Hereditary myopathy with lactic acidosis, succinate dehydrogenase and aconitase deficiency in northern Sweden: a genealogical study

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Abstract

A hereditary myopathy with lactic acidosis during physical exercise, low physical work capacity, and paroxysmal myoglobinuria (HML), called “Myopathy with deficiency of succinate dehydrogenase and aconitase” (McKusick 255125) has been described in 19 members of nine families who lived in two geographically separate areas in northern Sweden. By using the unique Swedish historical archives, including Catechetical Meeting Records from a number of northern Swedish parishes, it has been possible to trace ancestors of the nine families including all known 19 cases back in time to some key couples by making use of the unique Swedish historical archives, including Catechetical Meeting Records from a number of north Swedish parishes, and to strengthen the diagnosis in sporadic cases in some of the families. If they were found within the pedigree frame the sporadic cases were linked to familial cases.

Patients

Out of 19 diagnosed patients 16 are still alive. The 19 patients belong to nine separate families, A to I, with healthy parents and 26 healthy siblings. Of these families, A to E were described during the 1960s. Family A includes four patients out of eight siblings, family B two out of seven siblings, family C three out of three siblings, family D four out of eight siblings, and family E one out of seven siblings. Family F includes one patient out of two siblings, family G two out of six siblings, family H one of two siblings (unpublished results), and finally one patient of two siblings in family I, who was not included in the recent surveys of the disease, but was accepted as a patient with the disease because of a typical clinical history, hospital records and pedigree analysis. He died in hospital at the age of 14 years (table).

The case report of this boy, extracted from medical records, is presented below. A boy, born in 1972, had muscular weakness since early childhood. He was unable to take full part in physical exercise at school. Exercise tests on a bicycle ergometer at the age of 9 to 10 years showed low physical work capacity and high lactate blood concentration on high work loads. The nature of his disease was not understood.

At 14 years of age he performed heavy muscular work (pulling up a heavy boat) which led to an exacerbation of his disease. He developed severe muscular pain beginning in the calves

<table>
<thead>
<tr>
<th>Family</th>
<th>Total No of sibs</th>
<th>Affected sibs</th>
<th>Reference</th>
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<tbody>
<tr>
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<tr>
<td>D</td>
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</tr>
<tr>
<td>E</td>
<td>7</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
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</tr>
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<td>H</td>
<td>2</td>
<td>1</td>
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<tr>
<td>I</td>
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<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>45</td>
<td>19</td>
<td></td>
</tr>
</tbody>
</table>

Affected and healthy sibs in families A to I
and extending to the legs, stomach, and the whole body, with paresis of the legs. After one to two weeks his condition rapidly deteriorated. He was admitted to an intensive care unit almost unconscious with heart rate 120 beats/minute, high respiratory rate, anuria, blood pH 6-99, hyperkalaemia 7-10-3 mmol/l, CK level 300-1409 (NV<3-3) μkat/l, P-creatine 124-234 (N<100) μmol/l. On the second day he died in circulatory shock with falling blood pressure in spite of intensive care treatment with artificial ventilation, oxygen, bicarbonate (Trionat), albumin, plasma, peritoneal lavage, and dialysis.

Necropsy showed tubular nephritis (shock kidney). Microscopy showed striated skeletal muscles with marked variation of fibre size, some fibres without striation (hyalinised), and with vacuolisation.

Case reports of 12 other patients have been published. The symptoms, signs, and the clinical picture have been described in previous and recent papers. A short summary is given below.

All the patients had a very low physical work capacity compared with healthy people, starting in early childhood onwards. Even after such activities as walking slightly uphill, the patients developed palpitations, dyspnoea, and muscle pains. Several of them had hypotrophy of the calves. The disease had a chronic course but sometimes an acute phase of the disease developed.

The patients had a normal maximum voluntary isometric muscle strength. In spite of that an exercise test on a bicycle ergometer showed a low exercise tolerance, one third of normal or less. Even at a low work load in absolute terms, these patients had a high heart rate, a high respiratory frequency, and high lactate and pyruvate concentrations in the blood. The patients complained of pains in the leg muscles during and after the exercise.

In the chronic stage the patients had a normal circulation at rest and normal blood lactate values. However, on exercise their circulation became extremely hyperkinetic, that is, the cardiac output was high in relation to oxygen uptake. The arteriovenous oxygen difference was low and the lactate concentration in the blood was abnormally high in relation to the exercise intensity.

The heart function of these patients was normal and at maximum work intensity they reached a cardiac output and maximum heart rate comparable with that of normal subjects, although at a much lower physical work intensity. The oxygen saturation in blood from the femoral vein of exercising legs was high compared to normal subjects. The exercising muscles just could not take up the oxygen offered by the blood. The patient's work capacity on a bicycle ergometer was limited by the low arteriovenous oxygen difference, high heart rate, and a cardiac output which was maximal at a low work load, and also by intense pain in the working muscles.

Acute exacerbation of the disease may be initiated by prolonged straining exercise, starvation, or infections. It may be dramatic with extensive muscle paralysis, tachycardia, sometimes respiratory paralysis, severe acidosis, and myoglobinuria. In the most severe acute exacerbations of the disease circulatory shock may develop. Three out of our 19 known cases died in such an acute episode, in spite of intensive care, including attempts to buffer the metabolic acidosis, artificial respiration, and in one case plasmapheresis. Two other patients have been close to death in such acute attacks.

**Methods**

The genealogical analysis involved the following. After collecting information about ancestors born in the 20th century, parish registers converted to microfiches at the Research Archives, Umeå University, were studied. Families with affected subjects were then traced back as far as possible.

The establishment of the Swedish parish registers took place at the end of the 17th century according to a Church Law Act of 1686. The principal task was to keep cathechetical meeting records in order to promote the ability to read and to understand basic religious tenets of the Protestant State.
Church. These records, as well as other systematically related records of births, marriages, and deaths, make it possible to establish family links between people living more than 200 years ago and those of the recent past. The mode of inheritance was estimated according to the “sib” method.

Simulation studies were performed using the SLINK program package under the assumption of an autosomal recessive gene with full penetrance. The number of people analysed in the simulation was 156 of whom 51 were in the category marker available. One hundred replicates were generated with a four allele marker, with allele frequencies of 0·25 each, at 5 cM of the disease locus.

Results and discussion
All nine families with HML patients could be genealogically linked to one or two of the other families involved in the study, and pointed to some genetically interesting links with some of the key families back in time. Attempts to trace the nine families back in time to a common single key family have so far not been successful (fig 1). The mode of inheritance is in accordance with autosomal recessive inheritance (confidence limits 0·22–0·54) (table).

Genealogical research on a disease with proven dominant inheritance and with high penetrance often makes it possible to confirm the appearance of the disease among ancestors one or a few generations back in time using medical journals or other records or through interviews with family members. This is practically impossible when dealing genealogically with autosomal recessive diseases like the one studied here. In order to make use of genealogical data in this connection, there are some general, but hitherto seldom considered aspects to be discussed. The possibility of finding any key family of genetic significance will be maximised by using all known families with patients with the disease for genealogical studies. Pedigrees involving only a couple of families are a weak basis for genealogical analysis. The use of reference genealogical studies from the same geographical regions make it possible to discuss validity aspects of the results found. Owing to the relative immobility of the population in northern Sweden until lately, close links among people from the same region are common in rural areas, and could explain the consistency of pedigrees.

Certain specific ancestral lines are in general more reliable than others. For instance those involving close links between families studied are more reliable than those involving males, because of the effects of illegitimate children or discrepancies between de facto and de jure fathers.

The present genealogical analysis includes all known Swedish families with patients with HML. The number of links found between the separate families is not exceptional for genealogical studies in northern Sweden. However, the families studied do not come from one single geographical area. Both the separate patient families A to I and the key families O
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