Myopathy with mitochondrial inclusion bodies: histological and metabolic studies

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SYNOPSIS Apparently new information about a metabolic abnormality has been obtained by measuring biochemical and respiratory responses to controlled exercise in a patient with abnormal mitochondria in muscle fibres. A male patient (49 years old) presented with bilateral ptosis of 15 to 20 years' duration and weakness for one to two years. Biopsies from the deltoid and triceps muscles were subjected to histological, histochemical, and electron microscopical examination. Routine histology showed only minor changes; 2–5% of muscle fibres had pale borders in which there were aggregates of mitochondria and 1–5% of fibres showed atrophy. Histochemical examination showed increased activity of succinic dehydrogenase in mitochondria and lactate dehydrogenase in cytoplasm. Electron microscopy showed crystalline inclusions in many subsarcolemmal mitochondria. Metabolites were studied during and after exercise on an ergometer and revealed remarkable differences from normal. Blood lactate rose to 12.5 μmol/ml and pyruvate to 0.39 μmol/ml compared with up to 4.0 and 0.16 μmol/ml respectively in controls exercising to a comparable percentage of capacity. Concentrations of ketone-bodies and free fatty acids fell during and after exercise, while they rose in the controls. These observations imply a major mitochondrial defect which causes dramatic biochemical changes in fuel supply in blood during exercise. The changes suggest that fatty metabolism was accelerated (in this patient) and was related to a block of carbohydrate utilization as fuel.

Electron microscopy has allowed delineation of abnormalities of intracellular structures including mitochondria. The oculoskeletal myopathies is one group of disorders in which major defects are detectable in this way. These patients present with progressive ptosis, generalized muscle weakness, and sometimes paresis of extraocular muscles. The literature has been reviewed by Morgan-Hughes and Mair (1973) who reported an additional four patients. Although these authors have supplied detailed descriptions of the neuropathological changes which may occur, they did not investigate the metabolic effects of the disorder. They showed histochemical and ultrastructural changes in the mitochondria of muscle fibres. Mitochondria are the sites within the cells where enzyme systems subserving oxidative phosphorylation of both fat and carbohydrate are concentrated, and some authors have suggested that defects in metabolism in these pathways may occur in association with such abnormalities (Bradley et al., 1969).

We have had the opportunity of investigating another patient with oculoskeletal myopathy. We have evidence that the neuropathological changes are similar to those described by Morgan-Hughes and Mair (1973) and now report metabolic responses to exercise which have apparently not been investigated previously in patients with these clinical and histological abnormalities.

CASE REPORT

A.B. (Institute of Neurological Sciences, Glasgow, 807873) was a 49-year-old married man with two children, who was employed as a lorry driver. His mother died of carcinoma of the stomach at 62 years and his father of myocardial infarction at 67 years.
TABLE 1

HISTOCHEMICAL REACTIONS OF MUSCLE FIBRES IN PATIENT*

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Site of enzyme</th>
<th>Activity in muscle fibres</th>
<th>Distribution in abnormal fibres according to type</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Cytoplasm</td>
<td>Mitochondria</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal fibres</td>
<td>Abnormal fibres</td>
</tr>
<tr>
<td>Succinic dehydrogenase</td>
<td>M</td>
<td>t</td>
<td></td>
</tr>
<tr>
<td></td>
<td>d</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cytochrome oxidase</td>
<td>M</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3-hydroxybutyrate dehydrogenase</td>
<td>M</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glutamate dehydrogenase</td>
<td>C + M</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Isocitrate dehydrogenase</td>
<td>C + m</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>α-glycerophosphate dehydrogenase</td>
<td>NAD linked</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>NAD—tetrazolium oxidoreductase</td>
<td>C + M</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>NADP—tetrazolium oxidoreductase</td>
<td>C + M</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>Malate dehydrogenase</td>
<td>C + M</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>Lactate dehydrogenase</td>
<td>C</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>Glucose-6-phosphate dehydrogenase</td>
<td>C</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>6-phosphogluconate dehydrogenase</td>
<td>C</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>Alcohol dehydrogenase</td>
<td>C + m</td>
<td>++ +</td>
<td>++ +</td>
</tr>
<tr>
<td>Myosin ATPase</td>
<td>C</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Glucosan phosphorylase</td>
<td>C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leucine aminopeptidase</td>
<td>C + M</td>
<td>N</td>
<td>N</td>
</tr>
</tbody>
</table>

* Abnormalities in the activities of mitochondrial (M) and cytoplasmic (C) enzymes were seen in some type I and Ila fibres.
(+ + = moderately increased; + + + = greatly increased; N = normal activity; m = predominantly mitochondrial; c = predominantly cytoplasmic; t = triceps; d = deltoid.)

He had five siblings, three of whom were well. One died of alcoholic cirrhosis (50 years) and the other of 'stomach trouble' (51 years). There was no family history of muscle disease, ptosis, or diabetes. He complained of drooping of his eyelids for many years, and he was aware of trying to lift them with his forehead. He had also had very occasional double vision on lateral gaze for about two to three months. These symptoms were worse later in the day. Photographs indicated that he looked normal at 18 years, but a left ptosis had become obvious at about 33 years. He made no complaint about tiredness when first seen, but later admitted that this had been a problem for one to two years; it was not related to time of day.

His height was 176 cm and weight 62 kg. There was no obvious muscular wasting. He had wrinkling of his forehead with bilateral ptosis. There was weakness of upward conjugate gaze and slight weakness of conjugate gaze to either side. Convergence was poor. His voice had a nasal quality but the palate moved normally. In the limbs there was no muscle fatigability but there was slight weakness of both triceps muscles and of the small muscles of both hands. All reflexes were normal. The remainder of the physical examination was negative.

The following investigations were normal: erythrocyte sedimentation rate (7 mm/hr), haemoglobin (14.8 g/100 ml), white cell count (7,000/mm³), electrolytes (Na⁺ 141, K⁺ 4.6, Cl⁻ 99 mEq/litre), WR, blood urea (44 mg/100 ml), blood sugar (85 mg/100 ml), protein bound iodine (5.3 μg/100 ml), electrocardiograph, electroencephalogram, radiographs of skull and chest, and eredophonium bromide (Tensilon) test. Serum creatine phosphokinase was at the upper limit of normal (101 mU/ml; normal range for laboratory—100 mU/ml). Nerve conduction studies showed normal conduction velocities in right median (68 m/s) and ulnar (54 m/s) nerves but slight reduction in right lateral popliteal nerve (38 m/s), which was thought to be significant as limb temperature was satisfactory. Electromyography of proximal muscle groups showed an excess of myopathic units (short duration polyphasic units). There was also scanty fibrillation in the right deltoid, and polyphasic fasciculations in the right tibialis anterior and right orbicular oculi muscles suggesting a concomitant neuropathic disorder.

MUSCLE BIOPSIES Biopsies were taken from the left triceps and the left deltoid muscle under local anaesthesia. They were examined by light microscopy with routine stains. Glycogen distribution was identified by periodic acid Schifff staining (Hotchkiss, 1948) and fat by staining with Oil red O (Lillie and Ashburn, 1943). Specimens were also examined by electron microscopy (Philips 201 electron microscope), goniometry (Philips 301 electron microscope with double tilt goniometry), electron diffraction and
x-ray microchemical analysis (Edax—Energy-Dispersive Analysis by x-ray) with Philips 301 electron microscope. The enzymes shown in Table 1 were demonstrated histochemically. Succinic dehydrogenase was examined by the method of Seligman and Rutenberg (1951); myosin ATPase by that of Padykula and Herman (1955a, b); glucosan phosphophorylase by that of Takeuchi and Kuriaki (1955); and leucine aminopeptidase by that of Nachlas et al. (1957). The other enzymes were studied by the methods of Hess et al. (1958). Light microscopy with phase contrast and polarized light revealed virtually normal muscle and the abnormalities seen in stained preparations were minor. In the specimen from the left triceps muscle about 5% of the fibres were atrophic, some being exceptionally small in cross-sectional area (20 μm) with clusters of nuclei. In atrophic fibres there were also some nuclei which were larger than normal (15 μm diameter) and these had very large nucleoli (5 μm diameter). Some obviously abnormal fibres (approximately 2% of total) had unusually pale borders which had abnormal mitochondrial aggregates. The nuclei and nucleoli were larger in these areas. In some areas many of the fibres had internally placed nuclei but the remaining fibres appeared normal. Longitudinal section showed that some nuclei were concentrated close to capillaries, these nuclei appearing larger than normal. The changes in the specimen from the left deltoid muscle were similar but only 1% of the fibres were atrophic. Atrophic fibres appeared in groups of up to six but no supporting evidence of denervation was found in preparations displaying motor end plates and terminal innervation. There was a larger proportion of fibres with pale borders in this specimen than in that from triceps (about 5%).

In longitudinal sections large nuclei were frequently seen close to capillaries. In both specimens glycogen was found to be reduced in all fibres. The concentration of fat in type I fibres was reduced and it was virtually absent from type II fibres: the fat which remained was mainly peripheral in the fibres.

**ELECTRON MICROSCOPY** Electron microscopy (EM) of the specimens from deltoid and triceps revealed normal mitochondria (Fig. 1). The least abnormality

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**FIG. 1.** Electron micrograph of the crystalline inclusion bodies in mitochondria showing the patterns which are determined by the angle of incidence of the electron beam. Two mitochondria are shown which do not contain crystals but have abnormally arranged cristae. × 12,500.
was the arrangement of cristae in concentric circles in the mitochondria rather than transversely as is found normally. Some mitochondria also contained crystal-like inclusions. These mitochondria were most numerous in the pale peripheral borders of the abnormal fibres seen on light microscopy. The crystals were invested by the inner mitochondrial membrane, which was not incorporated in their structure. The crystals varied in length from 0.2 to 5 \( \mu \text{m} \) and were slightly curved. They were sometimes square but usually oblong. Each crystal consisted of bands about 0.04 \( \mu \text{m} \) in width separated from adjacent bands by a clear region 0.008 \( \mu \text{m} \) across. The bands had light and dark longitudinal stripes (0.008 \( \mu \text{m} \) wide). Although the majority of crystals had such a pattern, in some the bands were arranged obliquely and others had a grid-like structure. Goniometry confirmed that some of the crystals were curved and it was possible to show that the differences in the appearances of the bands depended upon the angle of incidence of the electron beam on the crystal. Electron diffraction and microchemical x-ray analysis were non-contributory because of the low mass of the crystals. EM confirmed that there was a general reduction in the fat content of the muscle fibres.

**HISTOCHEMISTRY** Histochemical examination with myosin ATPase showed fibres of types I, II, and Ila in the usual distribution. Fibre typing was more difficult with other enzymes. Marked abnormalities were observed in fibres of types I and Ila among which 20% of fibres were affected. No abnormalities were found in type II fibres. Abnormalities were principally seen in the activities of mitochondrial or predominantly mitochondrial enzymes but there were also changes in several enzymes in cytoplasm. In the biopsy specimen from triceps there was a general reduction in the activity of succinic dehydrogenase but there was a peripheral zone of considerably increased activity in type I and type Ila fibres. In the biopsy specimen from deltoid the activity of succinic dehydrogenase was also increased peripherally in type I and type Ila fibres (Fig. 2). Several other mitochondrial enzymes showed increased activity and similar changes were observed in cytoplasmic enzymes, including lactate dehydrogenase (Table 1).

**SKIN BIOPSIES** Biopsies of skin were taken at the same time as the muscle biopsies. No histological changes were found and on EM all mitochondria were normal.

**BIOCHEMICAL RESPONSE TO EXERCISE** The biochemical changes during exercise were studied on two occasions, during and after a 30 minute period of exercise. The results were compared with those from six controls—normal male subjects aged 28 to 44 years. The investigations were performed between 09:00 hr and 11:00 hr after an overnight fast. A catheter was placed in an antecubital vein and flushed with saline. Two resting samples of blood were taken. Work was performed for 30 minutes at 600 kpm on a bicycle ergometer (Elema Schönander constant load ergometer, EM369). Heart rate was recorded during the exercise and for an additional 15 minutes, using miniature chest electrodes with an electrocardiograph. Blood samples were taken at five minute intervals during exercise and then at 15, 30, 60, and 90 minutes afterwards. The controls were investigated in a similar way: their work load (600 \( \pm \) 100 kpm) was adjusted so that the increased heart rates were similar to that in the patient—namely, between 150 and 170/min. In the second investigation pulmonary ventilation was measured.

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**FIG. 2.** Abnormal succinic dehydrogenase activity in a type I muscle fibre. The intense zones of peripheral activity were characteristic of the abnormal fibres and in these areas the intramitochondrial crystals were found. \( \times 700. \)
and expired air was collected in a Douglas bag and analysed for O₂ and CO₂ to determine oxygen consumption, carbon dioxide output, and respiratory exchange ratio (R). Measurements were made at rest and during exercise and recovery.

Each blood sample was divided into parts. Five millilitres were deproteinized by addition to 5 ml 10% perchloric acid. The remainder was heparinized and the plasma separated by centrifugation. All specimens were then stored in ice. The deproteinized samples were analysed for glucose (Bergmeyer and Bernt, 1963), lactate and pyruvate (Hohorst et al., 1959), glycerol (Kreutz, 1962), acetoacetate and 3-hydroxybutyrate (Williamson et al., 1962). The plasma sample was analysed for free fatty acids (FFA) by a colorimetric method after chloroform extraction (Itaya and Ui, 1965), for human growth hormone (HGH), by radioimmunoassay using two antibodies (Morgan and Lazarow, 1963), and for immunoreactive insulin (IRI) by a charcoal immunoassay (Hunter, 1969; Hunter and Ganguli, 1971). The standards used were MRC Standard A for HGH and Wellcome MR71 for IRI.

Glucose concentrations (Fig. 4) at rest were similar in the patient and the controls but there was a marked difference during and after exercise. The

**FIG. 3.** Blood lactate and pyruvate ($\mu$mol/ml) in the patient (●—●) and six normal control subjects (○—○) (± SEM) during and after 30 minutes of exercise, indicated by the solid bar.

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1240

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blood glucose rose to 135 mg/100 ml in the patient at the end of exercise, whereas there was no significant change in the controls.

Lactate concentrations (Fig. 3) at rest were similar but with exercise there was a rapid elevation in the patient to 12.5 μmol/ml compared with 4.0 μmol/ml in the controls. The plasma concentration in the patient declined after exercise but had not returned to the resting value 90 minutes after the end of exercise.

Pyruvate concentrations (Fig. 3) at rest were similar and the exercise caused a rapid elevation. The highest plasma concentration in the patient (0.39 μmol/ml) was achieved 15 minutes after the end of exercise. After this the concentration declined but had not returned to the resting value by the end of the investigation. The maximum lactate/pyruvate ratio was greatly elevated in the patient (400% above resting) compared with the controls (230% above resting).

Plasma free fatty acids (FFA) concentrations (Fig. 5) at rest in the patient were 80% higher than in the controls on the first occasion but similar on the second. On both occasions concentrations declined during exercise compared with the controls, in which they rose.

Blood glycerol concentration (Fig. 5) at rest in the patient was higher than in the controls on the first occasion but similar on the second. In both studies it rose to nearly twice the mean level in the controls by the end of exercise. The concentration then declined in both the patient and the controls, the resting values being achieved by the end of the investigations.

Total blood ketone-body concentrations (Fig. 6) at rest in the patient were five times higher than in the controls on the first occasion and within the normal range on the second. The concentrations fell during and after exercise whereas they rose in the controls. On the first occasion the 3-hydroxybutyrate/acetoacetate ratio for the patient was 2.24 before exercise and rose to 4.40 after five minutes of exercise and then fell progressively to 1.0, 30 minutes after exercise. In the patient's second exercise investigation the changes were similar. In the controls the mean value at rest was 2.16, and it fell to 1.36 after five minutes of
exercise. After 30 minutes of exercise it was 2.83 and 30 minutes later it had risen to 3.8.

Plasma immunoreactive insulin (IRI) concentrations (Fig. 4) at rest were similar but exercise caused a fall in plasma IRI in all studies. The fall was greater in the controls after 20 minutes of exercise. After this the concentrations rose, both in the remaining part of exercise and for 15 minutes afterwards. The concentration was much greater 15 minutes after exercise in the patient (200% of resting) than in the controls (133% of resting). The concentrations returned to resting values by the end of the investigations.

Plasma human growth hormone (HGH) concentrations (Fig. 7) at rest were similar, but with exercise there was a rapid elevation in the patient to 40 µU/ml compared with 27 in the controls. The concentrations then declined but had not returned to the resting value by the end of the investigations.

Oxygen consumption at rest was 240 ml/min. The
values at six, 20, and 30 minutes during exercise were 1·54 l/min, 1·38 l/min, and 1·45 l/min. The oxygen consumption had returned to normal 20 minutes after the end of exercise. R was 1·04 while resting before exercise, indicating hyperventilation, and fell progressively during exercise to a minimum of 0·74 at the end. The value 20 minutes afterwards was 0·85 (Table 2).

**TABLE 2**

<table>
<thead>
<tr>
<th></th>
<th>Before exercise</th>
<th>6–8 min</th>
<th>20–22 min</th>
<th>28–30 min</th>
<th>20 min after</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate</td>
<td>76</td>
<td>150</td>
<td>168</td>
<td>174</td>
<td>100</td>
</tr>
<tr>
<td>Ventilation</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATPS/min</td>
<td>11·0†</td>
<td>49·3</td>
<td>48·5</td>
<td>54·5</td>
<td>9·5</td>
</tr>
<tr>
<td>Oxygen consumption</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>STPD l/min</td>
<td>0·24</td>
<td>1·54</td>
<td>1·38</td>
<td>1·45</td>
<td>0·19</td>
</tr>
<tr>
<td>R</td>
<td>1·04†</td>
<td>0·9</td>
<td>0·74</td>
<td>0·74</td>
<td>0·85</td>
</tr>
</tbody>
</table>

* The heart rate suggests that he was at nearly 80% VO₂ maximum so oxygen consumption is relatively low.
† Hyperventilating: end-expired CO₂ 30 Torr.

**BIOCHEMICAL RESPONSE TO EXERCISE AFTER NICOTINIC ACID** The biochemical changes during exercise were also studied 15 minutes after ingestion of nicotinic acid (200 mg). The patient exercised for 30 minutes at 600 kpm and the arrangements were similar to those on the previous occasion when ventilatory studies were also carried out. Heart rate changes were the same as those in the previous investigations of exercise (maximum 170/min). The average exercise oxygen consumption was also similar (1·43 l/min). The major difference was in blood glycerol which was depressed at rest by nicotinic acid (from 0·06 to 0·03 μmol/ml) and rose to 0·05 μmol/ml only during exercise. It then rose during recovery but even then the maximum value was 0·10 μmol/ml 45 min (compare Fig. 5). There was also a marked reduction in the blood glucose concentrations observed during and after exercise (maximum 110 mg/100 ml) and maximum blood lactate and pyruvate were also lower (9·87 μmol/ml and 0·34 μmol/ml respectively). Plasma FFA declined during exercise (lowest concentration 0·48 μeq/ml). Total ketone-bodies showed some depression during exercise but rose to 0·10 μmol at 45 min afterwards. Growth hormone changes were similar (maximum 36·0 μU/ml). The gas exchanges studies showed the same high R value at rest (1·05) but reduction was less marked during and after exercise: 0·86 at the end of exercise and 0·92–0·95 in the following 30 min.

**EFFECT OF HEPARIN UPON LIPOLYSIS** The investigations were carried out in the morning after an overnight fast according to the method of Fineburg et al. (1972). Corn oil (60 g) which had been emulsified with egg albumen (15 g) was given orally. This was followed three hours later by an intravenous injection of heparin (50 mg). Venous blood samples were taken before ingestion of the corn oil, at 30 minute intervals and then at 15 minute intervals after the injection of heparin for a further two hours. The test was repeated on another occasion omitting the corn oil. The blood samples were analysed for FFA and keto-bodies (acetoacetate plus 3-hydroxybutyrate) as already described. After the ingestion of the corn oil, plasma FFA concentrations remained unchanged and just before the injection of heparin the concentration was 0·80 μEq/l. The concentration 45 minutes after the injection of heparin was 2·4 μEq/l and it then fell to a value of 1·8 μEq/l 25 minutes later. In the control investigation on the patient, injection of heparin alone also caused a rise in the concentrations of FFA, but this was less than that obtained after ingestion of corn oil. The rise in FFA concentrations after heparin was associated with a considerable rise of total ketone bodies.

**GLUCOSE TOLERANCE TEST** A standard glucose tolerance test was performed after an overnight fast. Glucose (50 g) was given orally and blood samples were obtained at 30 minute intervals for 2½ hours. Lactate, pyruvate, glucose, and plasma IRI were estimated as in the exercise investigation. The rise and the subsequent fall of both glucose and plasma IRI were similar to those of normal subjects. The tolerance to a glucose load was therefore normal in the patient.

**DISCUSSION**

Abnormal muscle mitochondria have been reported in several apparently unrelated conditions. The first report of a muscle disorder with abnormal mitochondria was made in 1959 when increased metabolism was found. The patient's mitochondria contained large crystalline inclusions (Ernster et al., 1959; Luft et al., 1962). Mitochondrial abnormalities have been reported in children (Shy et al., 1966; Jerusalem et al., 1973) but studies showed no significant increases in metabolic rates. In one of these,
'megaconial myopathy', rectangular inclusion bodies were observed in the muscle mitochondria. Mitochondrial inclusions have also been reported in the skeletal muscle of adults (Shafig et al., 1967; Bradley et al., 1969), but these reports did not include assessments of the effect of the muscle disorder upon metabolism.

The present patient had an abnormal metabolic response to exercise compared with controls. Concentrations of lactate and pyruvate rose during exercise to values above those observed in the controls. There was also a rapid rise in blood glucose concentration. These observations suggest that, although glycolysis was proceeding, there was a block to complete oxidation of carbohydrate via acetyl CoA and the tricarboxylic acid cycle.

The patient's lactate/pyruvate ratio, during the first part of exercise, rose above the ratios observed in the controls. Some elevation is the usual response and is probably due to a rapid fall in the ratio of cytoplasmic nicotinamide adenine dinucleotide (NAD) and reduced nicotinamide adenine dinucleotide (NADH) (Krebs, 1967). This cytoplasmic activity is normally linked to the concentration of NAD and NADH in mitochondria but may be independent (Devlin and Bedell, 1960). These changes in the mitochondria may be reflected in the ratio of the concentrations of 3-hydroxybutyrate/acetocetate in the blood (Williamson et al., 1967). In the patient this ratio first rose and then fell, whereas the converse was observed in the controls. Such findings are obtained with relative cellular hypoxia (Alberti et al., 1972). The changes in these ratios are therefore in keeping with the suggestion already made of a block of oxidative metabolism for carbohydrate.

There was also a much greater rise in blood glycerol concentration in the patient compared with the controls suggesting greater fat mobilization in the patient. Lipolysis results in production not only of glycerol but also FFA. The rise in the concentration of plasma FFA during exercise which occurred in the controls, and is the normal finding, was, however, absent in the patient. Tests of lipoprotein lipase activity had shown that there was no block in the normal production of FFA from plasma triglycerides suggesting that normal transport of fat was occurring. The depression of plasma FFA in the patient during exercise may therefore have been due to either greater catabolism of FFA or to re-esterification to fat. There was no evidence from the biopsies that deposition of fat was occurring excessively in the muscles studied. Normally ketone-body concentrations rise during the post-exercise period (Courtice and Douglas, 1936; Johnson et al., 1969) but this change was absent in the patient. The blockage of carbohydrate metabolism could result from increased fat oxidation to acetyl CoA, a process which is known to block further pyruvate oxidation (Garland and Randle, 1964) and cause accumulation of pyruvate and lactate. This would also account for some of the elevation of glycerol, for this is normally metabolized through the glycolytic pathway via α-glycerophosphate.

The rapid rise of blood glucose implies both decreased carbohydrate utilization during exercise and also active glucose production by the liver. Pathways subserving carbohydrate storage as glycogen were normal, however, as adduced from the normal glucose tolerance test. The blockage of carbohydrate metabolism did not prevent the patient exercising and it is probable that energy production depended upon metabolism of fat, as mobilization was occurring and yet there was no accumulation of FFA during exercise and of ketone-bodies after exercise. Additional support for the conclusion that fat was actively metabolized by the patient was the finding that the patient had a relatively low R (CO₂ production/O₂ consumption) during exercise. R often rises in severe exercise to a value greater than 1-0. For this patient the exercise was severe relative to his capacity. The value above unity before the investigation was due to hyperventilation but it would be unlikely that this would have driven off so much CO₂ before exercise as to account for the low R 30 minutes later. Further evidence for active mobilization of fat as fuel during exercise by the patient was obtained from the study with nicotinic acid. Nicotinic acid is known to block fat mobilization (Carlson and Orå, 1962). The lower values of FFA and glycerol and higher R values we obtained after nicotinic acid during exercise by the patient are in keeping with the conclusion that fat mobilization, although active in the other studies, was blocked by the nicotinic acid. This argument is also supported by the histochemical
findings of increased activities of enzymes involved in oxidative metabolism such as succinic dehydrogenase and cytochrome oxidase in the abnormal mitochondria.

There were also differences in the hormonal responses to exercise of the patient compared with the controls. Plasma IRI concentration was depressed to a lesser extent during exercise in the patient compared with the controls and he had a greater increase after exercise. During exercise there is an increase in circulating catecholamines (von Euler and Hellner, 1952), which may explain the failure of the increased glucose concentration in the patient to stimulate insulin release, as adrenaline and noradrenaline inhibit insulin release (Kris et al., 1966; Porte and Williams, 1966). The higher concentrations of plasma IRI in the patient compared with controls after exercise could, however, be related to the increased blood glucose as concentrations of blood catecholamines return rapidly to normal, the fall being of the order of 60% in five minutes (Johnson et al., 1974).

Plasma HGH concentrations were higher in the patient during exercise. It has been suggested that HGH causes lipolysis during exercise and enhances the availability of FFA (Hunter et al., 1965) and therefore the difference in FFA response to exercise in our patient compared with the controls may have been related to the differences in HGH concentration. Although studies of patients with hypopituitarytism have shown that it is not essential for FFA release (Johnson et al., 1971), it does appear to cause FFA release in some situations (Johnson et al., 1973).

There appear to be considerable differences in the metabolic effects of disorders in which abnormal mitochondria are found. The patient with myopathy described by Bradley et al. (1969) had mitochondrial changes on EM which appeared similar to those in our patients. However, in contrast, they found excessive fat in muscle fibres. The two patients described by Coleman et al. (1967) had increased activity of succinic dehydrogenase and other enzymes but no EM studies were reported. These patients, however, also had abnormal accumulation of fat in muscle fibres. Other patients with excess fat in muscle fibres have been described by Worsfold et al. (1973) and in vitro studies of mitochondria from these patients showed that oxidative metabolism was depressed. Mitochondrial aggregates have also been described by Engel (1964) in the muscle fibres from three patients, but these mitochondrial aggregates lacked succinic dehydrogenase activity, thus differing from the abnormal mitochondria in our patient. The patient described by Luft et al. (1962) did have increased metabolic activity and they considered that this occurred in spite of a defect in mitochondrial enzyme organization. The increased fat metabolism during exercise which we now report is compatible with their findings.

Our observations indicate that exercise provides a useful tool for examining the metabolism of abnormal mitochondria. An abnormal metabolic response to exercise has been found and we suggest that studies of further patients may allow delineation of specific patterns of metabolism of value in diagnosis.

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