Alpha-hydroxybutyrate dehydrogenase activity in sex-linked muscular dystrophy

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SYNOPSIS In two families with severe sex-linked muscular dystrophy, high levels of α-hydroxybutyrate dehydrogenase (HBD), lactate dehydrogenase (LD), aspartate transaminase (AspT), aldolase, and creatine phosphokinase (CPK) were found in the sera of three young affected males. In both families the mother had raised a level of HBD activity. Four sisters of the three affected boys had raised serum enzyme levels, and they are regarded as presumptive carriers of the disease. Biopsy specimens of dystrophic muscle had LD and HBD contents which were significantly lower than those of control specimens, while the HBD/LD ratios were markedly greater. Muscle from two unaffected members of the same family also exhibited high ratios, indicating the presence of electrophoretically fast LD isoenzymes, and this was confirmed by acrylamide-gel electrophoresis.

Measurement of the serum α-hydroxybutyrate dehydrogenase activity has provided a convenient means for the detection of a relative increase in the electrophoretically fast-migrating lactate dehydrogenase isoenzymes, LD1 and LD2 (Rosalki and Wilkinson, 1960; Plummer, Elliott, Cooke, and Wilkinson, 1963), and has proved to be a useful diagnostic test in myocardial infarction and other conditions in which LD1 and LD2 are released into the circulation (Elliott and Wilkinson, 1961; Konttinen, 1961; Pagliaro and Notarbartolo, 1961; Rosalki, 1963). Elevated serum HBD activities have been reported in patients with the clinically severe type of progressive muscular dystrophy (Pagliaro and Notarbartolo, 1961; Elliott and Wilkinson, 1963), and several investigators have reported a relative increase in the proportions of LD1 and LD2 in biopsy specimens of skeletal muscle taken from patients with this disease (Richterich, Gautier, Egli, Zuppinger, and Rossi, 1961; Wiene and Herpol, 1962; Dreyfus, Démos, Schapira, and Schapira, 1962; Wiene and Laurysens, 1962).

We recently had the opportunity to study muscle biopsy specimens and sera from a family (family A) with presumptive evidence of sex-linked inheritance for severe muscular dystrophy, and sera from another family (family B) with known sex-linked inheritance. The relative and total HBD and LD activities in muscle extracts were compared with the isoenzyme patterns obtained after electrophoresis on cellulose acetate foil and in acrylamide gel. The serum aspartate and alanine transaminase (AspT and AIT, formerly known as glutamic-oxaloacetic and glutamic-pyruvic transaminase (SGOT and SGPT), aldolase and creatine phosphokinase (CPK) levels were also determined since abnormal activity of these enzymes in the serum is known to occur in progressive muscular dystrophy (Sibley and Lehninger, 1949; Schapira, Dreyfus, and Schapira, 1953; Pearson, 1957; Dreyfus, Schapira, and Schapira, 1958; Thompson and Vignos, 1952; Ebashi, Toyokura, Momoi, and Sugita, 1959; Dreyfus, Schapira, and Démos, 1960; Thomson, Leyburn, and Walton, 1960; Hughes, 1963).

METHODS

CLINICAL In the two families, all available members were clinically examined and full details of family histories were noted. In all the affected boys, including those deceased, the onset of the disease was in infancy with rapid progression to immobility in early adolescence.
All the affected individuals had a history of pseudohypertrophy of the calves with initial onset of the condition in the muscles of the pelvic girdle and spread to those of the shoulder girdle. In family A (Fig. 1) biopsies of the triceps muscle were obtained from one of the two affected sibs, III2 and III6, the unaffected male sib, III4, and the unaffected sister, II2. Sera were available from 12 members of this family including the mother's parents. There was no history of the disease in any previous generation.

In family B (Fig. 2) sera were available from only one of the two living affected sibs (III6) and from five other members, including the mother's sister (II5), who has an unaffected son and daughter.

Some difficulty was experienced in obtaining suitable control muscle, as it has recently been shown that the LD isoenzyme patterns of skeletal muscles differ according to whether they consist of predominantly red or white fibres (Dawson, Goodfriend, and Kaplan, 1964; Brody, 1964; Rosalki, 1964). In red muscle, e.g., soleus, most of the enzyme occurs in the fast fractions LD1 and LD2, whereas in white muscle, e.g., gastrocnemius, the slow fractions are the characteristic forms. Two specimens of triceps muscle, taken at necropsy, were used as controls and these were supplemented by seven biopsy specimens of latissimus dorsi taken from patients undergoing thoracic surgery. In none of these cases was there a history of cardiac, hepatic, haematological, or muscular disease.

Blood was collected by venepuncture into plain glass bottles. The serum, which was separated within an hour, was stored frozen at −18°C. during transit and until enzyme activity was determined.

DETERMINATION OF ENZYME ACTIVITY The activities of SLD and SHBD were determined spectrophotometrically at 25°C. by the methods of Wróblewski and LaDue (1955) and Rosalki and Wilkinson (1960) respectively. Muscle extract, diluted with 0-067M-Sørensen phosphate buffer (pH 7.4) (0.1 ml.), was mixed with phosphate buffer (2.7 ml.), reduced NAD (0.25 mg. in 0.1 ml. phosphate buffer) was added, and the resultant mixture was allowed to equilibrate at 25°C. for 30 minutes. The substrate (0.022N-sodium pyruvate or 0.1M-sodium α-oxobutyrate, both at pH 7.4 in phosphate buffer) (0.1 ml.) was then added and the extinction at 340 mμ was measured at 25°C. Activities are expressed as micromoles of NADH2 oxidized per minute (i.u.).

Aspartate aminotransferase activity was determined by the method of Karmen (1955) and alanine aminotransferase (ALT) by the method of Wróblewski and LaDue (1956) both at 25°C.

Aldolase activity was determined on non-haemolysed
sera that had only been frozen once. A Biochemica test combination, based upon the method of Beisenherz, Boltze, Büchner, Czok, Garbade, Meyer-Arendt, and Pfeiderer (1953), was used. Activity was determined spectrophotometrically at 37°C and results are expressed as micromoles NADH₂ oxidized per minute (i.u.).

Creatine phosphokinase activity was determined by the colorimetric method of Dreyfus and Schapira (1961) and results are expressed as micromoles of creatine liberated from creatine phosphate per minute (i.u.).

Acrylamide gel electrophoresis The method of Raymond and Weintraub (1959) was used in conjunction with the vertical electrophoresis apparatus of Raymond (1962).

0.09M-Tris buffer of pH 9.2 was prepared by dissolving 2-amino-2-(hydroxymethyl)-1,3-propanediol (0.43 g.), ethylenediamine-tetra-acetic acid (0.43 g.), and boric acid (0.43 g.) in deionized water and adjusting the volume to 1 litre.

Dimethylaminopropionitrile (300 mg.) was added to 150 ml. of a 5% w/v solution of Cyanogum 41 (acrylamide containing 5% methylene-bis-acrylamide) in 0.09M-tris buffer (pH 9.2). Ammonium persulphate (300 mg.) dissolved in 2 ml. tris buffer was added and the resulting solution was transferred without delay to the electrophoresis apparatus where it was allowed to gel. After about 30 minutes, the apparatus was filled with tris buffer, and water at 0 to 5°C. was circulated through the cooling plates.

Bovine albumin (20 mg./ml) and a trace of bromophenol blue were added to each enzyme solution to serve as markers, and 15-50 µl. (so as to give approximately equal total quantities) was transferred to each slot. A potential of 25 to 30 volts/cm. was applied for two to three hours, after which the gel was removed from the apparatus and immersed in the staining solution. This consisted of the tetrathiazolium salt, MTT (15 mg.), NAD (12 mg.), and phenazine methosulphate (0.5 mg.) dissolved in a mixture of 0.5M-sodium lactate (25 ml) and 0.09M-tris buffer (75 ml). After 30 to 40 min. the gel was extracted with water to remove unreacted tetrathiazolium salt and fixed in 10% v/v acetic acid.

Preparation of muscle extracts The muscle specimens were frozen at -18°C. until required for use. Portions were sent by refrigerated air freight from London to Philadelphia packed with dry ice in a thermos flask. Comparable results were obtained in both laboratories.

Specimens (200 mg.) were washed free from blood with 0.9% NaCl and were then homogenized at 0°C. with 0.067M-Sörensen phosphate buffer at pH 7.4 (0.6 ml.) and a Potter-Elvehjem homogenizer with a Teflon pestle. The extracts were centrifuged at 3,000 g at 4°C. for 10 min. and the supernatant fractions were separated. Electrophoresis and enzyme assays were carried out on freshly prepared extracts. For enzyme determination, samples were suitably diluted (usually tenfold) with phosphate buffer.

RESULTS

Table I shows the HBD and LD enzyme levels in the four extracts of muscle from the available relatives in family A and in the control muscles. In all three brothers the LD activities were lower than in any one of the control series, and in the two affected members the difference was marked, but the enzyme levels for the mother were apparently normal. However, the four members of the family, especially the affected boys, had muscle HBD/LD ratios that were much higher than any of the control values.

Figure 3 illustrates the LD isoenzyme pattern obtained by acrylamide gel electrophoresis of the extracts of normal (white) muscle, dystrophic muscle, and muscle of the mother, a presumptive carrier. In both affected children most activity was observed in the fastest bands LD₁, LD₂, and LD₃, whereas LD₄ was faint and LD₅ was not detectable. The patterns for the mother and the unaffected brother revealed larger proportions of the slow bands but in both cases LD₁, LD₂, and LD₃ predominated, while in normal muscle LD₃, LD₄, and LD₅ were the principal components.

Figure 4 shows the effect of replacing lactate by α-hydroxybutyrate, a substrate but little affected by the slow moving isoenzymes characteristic of normal (white) muscle. It gives a clear demonstration of the qualitative difference between the lactate dehydrogenases of normal and dystrophic muscle.

There appears to be an inverse relationship

<table>
<thead>
<tr>
<th>TABLE I</th>
<th>ENZYME ACTIVITIES IN MUSCLE FROM MEMBERS OF FAMILY A AND NORMAL SUBJECTS</th>
</tr>
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<tbody>
<tr>
<td>Family A</td>
<td>Specimen</td>
</tr>
<tr>
<td>Pedigree No.</td>
<td></td>
</tr>
<tr>
<td>III</td>
<td>Dystrophic</td>
</tr>
<tr>
<td>III</td>
<td>Dystrophic</td>
</tr>
<tr>
<td>III</td>
<td>Normal</td>
</tr>
<tr>
<td>II</td>
<td>Normal</td>
</tr>
<tr>
<td>II</td>
<td>Normal</td>
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<tr>
<td>II</td>
<td>Normal</td>
</tr>
<tr>
<td>Mean</td>
<td>1.29</td>
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<tr>
<td>S.D.</td>
<td>0.54</td>
</tr>
<tr>
<td>Range</td>
<td>0.84-2.14</td>
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</table>

C. F. Boehringer und Soehne, GmbH, Mannheim, Germany.
American Cyanamid Co.
trophic boys, who also showed grossly elevated levels of AspT, aldolase, and creatine phosphokinase as compared with normal adults. In the serum of children over the age of a few weeks, the AspT, LD, and HBD activities were similar to those of adult serum. (We are indebted to Dr. S. B. Rosalki for the information that in his experience creatine phosphokinase levels are raised in infancy but fall to normal adult values by the fifth year of life. Aldolase activities, however, may be 50-100% higher in children up to the age of 10.)

In family A, the mother (IIb) and maternal grandmother (Ib) had high levels of SLD, SHBD, and aldolase. The maternal grandfather’s serum had raised levels of SHBD and SLD but the normal transaminase, aldolase, and creatine phosphokinase values suggest a pathological condition other than muscle disease. The father (IIc) of the affected boys showed all enzyme levels within the normal range except for a markedly high value for creatine phosphokinase. The sera for his two unaffected sons (III1, III4) gave values within the normal range, but the four unaffected daughters (IIId, IIIb, III7, and IIIg) showed increased levels of SHBD and AspT, while III1 and III7 also showed a rise in the SLD and CPK levels, and IIIg an associated increase in the SLD and aldolase activity. However, since IIIg is only 4 years old, her aldolase level is not necessarily abnormal.

With the exception of aldolase all the enzyme levels of the affected boy (III4) in family B were grossly elevated. An increase in the SHBD and creatine phosphokinase levels was shown by the ‘carrier’

![Image](https://via.placeholder.com/150)

**FIG. 3.** Acrylamide gel electrophoresis of LD isoenzymes of extracts of normal muscle, dystrophic muscle (III1), and muscle from the mother of family A (IIb).

![Image](https://via.placeholder.com/150)

**FIG. 4.** Acrylamide gel electrophoresis of LD isoenzymes of extracts of normal and dystrophic muscle treated with α-hydroxybutyrate in place of lactate.

Table II shows the serum levels of LD, HBD, AspT, AIT, aldolase, and creatine phosphokinase of the available members in both families. There was a high overall incidence of abnormal levels of dehydrogenase activity, the highest occurring in the dystrophic muscle, dystrophic muscle (III1), and muscle from the mother of family A (IIb).

The values for the corresponding sera from both families were much lower than the controls, the values for the corresponding sera were appreciably greater than the control figures.

**Table II**

<table>
<thead>
<tr>
<th>Sex</th>
<th>Subject</th>
<th>Age (yr)</th>
<th>LD (i.u./l)</th>
<th>HBD (i.u./l)</th>
<th>HBD/LD ratio</th>
<th>AspT (i.u./l)</th>
<th>AIT (i.u./l)</th>
<th>Aldolase (i.u./l)</th>
<th>CPK (i.u./l)</th>
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<td>140</td>
<td>0.81</td>
<td>17</td>
<td>12</td>
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<td></td>
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<td></td>
<td></td>
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</tr>
<tr>
<td>M</td>
<td>I1</td>
<td>66</td>
<td>284</td>
<td>192</td>
<td>0.68</td>
<td>12</td>
<td>9</td>
<td>2.0</td>
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<td></td>
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<td>65</td>
<td>271</td>
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<td>8</td>
<td>4</td>
<td>6.0</td>
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<tr>
<td></td>
<td>II1</td>
<td>39</td>
<td>154</td>
<td>122</td>
<td>0.79</td>
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<td>6</td>
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<td>38</td>
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<td>6</td>
<td>4</td>
<td>7.8</td>
<td>17</td>
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<td>0.67</td>
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<td>333</td>
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<td>7</td>
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<td>III11</td>
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<td>470</td>
<td>336</td>
<td>0.72</td>
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<td>17</td>
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<td>III3</td>
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<td>218</td>
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<td>0.72</td>
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<td>3</td>
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<td>3.8</td>
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<td>Family B</td>
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<td>M</td>
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<td>149</td>
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<td>205</td>
<td>150</td>
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<td>10</td>
<td>4.5</td>
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<tr>
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<td>22</td>
<td>180</td>
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<td>0.67</td>
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<td>3.6</td>
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<td>7.5</td>
<td>3.1</td>
<td>2</td>
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*Propositi.*
mother (IIa) and the unaffected half sister (IIIa) who also had a raised AspT level; the unaffected full sister (IIIb) had normal values. The unaffected brother (IIId) and the mother’s sister (IIe) showed no increase in any of the serum enzyme levels.

In neither family did the HBD/LD ratio for the serum distinguish between the affected males, their unaffected relatives, and normal controls.

DISCUSSION

In these familial studies, the degree of LD and HBD enzyme activity found in the muscle of the different members cannot be reconciled with a simple sex-linked mode of inheritance, ncr do the results for sera reflect the enzyme activity in muscle. The LD activity of the extracts from the dystrophic muscle, expressed per milligram of soluble protein, were only 6 to 62% of the corresponding values for the controls. It was also marginally lowered in the extract from the unaffected male sib, but appeared to be normal in the maternal specimen. In all four muscle extracts, however, the HBD/LD ratios were markedly greater than those for the controls, and the highest values were obtained with the dystrophic muscles. The ratios for the mother and unaffected male sib were intermediate between those for the affected sibs and the controls. These findings suggest the presence of a biochemical abnormality in the muscle of the unaffected individuals in whom no clinical signs of the disease can be found. An increase in the HBD/LD ratio has been shown to indicate a preponderance of the fast migrating isoenzymes LD1 and LD2 (Rosalki and Wilkinson, 1960; Plummer et al., 1963), and we have been able to demonstrate by electrophoresis increases in those isoenzymes relative to normal muscle. This confirms the earlier electrophoretic studies of dystrophic muscle (Richterich et al., 1961; Wieme and Herpol, 1962; Dreyfus et al., 1962).

In the present work, the increased HBD/LD ratio in muscle clearly identifies the affected boys and the ‘carrier female’, but such interpretations are made with caution in view of the abnormal value shown by the unaffected male sib. The result in the unaffected boy was confirmed by independent determinations in London and in Philadelphia, but at present we cannot offer an explanation. A difference in the HBD/LD ratios of the order observed, however, is unlikely to be due to errors arising during enzyme determination, since the precision of the latter is within ± 5%.

Determination of muscle enzyme activities in dystrophic muscle has recently been criticized because proliferation of connective tissue might lead to errors. In this work the muscle extracts were centrifuged to remove insoluble matter and the enzyme activities related to the soluble protein content. The results would be invalid only in the unlikely event of a compensating increase in the soluble protein content of muscle.

Several workers (Schapira et al., 1953; Dreyfus et al., 1958; Vignos and Lefkowitz, 1959; Pennington, 1962) have reported that dystrophic muscle has lower activities of the glycolytic enzymes, aldolase and phosphorylase, and of creatine phosphokinase and adenylate kinase than normal muscle. Our finding of reduced LD activity is consistent with that of Heyck, Laudahn, and Lüders (1963) but conflicts with that of Ronzoni, Berg, and Landau (1960) who found the LD level to be increased though the aldolase and creatine phosphokinase levels were reduced. This difference may in part be due to the duration of the illness and the gross wasting of the musculature in both patients whom we studied. However, the low figures found in their asymptomatic brother and mother cannot be attributed to such a cause.

A decrease in the LD2 content of muscle has been reported after experimental nerve section (Wiebe and Laurysens, 1962; Dawson et al., 1964; Laurysens, Laurysens, and Zondag, 1964) and by Brody (1965) has recently shown that in guinea-pig muscle there is loss of LD1. It may therefore be that changes in the LD isoenzyme content in human progressive muscular dystrophy are not specific for the dystrophic process, but are the consequence of changes in membrane permeability resulting in the disproportionate loss of the slow isoenzymes.

Whatever may be the mechanism involved, there is no doubt that the LD of dystrophic muscle, as shown during this investigation, is different from that of comparable normal white muscle. Since LD1 and LD2 are sensitive to inhibition by excess pyruvate, they are not capable of sustaining prolonged anaerobic glycolysis (Plagemann, Gregory, and Wróblewski, 1960; Cahn, Kaplan, Levine, and Zwilling, 1962). Such isoenzymes occur in heart muscle which functions in an aerobic environment, while normal white muscle isoenzymes LD1 and LD2 can continue to convert pyruvate into lactate in the presence of excess substrate. The predominance of LD1 and LD2 in dystrophic muscle may partly explain the three-fold decline in glycolytic activity as compared with normal muscle (Dreyfus et al., 1959).

In both families the serum enzyme levels for the dystrophic boys were grossly elevated, but of equal importance were the elevated levels in the unaffected females, all of whom are genetically ‘at risk’ as carriers of the harmful trait. The enzyme activities of the unaffected male sibs of the propositi in both families were within the normal range.
The serum aldolase has not proved to be very satisfactory for detection of the carrier state (Leyburn, Thomson, and Walton, 1961; Clayton, Wilson, and Carter, 1963) and determination of creatine phosphokinase has been recommended as the most helpful biochemical test for this purpose (Aebi, Richterich, Colombo, and Rossi, 1961; Hughes, 1962; Wilson, Evans, and Carter, 1965). Of the potential female carriers in family A, IIIa and IIIb exhibited abnormally raised serum creatine phosphokinase levels which were accompanied by high LD and HBD activities, thus giving an ominous prognosis for these two girls. Conversely, since IIIc showed normal serum creatine phosphokinase and LD activities and her HBD level was only marginally elevated, this girl is not likely to carry the harmful gene.

The youngest girl, IIIa, had normal creatine phosphokinase activity but, in marked contrast, high LD and HBD levels. It is thus impossible to discriminate for or against the carrier state in this girl: her serum aldolase level, which is very similar to that of her mother, may not be abnormal in a child of 4 years. The question would probably be resolved by repetition of the enzyme studies when the child is older.

An interesting finding in family A is the high creatine phosphokinase activity observed on several occasions in the serum of the father (II1), which has no apparent clinical explanation. As he was unemployed at the time, his raised levels cannot be attributed to strenuous muscular work. Such values are more likely to be found in a heterozygote carrier of the trait, but there is no evidence for any mode of inheritance other than sex-linked.

In family B, with known sex-linked inheritance, all the serum enzymes of the affected boy have high levels of activity. The increased serum HBD, AspT, and creatine phosphokinase activities shown by the eldest half-sister (IIIa) is again a grave prognosis for the 'carrier state', but the youngest sister (IIIb), in whom normal serum enzyme activities were observed, is not likely to be a carrier of the harmful gene. There is a very marked difference between the high serum HBD and creatine phosphokinase level for the carrier mother (II1) and the normal level of her sister (II2) who has an unaffected son.

In neither family was there an increase in the SHBD/SLD ratio above the upper normal limit (0.81), nor was there any significant difference between these ratios in the affected, the unaffected, and the control series. This is contrary to previous findings of high ratios (Elliott and Wilkinson, 1963); the difference may be due to the greater degree of genetic and clinical homogeneity of the present data.

Although the serum LD and HBD activities are not specific for identifying carrier females they do provide a useful adjunct to the evidence provided by the serum creatine phosphokinase level in these women. Schapira and Démos (1962) have reported a slight but significant increase in the maternal serum LD levels of 27 patients with dystrophy; the present work suggests that measurement of the serum HBD activity would give a more pronounced increase that might possibly be of greater diagnostic value than the LD determination. No single biochemical test has a high degree of accuracy when used for this purpose and it would appear that all the relevant enzyme assays are required to assess the potential carrier genotype of those girls in families with sex-linked muscular dystrophy.

We are indebted to Professor P. R. Allison, Nuffield Professor of Surgery, Radcliffe Infirmary, Oxford, for providing the control muscle and sera; Dr. H. Ellis, Churchill Hospital, Oxford, and Dr. E. Addenbrooke, Gloucester Royal Hospital, for referring the families; Dr. J. Thorne, Neurological Department, Radcliffe Infirmary, for his assistance in taking the muscle biopsies from members of family A; Dr. C. W. Hyde for some of the creatine phosphokinase determinations, and to the E-C Apparatus Corporation, Philadelphia, Pennsylvania, for providing the vertical electrophoresis apparatus. Technical assistance by Miss Margaret Cromwell-Thomas, kindly financed by C. F. Boehringer und Soehne, G.m.b.H., Mannheim, Germany, and a grant for scientific assistance (to J.H.W.) by the Medical Research Council are gratefully acknowledged. One of us (J.H.W.) wishes to thank Dr. Robert F. Norris and the University of Pennsylvania for facilities to carry out some of the work during a visit to Philadelphia. This work was supported in part by U.S. Public Health Service Grant No. GM-10876 to Dr. Samuel Raymond, Pepper Laboratory of Clinical Medicine, Hospital of the University of Pennsylvania.

REFERENCES


hum. Genet.*, 25, 41.
Plagemann, P. G. W., Gregory, K. F., and Wróblewski, F. (1960),
Richterich, R., Gautier, E., Egli, W., Zuppinger, K., and Rossi, E.
ment. Dis.*, 38, 721.
— (1964). *Personal communication.
103, 551.
med. J.*, 2, 1276.
873.
J.*, 1, 750.