Mitochondrial malic enzyme in Friedreich’s ataxia: failure to demonstrate reduced activity in cultured fibroblasts

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SUMMARY Mitochondrial and cytosolic malic enzymes were assayed radiochemically in fibroblasts from six patients suffering from Friedreich’s ataxia in order to verify earlier reports of abnormalities in these enzymes. No abnormalities could be detected in the activities of either enzyme. On cellulose acetate electrophoresis a band of enzyme activity corresponding to the mitochondrial isoenzyme was detectable contrary to earlier reports. Possible explanations for the disparity of results between different laboratories are discussed.

Friedreich’s ataxia is a progressive degenerative disorder of the heart and nervous system which follows an autosomal recessive mode of inheritance. It is associated with diabetes mellitus,1 and abnormalities in glucose2 and pyruvate3 metabolism have been reported. However, attempts at identifying specific enzyme defects have led to contradictory results.4 5 Stumpf et al8 have reported a reduction in the activity of mitochondrial malic enzyme (MEM) (EC 1.1.1.40) in cultured fibroblasts. This enzyme is present in large amounts in human heart and brain tissue and the degree of reduction in activity reported is comparable to that commonly observed in autosomal recessive enzymopathies. Bottacchi and Di Donato7 reported a less pronounced deficiency in muscle MEM and presented evidence for the existence of two iso-enzymes differing in their cofactor dependence and response to ATP and succinate. Both groups of authors also reported elevated cytosolic malic enzyme activity in Friedreich’s ataxia patients. However, Chamberlain and Lewis8 reported normal levels of MEM activity in fibroblasts from three patients. Furthermore studies on obligate heterozygotes9 have revealed levels of activity substantially lower than the 50% of normal which would be anticipated. In order to clarify these problems the activity of MEM was measured in fibroblasts from Friedreich’s ataxia patients using a new assay technique involving the release of carbon dioxide from radiolabelled malate.

Furthermore, cellulose acetate electrophoresis was carried out on sonicated cell extracts to test the report6 that a MEM band is not visible even when the membrane is overloaded with extract.

Patients

Five previously diagnosed patients with Friedreich’s ataxia were selected from the register of the Friedreich’s Ataxia Group and the records of the Department of Neurology at the Royal Hallamshire Hospital, Sheffield. All the patients were visited personally at their homes and informed consent obtained for any clinical examinations or the taking of samples. A detailed clinical examination was carried out by one of the co-authors (DK) which in each case confirmed the diagnosis of Friedreich’s ataxia. The criteria for diagnosis were cerebellar signs, absent or diminished deep tendon reflexes, muscular weakness of the lower limbs with or without distal muscular wasting, pes cavus and cardiomyopathy. The association of skeletal and cardiac manifestations was not considered essential for the diagnosis of Friedreich’s ataxia. Neurophysiological studies of motor and sensory nerve conduction velocities were not performed since all the patients met the criteria for clinical diagnosis. A skin biopsy was taken from the forearm or iliac region for cell culture. Fibroblasts from a previously reported patient10 with Friedreich’s ataxia were supplied by the MRC Clinical and Population Cytogenetics Unit in Edinburgh.

Methods

Cells were cultured initially in Ham’s F10 with 20% fetal bovine serum and then transferred on to Eagle’s Minimum Essential Medium with added non-essential amino acids and 10% fetal bovine serum. All media were buffered with HEPES and contained penicillin (200 Uml⁻¹) and streptomycin (100 μgml⁻¹). The cells were routinely screened for mycoplasma contamination by a microscopic fluorescence...
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Table 1 Clinical features of the Friedreich's ataxia patients used in this study

<table>
<thead>
<tr>
<th>Clinical features</th>
<th>Patient</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Sex</td>
<td>M</td>
</tr>
<tr>
<td>Date of birth</td>
<td>24.10.75.</td>
</tr>
<tr>
<td>Age of onset (yr)</td>
<td>6</td>
</tr>
<tr>
<td>Ataxia gait</td>
<td>Present</td>
</tr>
<tr>
<td>Incoordination (Finger-nose and Knee-heel)</td>
<td>Present</td>
</tr>
<tr>
<td>Dysarthria</td>
<td>Present</td>
</tr>
<tr>
<td>Nystagmus</td>
<td>Absent</td>
</tr>
<tr>
<td>Tendon reflexes</td>
<td>Decreased</td>
</tr>
<tr>
<td>Plantar reflexes (Extensor)</td>
<td>Increased</td>
</tr>
<tr>
<td>Romberg's test</td>
<td>Abnormal</td>
</tr>
<tr>
<td>Vibration test</td>
<td>Abnormal</td>
</tr>
<tr>
<td>Sensory loss</td>
<td>None</td>
</tr>
<tr>
<td>Muscle tone</td>
<td>Decreased</td>
</tr>
<tr>
<td>Muscle weakness</td>
<td>Absent</td>
</tr>
<tr>
<td>Upper limbs</td>
<td>Grade III</td>
</tr>
<tr>
<td>Lower limbs</td>
<td>Present</td>
</tr>
<tr>
<td>Pes cavus</td>
<td>Absent</td>
</tr>
<tr>
<td>Scoliosis</td>
<td>Absent</td>
</tr>
<tr>
<td>Cardiovascular Abnormalities</td>
<td>Absent</td>
</tr>
</tbody>
</table>

*Taken from Evans et al.10

The mitochondria were isolated by a modification of the method of Rhead and Tanaka.11 Cells from each line were grown to confluence in eight flasks of 175 cm² growing area, harvested with trypsin-EDTA and washed twice in isotonic saline. The pellet was resuspended in 2.5 ml of a cold mannitol:tris:EDTA (MTE) solution (0.27 mol.l⁻¹ mannitol, 10 mmol.l⁻¹ tris HCl, 0.1 mmol.l⁻¹ EDTA, pH 7.4) with 2.5 µg of Protease VII (Sigma Chemical Co. Ltd.) per 400 mg wet weight of cells. After incubation for 90 s on ice the cells were homogenised with 30 passes of an all glass Duarte homogeniser. The homogenate was centrifuged at 700 g for 10 minutes and the supernatant centrifuged at 15,000 g for 10 minutes. The supernatant was removed and used for the assay of cytosolic malic enzyme. The pellet was resuspended in 0.5 ml of a 0.8 mmol.l⁻¹ digitonin solution in (MTE) buffer and incubated on ice for 20 minutes. 4 ml of (MTE) buffer was added and the mixture centrifuged at 15,000 g for 10 minutes. The pellet was washed once in 4 ml (MTE) buffer, resuspended in 0.15 ml of distilled water and sonicated on ice with two 20 second bursts. The sonicate was centrifuged for 10 min at 33,000 g and the malic enzyme in all extracts assayed within 2 hours of preparation. Malic enzymes were assayed by determining the release of CO₂ from L-(U-C14) malic acid (Amersham International plc). 50 µl of extract (30-114 µg protein) were incubated with 50 µl of a pH 7.5 tris buffer containing labelled malate and all relevant cofactors (final concentrations tris 0.235 mol.l⁻¹, NADP 0.2 mmol.l⁻¹, MgCl₂ 20 mmol.l⁻¹, mercaptoethanol 2 mmol.l⁻¹, L-malate 3 mmol.l⁻¹ at 6 Ci.mol⁻¹). Incubations were carried out for 1 hour at 37°C in 5 ml serum tubes with circles of Whatman No. 1 filter paper soaked in 20 µl of 3.5 N NaOH placed in the caps. It was found that acidification was not necessary in order to achieve total trapping of the released CO₂. The papers were counted in a standard scintillation fluid. In order to correct for contamination of the mitochondrial fraction with cytosolic malic enzyme (MEC) lactate dehydrogenase activity was measured in both sets of extracts12.

Cellulose acetate electrophoresis was performed for 2 hours at +4°C using the conditions described by Stumpf et al6 and enzyme activity visualised by the method of Siebert et al.13 The cells from a 175 cm² flask were harvested, washed, frozen and thawed once in 20 ml of distilled water and sonicated on ice with two 20 second bursts. The sonicate was centrifuged at 33,000 g and 10 µl of the supernatant (71-218 µg of protein) added to the Cellogram II membranes (Shandon Southern Products Ltd.). Protein was measured by the reaction with the Folin-Ciocalteau reagent. All control cell lines were from individuals with no evidence of neurological abnormalities and were of equivalent passage number to the patients' lines.

Results

The detailed clinical features of the five cases examined by us are given in table 1. The mean age of onset of symptoms was 10-13 years and two siblings (1 and 2) were seen within 6 years of the onset of symptoms. Three patients had marked pes cavus and scoliosis was noted in two. All cases demonstrated absence of deep tendon reflexes and ataxia of both upper and lower limbs. In two cases grade II muscular weakness was noted associated with marked hypotonia in the lower limbs. Three cases showed signs of posterior column involvement whilst a positive Rhomberg's test was observed in five patients. Two cases showed nystagmus. Details of the patient from Edinburgh were obtained from the publication of Evans et al.10 This
patient, unlike the five patients we investigated, showed signs of cardiac abnormalities. Apart from patients 1 and 2 who were siblings all the other patients were genetically unrelated.

The assay of MEm and MEc gave substantial radio-active counts (600-5,000 cpm) at the specific activity employed with consistent blank values of approximately 800 cpm. The final mitochondrial sonicate contained 0-03-0-70% cytosolic contamination. This accounted for 5-40% of the apparent total MEm activity. After correction of the MEm activity for the contribution due to cytosolic MEc and protein the MEm activity was found to account for from 0-53-5-1% of the total activity. In view of the small contribution MEm contamination would make to the MEc activity a mitochondrial marker enzyme was not used to correct the latter. Table 2 shows the results of assays of MEm and MEc on cells from the patients and the controls. When values were assessed by a Student t test no significant differences could be found between the patients and controls in terms of corrected or uncorrected MEm MEc or the ratio of MEm/MEc specific activities for each individual. In terms of MEm and MEc individually all values for the patients were within the control range. With the MEm/MEc ratios only one value was slightly below the normal range.

Electrophoresis of sonicates from five of the patients including patient 6 revealed a distinct MEm band (fig). This band showed the polymorphism characteristic of human MEm and migrated in the same position as that of a reference human brain sample. The band was still present after incubation of one Friedreich’s ataxia line and a control line in stain containing 2-5 m.mol.l\(^{-1}\) ATP although it was difficult to assess whether there had been a specific reduction in intensity.

### Table 2  Malic enzyme activities in cultured fibroblasts from Friedreich’s ataxia patients

<p>| pmoles CO₂ min (^{-1}) mg (^{-1}) protein | Mean (± SD) |</p>
<table>
<thead>
<tr>
<th>MEm (Uncorrected)</th>
<th>MEm (Corrected)</th>
<th>MEc</th>
<th>MEm/MEc</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls(7)</td>
<td>186(±120)</td>
<td>173(±110)</td>
<td>645(±310)</td>
</tr>
<tr>
<td></td>
<td>(63-4-342)</td>
<td>(54-9-314)</td>
<td>(293-1226)</td>
</tr>
<tr>
<td>Patients(6)</td>
<td>202(±85)</td>
<td>182(±91)</td>
<td>671(±188)</td>
</tr>
<tr>
<td></td>
<td>(92-4-305)</td>
<td>(69-3-287)</td>
<td>(488-1011)</td>
</tr>
</tbody>
</table>

All assays were performed in duplicate.

**Discussion**

The failure to demonstrate either a reduction in MEm activity or an elevation in MEc activity is in agreement with Chamberlain and Lewis \(^8\) but at variance with Stumpf et al \(^6\) and Bottachi and Di Donato. \(^7\) There are a number of possible explanations for these disparities.

Firstly there may be differences between the authors in the criteria for defining Friedreich’s ataxia. Certainly Friedreich’s ataxia is a disorder which can easily be misdiagnosed and there is a plethora of criteria by which to define it reported in the literature. \(^14\)-\(^16\) This is why we believe it is necessary to report details of the clinical features of the cases used in studies on this disease. Our patients fitted the prime criteria of reduced tendon reflexes and ataxic gait. Together with the other abnormalities reported we are confident that this picture represents classic Friedreich’s ataxia even though one patient showed his first symptoms at 30 years of age.

Secondly it may be that there is genetic heterogeneity within Friedreich’s ataxia which is expressed on a geographical basis. Certainly inherited diseases classified on clinical grounds have often been later subdivided on the basis of investigations of the primary biochemical defects. Such a possibility is testable by the exchange of material between groups in different countries.

A third possibility is that technical artefacts have led to the disparities. The assay used by Stumpf et al follows the formation of NADPH and there are a number of other enzymes within the cell capable of reducing NADP. However, Chamberlain and Lewis \(^8\) measured this parameter in their assays and the stain used in the electrophoretic method reported here also detects NADPH formation. Thus it seems unlikely that the assay is the source of the problem. The values for the specific activity of MEm reported here are approximately 15% of those reported by Stumpf et al \(^6\) and Chamberlain and Lewis. \(^8\) This is partly a result of the lower substrate concentration used and correction.
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for this based on the published kinetic parameters of human MEm\textsuperscript{17} gives values 25% of that of Stumpf et al.\textsuperscript{8} This may indicate that the dehydrogenation assay is measuring dehydrogenases other than malic enzyme. The decarboxylation assay should be inherently very specific since malic enzyme is the only human enzyme known to decarboxylate L-malate. A further complication is the report by Bottachi and Di-Donato\textsuperscript{7} that there exists in human muscle mitochondria a malic enzyme which can utilise NAD and NADP, shows sigmoidal kinetics, is inhibited by ATP and is activated by succinate. This work follows earlier reports of the existence of such an isoenzyme in bovine adrenals\textsuperscript{18} rabbit heart,\textsuperscript{19} rat liver\textsuperscript{20} and human placenta.\textsuperscript{21} It is not clear whether this enzyme exists in human fibroblasts and if so, in what proportion it is to total MEm activity. Certainly the failure to demonstrate substantial ATP inhibition indicates that it is not the predominant isoenzyme. Bottachi and Di-Donato\textsuperscript{7} indicate that it is the non-NAD dependent isoenzyme which is deficient in Friedreich’s ataxia muscle suggesting the possibility that slight differences between culture conditions and separation procedures could lead to preferential inactivation/enrichment of one isoenzyme or the other. However the culture conditions used were very similar to those reported by Stumpf et al.\textsuperscript{6} Differences due to variations in the degrees of confluence when the cells were harvested are unlikely since our cells were harvested when at confluence whilst those used by Chamberlain and Lewis were harvested in the logarithmic growth phase (S Chamberlain—personal communication). In one experiment the isolation method reported by Stumpf et al\textsuperscript{6} was used on a control line and the assay performed with a similar substrate concentration. The values obtained were within the control range previously established. Despite these observations, the possibility exists that slight variations in culture conditions or isolation procedures may dramatically influence the MEm isoenzyme pattern.

Finally it needs to be stressed that all cell lines should be tested for mycoplasma-contamination by a reliable non-microbiological method as this can lead to spurious results in various enzymological tests.

Friedreich’s ataxia has proved to be a difficult disorder to define as an enzynopathy. This is probably partly due to problems in accurately defining clinical diagnostic criteria, cell culture conditions and assay methodology. If indeed it is caused by an inherited enzyme deficiency then it will have proved to be one of the most difficult to define of all the inborn errors of metabolism.

References

18 Sauer LA. Mitochondrial NAD—dependent malic

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