Bent Spine Syndrome: A Phenotype of Dysferlinopathy or a Symptomatic DYSF Gene Mutation Carrier

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Dear Sir,

A bent spine can result from paraspinal muscle weakness of different causes. The symptom has also been described as bent spine syndrome or camptocormia [1, 2]. More or less incomplete fatty degeneration of the erector spine muscles has been found in these cases [3, 4]. While camptocormia is an age- and disorder-related symptom, the primary degeneration of the erector spine muscles is regarded as a muscle disorder, and denominated as bent spine syndrome. Different etiology could result in paraspinal muscle weakness, such as inflammatory myopathies [5, 6], metabolic disorders [7, 8] and, rarely, genetically determined degenerative muscle disorders such as ryanodine receptor gene (RYR1) mutation [9], D4Z4 gene mutations (which characterize facioscapulohumeral dystrophy [4, 10]) and, as described in one case, a DYSF gene mutation [3]. We present the history of a 67-year-old woman with bent spine syndrome, in whom the mutation of the DYSF gene also was the probable background of bent spine syndrome.

Case History

This 67-year-old woman had been followed up at our department for about 5 years. There were no complaints or symptoms of a similar nature in her family history. She had good muscle prowess in her childhood, but was not very active in sport before the onset of her complaints. Her symptoms started approximately 15 years ago with a slowly progressive gait disturbance. It had become difficult to climb stairs and to stay upright. Later in the course, she also noted mild muscle weakness in her shoulder girdle.

The physical neurological investigation revealed severe thoracic kyphosis in the standing position, without physical signs of spondylosis or muscle spasticity (fig. 1a). A moderate paresis was also found in the abductor muscle of the shoulder girdle. Tendon reflexes were weak but symmetric. No limb ataxia, trunk ataxia or sensory loss was present. The physical symptoms were characteristic for bent spine syndrome. Routine laboratory parameters like liver and muscle enzyme levels, including serum CK level, were within normal range.

Electromyography (EMG) of the vastus lateralis and tibialis anterior muscles did not show any signs of denervation. The interference pattern was normal, and a mixture of both high- and low-amplitude muscle action potentials was found, with an increase of polyphasic muscle action potentials. EMG of the paraspinal, infraspinalus and deltoid muscles, however, showed a typical myopathic pattern with low-amplitude, thin muscle action potentials. No denervation activity was seen. Tibialis somatosensory-evoked potentials were normal.

MRI investigation demonstrated atrophy and fatty degeneration of the erector spine muscles, with dominance on the right side (fig. 1b). Muscle biopsy of the deltoid muscle demonstrated moderate signs of myopathy with an increased number of internal nuclei, some rod body-like inclusions and lobulated muscle fibres (fig. 1c). Dysferlin immunostaining showed a moderately decreased, uneven, sarcolemmal staining in about 30% of the muscle fibers (fig. 1d). Major histocompatibility complex I immunostaining demonstrated scattered, focal sarcolemmal upregulation of this protein. Electron microscopy showed focal thickening of the basal membrane (fig. 1e), microvilli-like projections of the sarcolemma with numerous subsarcolemmal vesicles, scattered degeneration of the myofibrillar structure, mitochondrial proliferation and amorphous electrodense lipid structures (fig. 1f). Furthermore, interstitial focal accumulation of amyloid filaments was found, as demonstrated by Congo red staining (fig. 1g) and electron microscopy (fig. 1h).

The DYSF gene was analyzed by PCR and sequencing of both DNA strands of...
the entire coding region and the highly conserved exon-intron splice junctions. In addition, MLPA analysis was performed to test for deletions or duplications. Molecular genetic investigation found a c.3065G>A heterozygote mutation in exon 29 of the DYSF gene resulting in arginine-glutamine exchange (p.R1022Q). PMP22, SMN1 and D4Z4 genes were also investigated, but there were no deletion or multiplication found with multiple ligation probe analysis.

Discussion

Our patient presented with a progressive course of severe paraspinal muscle weakness and moderate shoulder girdle weakness, with the appearance of bent
It has been postulated that an active sport life could contribute to the appearance of the symptoms [11], and impaired sarcolemmal resealing of damaged myofibers could be a key pathomechanism in this disorder [12]. Our patient had normal muscle prowess before the onset of the symptoms. She did not have an active sport life; however, the symptoms first appeared when she was in her fifties.

The laboratory and enzyme investigations did not show abnormalities. EMG of the paraspinal and deltoid muscles demonstrated myopathy, and the MRI investigation showed severe fatty degeneration of the erector spinae muscles. Muscle biopsy showed myopathy with lobulated fibers and slightly abnormal dysferlin staining. Unfortunately, the quantitative dysferlin evaluation by Western blot analysis [13] was not successful. The molecular genetic investigation found a heterozygous DYSF gene mutation, which has been shown to be responsible for muscular symptom [14]. We could not confirm that the patient was suffering from autosomal recessive limb-girdle muscular dystrophy type 2B due to mutation in the DYSF gene, because we found no second mutation. It is possible that the patient was a symptomatic DYSF gene mutation carrier; a few cases have been described [15].

We completed our investigations with electron microscopy. Here we found quite similar pathological changes to what has been described and which seem to be characteristic for dysferlinopathy, such as focal myofibrillar disorganisation, mitochondrial proliferation, microvilli-like sarcolemmal projections and proliferation of the basal membrane [16, 17]. We also found amyloid accumulation, a recently recognized pathological finding in dysferlinopathy which might have a therapeutic impact [18].

Mutations in the gene encoding for dysferlin lead to distinct phenotypes, mainly limb-girdle muscular dystrophy type 2B or Myoshi myopathy. To our knowledge, only one previous case history of bent spine syndrome has been published where a DYSF gene mutation could be accounted for this rarely seen symptom. Our patient is the second case where the probable genetic background of a muscle dystrophy presenting as bent spine syndrome is a DYSF gene mutation.

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**References**