 patients (90%) presenting with necrotic and regenerative fibers and 9 of the 10 patients (90%) presenting with the expression of major histocompatibility complex, class 1 (only sparse fibers or groups of fibers in 1 patient, on the surface of most/all fibers in 3 patients, and perifascicular or diffuse with reinforcement in the perifascicular region in 6 patients) (17). These features were consistent with the findings in our case. However, another study of four anti-PM/Scl antibody-positive cases suggested that fibrosis was significantly associated with anti-PM/Scl antibodies (18). A recently published paper showed that perivascular inflammation was the most commonly observed feature among anti-PM/Scl antibody-positive patients (6). Different histological features have been reported in the literature, and the pathological findings of anti-PM/Scl antibody-positive myositis have yet to be definitively established. Applying the current classification criteria to our case was difficult, and we diagnosed the patient with inflammatory myopathy. The present report was from an area in which the onset of anti-PM/Scl antibody-associated disease is rare. Further cases need to be accumulated and examined not only from a serological perspective but also from a pathological point of view.

Several limitations associated with the present study warrant mention. Testing for anti-PM/Scl-75 antibody was performed by a commercial center in Japan. The antibody titer was not known because the results were qualitative in nature. The reliability of measurements performed by external companies is not necessarily high, but the same measurement method has also been used in recently published research (6).

In conclusion, we encountered an anti-PM/Scl-75 antibody-positive Japanese patient who showed inflammatory myopathy with an apparent clinical classification of PM. We reached the eventual diagnosis using clinical findings, laboratory tests, needle electromyography, and a muscle biopsy. Physicians should be aware that patients with anti-PM/Scl antibodies may be present among those with suspected myositis who have not yet received a confirmed diagnosis.

The authors state that they have no Conflict of Interest (COI).

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