Familial cramp due to potassium-aggravated myotonia

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Abstract
Clinical, electrophysiological, and molecular genetic features were investigated in two patients from a family with dominantly inherited myotonic disease, characterised by painful cramps, stiffness without weakness, fluctuation of symptoms, and cold sensitivity. A reduction in amplitude of the compound muscle action potential was demonstrated on cooling and administration of potassium, although no clinical exacerbation was seen. A heterozygote mutation Val1589Met was identified in the \( \alpha \)-subunit of the skeletal muscle sodium channel gene in both patients, consistent with the diagnosis of potassium-aggravated myotonia. The phenotype in this family is much milder than that previously described in another family with a mutation at this site. (J Neurol Neurosurg Psychiatry 1998;65:569–572)

Keywords: cramp; sodium channel disease; potassium-aggravated myotonia

Myotonia is a stiffness of the muscles, with inability to relax after contraction, induced by mechanical or electrical excitation. It is due to hyperexcitability of the muscle fibres, resulting from changes in ion flux homeostasis at the muscle sarcolemma, and is a manifestation of various primary muscle diseases including myotonic dystrophy, hyperkalaemic periodic paralysis, paramyotonia congenita, and potassium-aggravated myotonia. Myotonia may be described in several ways by patients, including a complaint of recurrent "cramps".

Molecular diagnosis has allowed the pathogenetic classification of these phenotypes, in particular the trinucleotide repeat expansion in the DMPK (myotonin-protein kinase) gene in myotonic dystrophy, the muscle chloride channel mutations of myotonia congenita, and the muscle sodium channel mutations of hyperkalaemic periodic paralysis, paramyotonia congenita, and potassium-aggravated myotonia. Potassium-aggravated myotonia is the most recently defined entity, having initially been termed sodium channel myotonia, and includes the conditions acetazolamide responsive myotonia, myotonia fluctuans, and myotonia permanens, all associated with mutations in the \( \alpha \)-subunit of the sodium channel gene (SCN4A). We describe a family with a much milder phenotype of potassium-aggravated myotonia than previously reported for a mutation at this site.

Case report
A 20 year old university student (IV.11, fig 1) presented with difficulty in prolonged writing, especially during examinations, and symptoms of sensory disturbance in the forearm, characterised by painful pins and needles. These symptoms had first developed several months earlier. She had experienced cramps in the toes, fingers, and eyelids, especially when tired or cold, throughout her life. Physical examination disclosed very mild orbicularis oculi myotonia, and was normal except for weakness, or wasting of the muscles.

Her 54 year old mother (III.11) had a long history of cramp-like sensations in the muscles...
of the legs (especially calf and toes) and noted difficulty with opening her eyes after tight closure. She also noticed a sensation of muscle spasm in the eyelids in cold weather and when she was tired. She had noticed difficulty in alternating between extremes of gaze. The symptoms could be more noticeable when exerting strenuously or in the cold, and when swimming. She had never experienced episodes of weakness or paralysis. She had experienced symptoms since birth, apparently being born with one arm in a fixed flexion posture. The symptoms had eased a little with age. On examination she had more marked orbicularis oculi myotonia than her daughter, and percussion myotonia of the thenar eminence, with grip myotonia. The grip myotonia increased in the first two to three movements and then eased, and the orbicularis oculi myotonia also increased initially and then eased on repetitive action. There was no weakness or wasting of the limbs. General investigation including serum sodium, potassium, and ECG were normal. Serum creatine kinase was mildly raised at 402 U/l (normal 0–150).

The mother had accepted her symptoms as normal as other members in four generations of her family (of white origin) had also experienced similar symptoms, which had not previously caused appreciable disability (fig 1). Family member III.14 experienced similar symptoms in the hands and feet, with difficulty opening her hands in the cold. In cold weather she experienced transient diplopia on rapid eye movement. She also experienced cramps "in the diaphragm". Other relatives with similar symptoms were I.1, II.6, II.8, II.10, II.12, and II.17. None had sought medical advice for their symptoms.

Methods

**ELECTROPHYSIOLOGICAL INVESTIGATION**

Nerve conduction studies and EMG were performed in conventional manner. A short exercise test was performed by the proband (IV.11) and her mother (III.11). This involved recording the abductor digiti minimi compound muscle action potential (CMAP) after stimulation of the ulnar nerve at the wrist. Disc surface electrodes (5 mm) were taped to the skin over the abductor digiti minimi (the active electrode over the hypothenar eminence and the reference over the proximal part of the fifth finger), and the ulnar nerve was stimulated with surface electrodes applied at the wrist. The electrodes were firmly secured in position and the finger, arm, and hand were immobilised using an arm board. Supramaximal stimulation of the ulnar nerve was performed (ambient temperature 24°C, skin temperature 32°C) with the peak to peak CMAP amplitude recorded at 30 second intervals for 2 minutes to establish a baseline. The patient then contracted ADM isometrically for 20 seconds. The CMAP amplitude was recorded every 10 seconds after exercise until no further decrease in amplitude was seen. The percentage decrement of CMAP amplitude after exercise was calculated when present. The limb was then cooled by immersion, with electrodes and splint in situ, within a polythene bag in a bucket of iced water. The skin surface temperature was reduced to 16°C, and the limb removed and allowed to warm to 25°C before repeating the exercise test. The tests were repeated in a normal volunteer with no evidence of neuromuscular disease.

**MOLECULAR GENETICS**

DNA was extracted from whole blood in EDTA using standard methods. Exons 1–24 of SCN4A, the gene encoding the α-subunit of the sodium channel protein, were amplified by polymerase chain reaction (PCR), and the products examined for mutations by single stranded conformation polymorphism analysis (SSCP), and by direct PCR sequencing, as previously described. In figure 2, the primers used were 5’ CCT CCT CCT CCT GGT CAT and 5’ GGG CTC GCT GCT GCT.
Results of the short exercise test performed at room temperature and after cooling of the arm to 16°C, in the proband (IV.11), her mother (III.11), and a normal unaffected person, as described in the text

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<th>Proband</th>
<th>Mother</th>
<th>Normal person</th>
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<tr>
<td><strong>Room temp</strong></td>
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<td><strong>Cooled</strong></td>
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<td>Minimum CMAP amplitude before exercise (mV)</td>
<td>6.1</td>
<td>6.0</td>
<td>6.3</td>
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<tr>
<td>Minimum CMAP amplitude after exercise (mV)</td>
<td>5.8</td>
<td>4.8</td>
<td>5.4</td>
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<td>Decrement of CMAP after exercise (%)</td>
<td>15.8</td>
<td>16.3</td>
<td>17.7</td>
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<tr>
<td>Maximum CMAP amplitude after exercise (mV)</td>
<td>15.3</td>
<td>11.7</td>
<td>15.8</td>
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MOLECULAR GENETICS

The mutation of guanine to adenine at base 4765, with substitution of methionine for valine at codon 1589, was identified in exon 24 of SCN4A (fig 2), in both III.11 and IV.11. No other mutation of SCN4A was identified.

Discussion

Mutations in genes encoding ion channels have been identified as the cause of a wide range of diseases including myotonias and periodic paralyses, episodic ataxias, seizures, long QT syndrome (in the heart), hypertension (Liddle syndrome), cystic fibrosis, startle disease, and nephrogenic diabetes insipidus. The human muscle sodium channel is a voltage dependent channel, composed of a major α-subunit and a β-subunit. The α-subunit gene (SCN4A) is on chromosome 17q23.1–25.3 and comprises 24 exons. It encodes 1836 amino acids, forming a 260 kDa glycoprotein. The α-subunit consists of four homologous domains (DI-IV) each containing six transmembrane segments (S1–6). Around 20 different mutations of the human muscle sodium channel have now been identified, and the resulting phenotypes (the sodium channel diseases) can be classified as four groups: hyperkalaemic periodic paralysis, normokalaemic periodic paralysis, paramyotonia congenita, and potassium-aggravated myotonia. These diseases show autosomal dominant inheritance, with variable penetrance, and de novo mutations may occur. Hyperkalaemic and normokalaemic periodic paralysis are associated with episodes of paralysis and in some cases may be associated with clinical or electrophysiological myotonia, especially at the beginning of an attack. Paramyotonia congenita manifests with paradoxical myotonia (increasing with exercise, as opposed to the normal “warm up” improvement with exercise in classic myotonia), which is exacerbated by cold, and shows predilection for face, neck, and long muscles of the hands, together with weakness after protracted exercise and exposure to cold. Some patients may also have episodes of muscle weakness with hyperkalaemia. Potassium-aggravated myotonia may be confused clinically with myotonia congenita, and potassium-aggravated myotonia (stiffness rather than weakness) may reflect the extent of sodium channel inactivation; a slight sustained depolarisation causing membrane hyperexcitability with resultant muscle stiff-
ness, and a larger depolarisation causing membrane inexcitability with weakness or paralysis. The short exercise test determines the excitability of the muscle membrane, and shows the sensitivity to cold and potassium even when there is no evident weakness. In the patient we tested with potassium load the response was mild, with no clinically apparent change, although altered excitability was detected on the short exercise test. A patient from the family reported by Heine et al with the same mutation (Val1589Met), was given the same dose of potassium (80 mmol) and “the muscles became very stiff within 30 minutes ... and the patient was unable to rise or walk”. The Val1589Met mutant sodium channel identified in our patients has been expressed and studied in human embryonic kidney (HEK293) cells, in which an instability of the inactivated state was shown. In view of the insignificant clinical response in our patient to oral potassium load, it is interesting that in these studies of the mutation in HEK293 cells, increasing the extracellular potassium concentration did not affect the current recorded.

These patients also exhibit variability of symptoms from day to day, which has previously been described as myotonia fluctuans. An unusual feature is the occurrence of painful cramps, as was the presenting feature in our proband. Indeed, the first linkage of potassium-aggravated myotonia (or sodium channel myotonia) to the sodium channel locus was performed in patients originally thought to have an autosomal dominant form of myotonia with painless myotonia and acetazolamide responsiveness. They also had prominent orbicularis oculi myotonia which was present in our patients. Symptomatic treatment of potassium-aggravated myotonia includes acetazolamide, and membrane stabilisers such as mexilitine and tocainide, but these were not tolerated or needed by our patients, who had mild symptoms.