Congenital neuromuscular disease with type I fibre hypotrophy, ophthalmoplegia and myofibril degeneration

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SUMMARY We report a 7-year-old boy with progressive, early onset somatic and cranial muscle weakness associated with external ophthalmoplegia, facial weakness, type I fibre hypotrophy and myofibril degeneration. We separate this condition from congenital fibre type disproportion because of the facial weakness, ophthalmoplegia, central nucleation, and lysis in type I fibres. The case, which is similar to that described by Bender and Bender (1977), nosologically should be classified between the centronuclear myopathies and congenital fibre type disproportion, and most likely represents a congenital or neonatal disturbance of trophic interaction between nerve and muscle.

Selective smallness of Type I fibres has been noticed in various stationary or slowly progressive congenital neuromuscular disorders. Although cranial and somatic muscles are commonly involved, eye muscle weakness, or ophthalmoplegia is unusual except in myotubular myopathy1-11 and congenital myasthenia gravis. Bender and Bender14 described an infant with severe congenital somatic and cranial muscle weakness associated with complete external ophthalmoplegia, type I fibre hypotrophy and intact neuromuscular junctions. In this report we present a second case of this rare type of congenital neuromuscular disease and compare its clinical characteristics with the other similar conditions. We distinguish this congenital neuromyopathy from myotubular (centronuclear) neuromyopathy and congenital fibre type disproportion on clinical and histological grounds. Our 7-year-old male patient presented with generalised weakness associated with blepharoptosis, ophthalmoplegia, type I fibre hypotrophy and focal zones of myofibrillar lysis.

Case report

The patient, a 7½-year-old boy, was delivered by Caesarian

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Received 23 November 1981
Accepted 30 January 1982

section due to foetal distress. During the pregnancy, foetal movement was normal. He had asphyxia at birth and required intensive care for one month because of poor sucking and hypotonia. Body weight gain was slow and weak sucking remained throughout infancy. During infancy, a neuromuscular problem was suspected. He walked without support at 14 months but was clumsy and fell frequently. Recurrent infection, including upper respiratory and gastrointestinal infections, required hospitalisation for pneumonia and gastroenteritis. After entering school he revealed an inability to keep up with his peers, frequently falling with difficulty arising and in climbing stairs. Blepharoptosis became apparent in the past year and this problem worsened with fatigue. There was no positive family history of any type of neuromuscular disorder, but two male siblings died in infancy of unknown causes.

The patient was thin with small muscle bulk. He had mild bilateral blepharoptosis, complete ophthalmoplegia and mild lordosis. He showed a waddling gait, was able to climb stairs only with great difficulty and evidenced symmetrical proximal weakness, especially in the legs. Deep tendon reflexes were absent but there were no bulbar signs. Tensilon test was negative. The following laboratory studies were normal; blood count, creatine kinase, transaminase, serum electrolytes, lactate, pyruvate, sedimentation rate, EEG and CT scan of the head. Routine needle electromyography revealed low amplitude potentials but no myotonic discharges. Nerve conduction studies were normal for age.

Materials and methods

A left deltoid muscle biopsy was prepared for routine histochemistry and electron microscopy.13 14 Electron micros-
copy was performed on $10 \times 2$ mm cylinders of longitudinally oriented muscle elongated on Whatman filter paper and fixed in 0.1 M sodium cacodylate buffered 2.5% glutaraldehyde. Approximately 1 mm cylinders were cut with razor blades and secondarily fixed in 1% osmium tetroxide, dehydrated with graded methanol-propylene oxide and embedded in Epon 812 resin. One micron sections were stained with toluidine blue and subsequent thin sections stained with lead citrate-uranyl acetate.

Results

LIGHT MICROSCOPY
There was minor distortion of the fascicular architecture with moderate variation in fibre size. Approximately 12% of fibres had internal sarcolemmal nuclei (fig 1). No increase in connective tissue or evidence of fibre necrosis, regeneration or inflammation was seen. Morphohistometric analysis of the fibre diameters performed on routine myofibrillar ATPase stained sections (fig 2) showed type I fibre predominance (type I, 86.7%; type II, 13.3%) and small type I fibres (figs 3, 4). In approximately 5% of type I fibres, myofibrillar ATPase and dehydrogenase reactions revealed non-reactive central regions consistent with myofibril degeneration. These areas were single, often eccentric, and occasionally replaced two-thirds or more of the transverse fibre diameter, but were not revealed in PAS, Gomori

Fig 1 Cryostat section revealing fibre size variation and occasional central nucleation. Note absence of endomysial fibroplasia. H&E. × 280

Fig 2 Frequency histogram of minimum fibre diameters. Analysis performed on sections reacted with myofibrillar ATPase, pH 9.4.

Fig 3 Note preferential type I fibre size variation and hypotrophy. Myofibrillar ATPase, preincubated pH 9.4. × 120
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trichrome, H and E or phosphorylase reactions. The incidence (fig 5) of the lesion was similar in myofibrillar ATPase and dehydrogenase reactions (table 1). Minor disruption of the sarcotubular network was noted but no other specific architectural changes, whorling or ring fibre change was identified. The typical histochemical changes associated with myotubular myopathy were absent. No abnormality of acid or alkaline phosphatase reaction was seen.

ELECTRON MICROSCOPY
Of 27 fibres examined, seven revealed areas of myofibrillar disorganisation. The transition zone between normal myofibrils and areas of disorganisation was irregular and indistinct. Often finger-like processes of myofilament lysis accompanied by irregular condensations of osmiophilic Z band

Table 1 Characteristics and incidence of myofibrillar degeneration

<table>
<thead>
<tr>
<th>Total fibres</th>
<th>Myofibrillar degeneration</th>
<th>Incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAS</td>
<td>118</td>
<td>0</td>
</tr>
<tr>
<td>Myofibrillar ATPase</td>
<td>682</td>
<td>28</td>
</tr>
<tr>
<td>NADH tetrazolium reductase</td>
<td>627</td>
<td>32</td>
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<tr>
<td>Phosphorylase</td>
<td>310</td>
<td>2</td>
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Fig 4 Reversal myofibrillar ATPase after preincubation, pH 4.3. Note type I fibre hypotrophy and presence of type IIC fibres. × 120

Fig 5 Central and eccentric zones of myofibrillar degeneration in type I fibres. Myofibrillar ATPase after preincubation, pH 4.3. × 280

Fig 6 Transverse section of myofibre showing central myofibrillar disorganisation. The sarcolemmal membrane is intact. Irregular dense osmiophilic material arises at the sarcolemmal membrane and merges with central zones of lysis. Intact hexagonal myofilament masses are identified. Lower portion of picture reveals the region of myofibrillar lysis. × 5500
material extended to the sarcolemmal membrane (fig 6). The central areas of myofibril disorganisation contained small tenuous osmiophilic zones of presumed disrupted Z band in which the terminal actin filaments had lost their complex interdigitations. In these zones, there was considerable preservation of the paracrystallloid architecture of thick and thin filaments (fig 7). Only rare mitochondria were noted but glycogen granules were easily identified. Sarcoplasmic reticulum triads were spared and commonly found in the disorganised myofibril zones.

Discussion

Selective smallness of type I muscle fibres has been described in several neuromuscular disorders, and includes myotonic dystrophy, nemaline myopathy, centronuclear myopathy, congenital fibre-type disproportion, rheumatoid arthritis, cerebellar degenerations, and other unusual congenital or early onset neuromuscular disorders. Our patient showed no fasciculations, bulbar signs, evidence of myotonia, cerebellar signs, and no joint pain or swelling. There was no elevation of serum CK, EMG alterations, and no inclusions within muscle fibres detected by electron or light microscopy. The unusual combination of early onset somatic muscle weakness and ophthalmoplegia suggests a differential diagnosis that includes centronuclear myopathy, congenital fibre-type disproportion and the Benders’ report of congenital myopathy with type I muscle fibre hypotrophy and intact neuromuscular junctions.

Our case is most similar to that of Bender and Bender.18 Brooke14 delineated the entity of congenital fibre type disproportion and reported 14 cases whose biopsies were characterised by disproportion in size of fibres with small type I and large type II fibres. No eye muscle involvement was seen in these cases. In 26 cases of centronuclear myopathy1–7 14–18 and 23 cases of congenital fibre type disproportion15 –24 analysed from the literature, we compared the clinical features of centronuclear myopathy with congenital fibre type disproportion (table 2). The histochemical findings in myotubular myopathy are distinctive. In particular, there is central accentuation or loss in PAS, myophosphorylase and dehydrogenase reaction, consistent perinuclear absence of myofibrillar ATPase and an abnormal radial orientation of the sarcoplasmic reticulum in PAS and NADH tetrazolium reductase. These findings were not present in the central nucleated fibre of this case. While we have made no attempt to separate cases of myotubular myopathy with or without type I fibre smallness, we are convinced that some overlap will exist. In particular, the case studies of Inokuchi,25

<table>
<thead>
<tr>
<th>Delayed motor development</th>
<th>Centronuclear myopathy</th>
<th>This case Bender and Bender</th>
<th>Congenital fibre type disproportion</th>
</tr>
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<tbody>
<tr>
<td>50%</td>
<td>yes</td>
<td>&gt;90%</td>
<td></td>
</tr>
<tr>
<td>Ophthalmoplegia</td>
<td>&gt;50%</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Facial weakness</td>
<td>70%</td>
<td>yes</td>
<td>&lt; 5%</td>
</tr>
<tr>
<td>Skeletal dysfunction</td>
<td>&gt;50%</td>
<td>yes</td>
<td>&lt; 20%</td>
</tr>
<tr>
<td>Kyphoscoliosis</td>
<td>&lt;10%</td>
<td>no</td>
<td>50%</td>
</tr>
<tr>
<td>Hip dislocation</td>
<td>&gt;85%</td>
<td>yes</td>
<td>&gt; 5%</td>
</tr>
<tr>
<td>DTR-absent</td>
<td>50%</td>
<td>no/yes</td>
<td></td>
</tr>
<tr>
<td>Myopathic EMG</td>
<td>12%/occasional</td>
<td>25%</td>
<td></td>
</tr>
<tr>
<td>% of fibres with central nuclei</td>
<td>≥50%*</td>
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*Except for a single case which has 35% fibres with central nuclei.*

Fig 7 Detail of myofibrillar lysis with disrupted but identified masses of thick and thin filaments, preserved, slightly dilated sarcotubular triads with mitochondria, granular glycogen and moderately electron dense ill-defined masses of presumed Z band material. × 23 000
Karpati,19 Brooke,15 Williamson,36 and Engel9 report variants of centronuclear myopathy, while the studies of Prince et al28 and Seay et al32 may be classified intermediate variants of congenital fibre type disproportion. Our case and that of Bender and Bender12 falls into this overlap or intermediate group with the unique feature of ophthalmoplegia and may represent a maturation defect occurring later than at the time of myotube formation. Although morphological alterations in muscle fibres are rare in centronuclear myopathy30 and congenital fibre type disproportion,13,22 they have some similarities to the changes in core/multi-core disease,31,32 target formations following denervation, tenotomy,28 malignant hyperpyrexia44 and fructose 1,6-diphosphatase deficiency.14 An early focal decrease of mitochondria, myofibrillar degeneration with disintegration of Z line and considerable decrease in glycogen, gives rise to the characteristic histochemical and electron microscopic findings of typical unstructured cores.33 The lesions in our case are more sporadic, involve approximately 5% of fibres, are only found in type I fibres in contrast to multi-core disease, are not revealed in PAS or phosphorylase reactions in contrast to central core and core-targetoid change after denervation and ultrastructurally reveal greater preservation of A band material and sarcoplasmic triad organisation. The regions of myofibrillar degeneration are central in contrast to the peripherally situated zones found in familial myopathy with myofibrillar lysis of type I fibres.36 While we cannot determine whether the myofibrillar disorganisation is a primary or secondary degeneration, the change is limited to type I fibres, suggesting a strong relationship to the pathogenesis of fibre hypotrophy. Whether the patient's condition is derived from neurogenic or myopathic disease is unclear, but the lack of fibre necrosis, regeneration, phagocytosis, or endomysial connective tissue proliferation with normal CK levels excludes a "myopathic" change in the ordinary sense. Engel et al9 presented a necropsy case of type I fibre hypotrophy with central nuclei, with no abnormality in the central or peripheral nervous system. However, a large proportion of patients with centronuclear myopathy or congenital fibre type disproportion had absent deep tendon reflexes even though they probably had enough muscle power to elicit joint movement. The pathogenesis of type I fibre smallness might be explained by some defect of the monosynaptic reflex arc which would probably influence type I fibre maturation.36

The combination of type I fibre hypotrophy, predominance, sporadic myofibrillar degeneration, and ophthalmoplegia are the cardinal features in our patient and closely approximate the case of Bender and Bender.12 The prognosis should be guarded. The Benders' case was more severely involved and died during early infancy (personal communication).

This work was supported in part by USPHS HD 05615, HD 04612, and BRSG RR 05756.

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