Association of Lupus with Spinal Muscular Atrophy and Suspected Bronchial Cancer

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Introduction

SLE is an autoimmune disease with multiple organ involvement. It may affect musculo-skeletal and nervous systems resulting in muscle weakness and atrophy. Classification criteria for the diagnosis of SLE were defined by the American College of Rheumatology. Spinal muscular atrophy (SMA) is characterised by degeneration of motor neurons of the spinal cord, which also may result in muscle weakness and hypotonia. Earlier, the diagnosis was based on electromyography (EMG) and muscle biopsy findings. Electromyographic characteristic of the disorder shows features of denervation with spontaneous activity of positive sharp waves, fibrillation, and occasional fasciculations. Motor unit action potential shows high amplitudes and long durations coupled with decreased recruitment. Histopathology of skeletal muscle usually shows atrophic fibres with islands of group hypertrophy, and severe loss of motor neurons in the spinal cord [1,2]. It is caused by homozygous disruption of the survival motor neuron 1 (SMN1) gene in form of deletion, conversion, or mutation. SMN1 is expressed in all tissues, but almost exclusively affects lower motor neurons. SMA is divided into four clinical types: severe type I, intermediate type II, mild type III, and adult-onset type IV has been added to include very mild disease. Patients with type IV disease typically have onset of weakness in the second or third decades of life [3-5].

Case Report

Hereby, we report the case of a female patient who had weakness of the flexor muscles of the pelvis at the age of 18. She was treated with high-dose corticosteroid at that time, as polymyositis was supposed. Due to the steroid therapy she became better and stronger, but was not symptom-free. Later the weakness progressed appearing in her lower and upper extremities, too. The EMG detected neurogenic failure; the electroneurography (ENG) showed axonal type of neuropathy. Based on these, SMA was suspected, but genetic examination could not be performed at that time.

When she was 30 years of old symmetric polyarthritis involving the small joints of the hands, butterfly rash, leucopenia, lymphopenia, ANA (3+ positive with homogenous pattern in 1:400 dilution of serum) and anti-dsDNA (349 WHO U/mL) positivity referred to SLE. Later grand mal epilepsy, pleurisy, pericarditis, and autoimmune haemolytic anaemia developed. Associating sicca syndrome was also diagnosed based on glandular signs and aENA, a-SS-A, a-SS-B positivity. Antiphospholipid syndrome was suspected as she had livedo reticularis, IIIrd degree aortic valve insufficiency, pre-termed ischemic heart disease, acute myocardial infarction (with positive troponin I test result and inferior hypokinesia on echocardiography), and multiple vascular lesions by cranial MRI. However, she was more times negative for anti-phospholipid antibodies (PL Ab) including anti-cardiolipin, anti-beta2-glycoproteins I and lupus anticoagulant. She was treated with steroid and azathioprine, and required cyclophosphamide cycles to control haemolysis. Aspirin prophylaxis was given after MI. Besides all of these, oesophageal dismotility and Raynaud’s phenomenon was also present. Capillary microscopy picture was characteristic for SLE and not for systemic sclerosis.

At the second year with SLE thoracic CT was performed because of effort dyspnoea and coughing. CT showed reticulo-nodular bunch and lymphadenopathy in the axillary regions. Mammography was negative. Findings were attributed to the consequence of smoking. Novel autoantibodies had not been detected.

At the age of 37 her muscle weakness worsened leading to muscle atrophy of the lower extremities with mild paraparesis. Activity of CPK was normal, but LDH was slightly elevated. To differentiate between myositis, steroid myopathy, SMA progression further investigations were performed. Muscle biopsy revealed a combination of discrete perivascular inflammation and neurogenic damage of muscle fibers (Figure 1 and Figure 2). The genetic testing detected the deletion of the SMN1 gene, confirming previous SMA diagnosis. Due to progressive muscle weakness, unusual in type IV SMA, and also weight-loss and anaemia appropriate examinations were performed to exclude paraneoplastic origin of symptoms.

Chest X-ray and pulmonary CT detected a round-shaped, spiculated infiltration in the lower lobe of the right lung. Broncho-fiberscopy was negative. Biopsy with a video assisted thoracoscopy (VATS) has not confirmed bronchial cancer. Histopathology findings were characteristic for postinfectious residues.

Figure 1: The variability of the fiber-diameter is more than as usual, there are some rounded, angular fibers as well. This picture is not typical for spinal muscular atrophy.
Conclusions

Lupus may cause a wide variety of organ manifestation, as in case of our patient. Although neuro-muscular symptoms were the most prominent in her case, those could be attributed mainly to an associated disease; spinal muscular atrophy. The co-existence of the two diseases is very rare and makes the differential diagnosis difficult because both may result in similar phenotype. Paraneoplastic syndrome may further complicate the differential diagnosis, as resulted in similar symptoms as well. However, the anti-Hu antibody, as antineuronal antibody could be as an indicating marker. In cases of anti-Hu related paraneoplastic syndromes usually associated with small cell lung cancer. About 20% have prominent lower motor neuron signs but almost always accompanying involvement of other areas of the nervous system [14,15]. Unusual clinical progression and non-specific histopathology as well as refractivity to adequate therapy suggested us to look for the third disease and focus on the regular exclusions of the malignancy. The most difficult is to find adequate intervention to treat such patients. Fortunately, malignancy was excluded by histology.

References