Late components of motor unit potentials in central core disease

J. M. LOPEZ-TERRADAS AND M. CONDE LOPEZ

From the Ciudad Sanitaria de la Seguridad Social “La Paz”, Department of Pediatrics, Service of Neurology, Madrid, Spain

SUMMARY Electromyographic studies in five patients suffering from central core disease are presented. A variable amount of late components of motor unit potentials were found in all of them, as others have found in Duchenne muscular dystrophy. This suggests the existence of collateral innervation of the resultant fragments of the muscular fibre splitting present in this disorder.

Since the original description in 1956 by Shy and Magee of central core disease as a nonprogressive myopathy, several ideas have been suggested on the nature of the disease. The histochemical similarity between target fibres and central cores led Engel (1967) to postulate a neurogenic factor in the genesis of central core disease. The finding of an increased terminal innervation ratio in this myopathy could also be considered to indicate a neural aetiology (Telerman-Toppet et al., 1973; Isaacs et al., 1975).

Longitudinal splitting of muscle fibres in this disease, mainly in type 2 fibres (Telerman-Toppet et al., 1973) with slow replacement with type 1 suggests that the disease is not progressive.

In this report we describe late components of motor unit potentials in central core disease and attempts to correlate them with the morphological changes.

Methods

Motor unit potentials were recorded from the biceps brachialis and extensor digitorum communis muscles of five patients from two families suffering from central core disease, with an age range of 13 to 43 years. Three of these patients have been reported previously (Pascual Castroviejo et al., 1974). We used a three channel DISA electromyograph and concentric needle electrodes for the recording, according to the usual technique.

Twenty motor unit potentials were recorded from each muscle.

Special emphasis was applied to the recording and analysis of the morphology of the motor unit potentials, defined as those which maintain all the components invariable in 10 or more consecutive discharges. A late component of a motor unit potential was considered to be one which was separated by at least 10 ms from the start of the potential. The number of motor unit potentials presenting one or more late components was expressed as a percentage of the total number of potentials obtained for each muscle and case (see Table).

<table>
<thead>
<tr>
<th>Case</th>
<th>Age (yr)</th>
<th>Sex</th>
<th>LCBB (%)</th>
<th>LCEDC (%)</th>
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<tr>
<td>1</td>
<td>14</td>
<td>M</td>
<td>16</td>
<td>12</td>
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<tr>
<td>2</td>
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<tr>
<td>5</td>
<td>43</td>
<td>M</td>
<td>21</td>
<td>14</td>
</tr>
</tbody>
</table>

M = male; F = female; LCBB = late components, biceps brachialis muscle; LCEDC = late components, extensor digitorum communis muscle.

Case reports

FAMILY 1

Case 1 A 14 year old boy (MMR) started walking at the age of 14 months. He runs or goes upstairs with some difficulty, but without any help. Clinical
examination disclosed mild bilateral paresis and atrophy of the deltoid muscle. Reflexes were diminished and the Gowers’ manoeuvre was positive. There was bilateral pes cavus.

Case 2 A 17 year old girl (EMR) started walking at the age of 13 months. She manages to walk correctly, but she cannot jump or run. Clinical examination showed muscles with marked fatty infiltration. Mild paresis was mainly proximal. Standing up from the lying position was difficult. Gowers’ manoeuvre was positive, Tendon reflexes were diminished. There was bilateral pes cavus.

Case 3 A 24 year old woman (JMR) started walking at the age of 19 months with clumsy gait. She now walks well, but needs some help to go upstairs. Clinical examination revealed mild paresis predominantly in the proximal muscles, and difficulty of dorsiflexion of both feet, Gowers’ manoeuvre was positive, and the reflexes were markedly diminished. There was bilateral pes cavus.

FAMILY 2

Case 4 A 13 year old boy (MCM) started walking at the age of 18 months. He walks without any problem, but has some difficulty in running or climbing stairs. He does not need any help. On examination he had mild generalised paresis with diminished tendon jerks and kyphoscoliosis. He wears an orthopaedic corset.

Case 5 A 43 year old man (MCL) was probably late to begin walking. He eventually managed to walk correctly, but since he was very young he has felt weakness soon after exercise. Examination showed mild paresis, mainly proximal, with diminished reflexes, no muscular atrophies, and no difficulty in standing up from the floor.

Results

HISTOLOGICAL FINDINGS

The diagnosis was established by histological study. Biopsies of the vastus lateralis muscle were performed in all patients. All samples showed a variation in fibre size. Fibrous and fatty tissues were present in increased amounts. Histochemical study revealed a predominance of type 1 fibres, being almost entirely type 1 in case 5 (90%). The type 1 fibres showed evidence of central areas devoid of oxidative enzymes in a variable percentage from case to case (varying between 15 and 80%). Muscle fibre necrosis was sparse.

ELECTROMYOGRAPHY

Electromyographic investigation disclosed late components appearing more than 10 ms after the
terminal sprouts which, growing from preserved axons, would reinnervate the previously denervated muscular fibres.

Nevertheless, the presence of motor unit potentials with late components is not exclusive to neurogenic disorders. They may be present also in muscular dystrophies, mainly in the Duchenne type (Desmedt and Borenstein, 1973, 1975, 1976; Stalberg et al., 1974).

This feature can be interpreted in several ways. On the one hand small groups of fibres in regeneration may receive collateral innervation from neighbouring axonal shoots (Reznik and Engel, 1970). On the other hand, a viable fragment of a muscular fibre may become separated from its primitive innervation because of focal necrosis (Denny-Brown, 1960), with later reinnervation (Miledi, 1962). Lastly, longitudinal splitting of the muscle fibres may lead to the formation of viable fragments entirely separated from the original fibre (Dubowitz and Brooke, 1973; Swash and Schwartz, 1977), and then receive innervation from the neighbouring axons, occasionally changing their histochemical type as a response to their new innervation. (Aloisi et al., 1974).

Whatever the mechanism, an increase of the terminal innervation ratio should be expected. However Coërs and Telerman-Toppet (1977) have found a normal terminal innervation ratio in Duchenne type muscular dystrophy. This increases the probability that the reinnervation in this disorder takes place by growth of the nerve endings which become isolated by the necrosis of the muscular fibres innervated by them, and not by formation of collateral sprouts.

In our cases of central core disease, the presence of polyphasic motor unit potentials, with a high percentage of late components suggests the possibility of the existence of collateral reinnervation in this disorder, as has been suggested by other authors who found an increase of the terminal innervation ratio (Telerman-Toppet et al., 1973; Isaacs et al., 1975; Coërs et al., 1976).

As muscle fibre necrosis is scanty in central core disease, the free terminal axons are also very limited in number. The collateral reinnervation would, therefore, be provided by the formation of new terminal sprouts.

Muscle fibre splitting has also been described in central core disease (Telerman-Toppet et al., 1973). This finding, with the increased terminal innervation ratio, strongly suggests the existence of collateral reinnervation of the resulting fragments of the splitting and satisfactorily explains the finding of late components in the EMG.

Discussion

In muscular disorders of neurogenic origin, polyphasic motor unit potentials with a great number of late components are frequently observed (Borenstein and Desmedt, 1973; Stalberg et al., 1975). This can be related to the formation of
References


