Application of new ultra-micro automated spectrophotometric determination for serum hydroxybutyrate dehydrogenase activity to diagnosis of myocardial infarction

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An automated spectrophotometric assay for serum $\alpha$-hydroxybutyrate dehydrogenase (SHBD) activity has been developed. This utilizes the change in absorbance at 340 nm consequent upon oxidation of NADH. Determinations are made at 38°C. at a rate of 60 per hour, and measurement of test and blank requires only 15 ml of serum. Sera of known SHBD activity assayed by a manual kinetic spectrophotometric technique at 25°C. are used as standards.

The normal range determined for 648 subjects was 40–200 mIU/ml. There was no difference between the sexes, and in both, SHBD activity increased significantly with age. Sera from 528 patients suspected of having sustained myocardial infarction were assayed. Of 197 patients in whom infarction was established on clinical and electrocardiographic criteria, 150 had raised SHBD activity. Of 286 patients in whom infarction was ultimately excluded, 67 had raised SHBD activity. The accuracy of SHBD activity in confirming the diagnosis of myocardial infarction was comparable with that of aspartate aminotransferase activity as determined spectrophotometrically in the same subjects, and there were advantages in carrying out both procedures. Storage of sera at $-20$°C. resulted in a 15 per cent increase in SHBD activity, whereas a fall of 5 per cent occurred after 7 days at $4$°C.

Human serum contains an enzyme catalysing the reversible reduction of $\alpha$-oxobutyrate according to the following equation:

$$\alpha$$-oxobutyrate + NADH$_2$ \rightleftharpoons $\alpha$-hydroxybutyrate + NAD

The enzyme concerned has been named serum hydroxybutyrate dehydrogenase (SHBD), but its activity is a function of the lactate dehydrogenase isoenzymes (Rosalki and Wilkinson, 1960). These are composed of two molecular subgroups, and it is the faster-moving isoenzymes rich in the B subgroup that mainly account for SHBD activity (Latner and Skillen, 1968).

The role of SHBD determinations as an aid to the diagnosis of myocardial infarction has been established (Elliott and Wilkinson, 1961; Elliott, Jepson, and Wilkinson, 1962; Konttinen and Halonen, 1962; Rosalki, 1963; Rosalki and Wilkinson, 1964), though raised levels are known to occur in other conditions, mainly diseases of the liver, megaloblastic anaemias, and the muscular dystrophies (Elliott and Wilkinson, 1961, 1963; Johnston et al., 1966). But the availability of other serum enzyme tests such as aspartate and alanine aminotransferase (SGOT and SGPT) and ATP:creatine phosphotransferase (CPK) has rendered the position of SHBD controversial; some authors consider that its determination in suspected cases of myocardial infarction is most advantageous (Preston, Batsakis, and Briere, 1964; Stuart et al., 1965), while others are of the opinion that it does not contribute more than alternative estimations to diagnostic accuracy in this condition (Nissen, Ranlev, and Weis-Fogh, 1965; Griffiths, 1966; Smith, 1967).
In the belief that an automated method for SHBD determination would enable a more rapid and economical throughput of samples, we have devised an ultra-micro automated spectrophotometric procedure for SHBD determination which has many advantages over previous techniques (Schwartz, Kessler, and Bodansky, 1961; Strandjord and Clayson, 1966; Dube, Hunter, and Knight, 1968). This report describes its application to a large series of non-hospitalized control subjects, and to patients in whom the possibility of myocardial infarction was raised by the physician. The diagnostic accuracy of SHBD has been compared with that of the aminotransferases in the latter group.

Material and method

Apparatus The following items were purchased from Technicon Instruments Co. Ltd., Chertsey, Surrey: proportioning pump module; sampler II module; 120/hr. cam with 1:2 sample: wash ratio; glass coil (105-1173-1) length 1219 cm., 2.0 mm. I.D.

The following items were purchased from Beckman Instruments Ltd., Glenrothes, Fifeshire, Scotland: DB spectrophotometer (Cat. No. S1402H); 5" linear/log recorder (Cat. No. S9307H); scale expander (Cat. No. 151060); micro-flowcell (Cat. No. 97290).

Reagents

(1) 0.067 M phosphate buffer, pH 7.40

This is prepared by dissolving 7.55 g. Na2HPO4 (anhydrous) and 1.81 g. KH2PO4 (anhydrous) in water, adjusting the pH to 7.40 and the volume to 1 litre; stable for one week at 4°C.

(2) Substrate solution

21.0 mg. β-NADH (disodium salt) and 100 mg. α-oxo-α-butyrate (monosodium salt) are dissolved in 100 ml. of above buffer to give final concentrations of 0.3 mM and 8.1 mM respectively. This is sufficient for 70 analyses, and is prepared fresh each day.

(3) Standards

Chemtrol (Clinton Laboratories, Los Angeles), approximate SHBD activity 900 milli-International Units per ml. (mIU/ml.) at 25°C., was diluted in 0.15 molar NaCl to give 4 samples covering the range 100-500 mIU/ml. at 25°C., each being assayed in duplicate by the manual spectrophotometric technique of Rosalki and Wilkinson (1960); they may be kept at 4°C. for up to one week but must be reassayed each day of use.

Technique

The manifold is shown in Fig. 1. The sampler II is governed by a 120/hr. cam with a sample time of 10 seconds and a wash time of 20 seconds. The bubbler consists of an AO Technicon T-piece with small tygon pump tubing inserted in the side-arm and adjusted to give not less than 10 bubbles in 10 seconds. The sample stream is introduced through a standard A6 fitting, and equilibrates with the reagent stream in a single mixing coil. The volume of the incubation coil is 35 ml. and the incubation time at 38°C. is approximately 10 minutes.

The absorbance of the substrate solution at 340 nm is approximately 1.45. The range-expander is set to give a recorder scale of 0-5-1.5. The system has a range equivalent to SHBD activities of 0-600 mIU/ml. at 25°C. When sera are pumped with substrate solution, the reduction in absorbance at 340 nm. is proportional to SHBD activity, the latter being determined using a calibration curve drawn from the peaks given by the standards. Blank determinations are carried out by pumping sera with buffer and NADH alone, the final NADH concentration being 0.2 mM to give an absorbance at 340 nm. of approximately 1.0. The effect of serum is to increase the apparent absorbance of the buffered NADH, and this increment, which in clear sera corresponds to about 10 mIU/ml. and in turbid sera to as much as 100 mIU/ml., is converted to SHBD activity from the calibration curve and added to that obtained with the same serum in the presence of substrate solution. The amount of serum sampled is 7.5 μl. for the test and 7.5 μl. for the blank.

Carry-over This was assessed by sampling a serum of low activity before and after a serum of high activity (Fig. 2). Nine replicates of the high serum gave a mean of 475 mIU/ml. (SD 5.8 mIU/ml.). Five replicates of the low serum when sampled before the high serum gave a mean of 101 mIU/ml. (SD 4.2) and when sampled after the high serum a mean of 117 mIU/ml. (SD 4.5). The carry-over was thus 3.4 per cent of the high-activity serum.

FIG. 1 Manifold for automated spectrophotometric SHBD determination. Flow rates given as ml./min. SMC represents Single Mixing Coil. AO and A6 are standard Technicon equipment modified only where indicated in text.
FIG. 2 Reproducibility and carry-over in automated spectrophotometric SHBD determination. Recorder tracing of peaks given by low-activity serum sequentially, high-activity serum sequentially, and the two sera when run alternately.

Comparison of manual and automated method When 36 sera were assayed by both techniques at 38°C, the automated method gave higher results (mean 150 units higher, SD 14·8). This represented 5 per cent of the average activity, and the difference was not statistically significant.

Precision Thirty samples whose HBD activity ranged from 115 to 705 mIU/ml, were analysed on two independent runs carried out the same day. The average difference between the duplicates was ±1·6 per cent and the largest difference in any one pair ±4·1 per cent.

Stability One hundred and two samples were analysed fresh, and again 7 days later after storage at −20°C. On average, HBD activity rose by 15·4 per cent after storage, 89 samples showing increased activity, 10 decreased activity, and 3 unaltered activity. This rise when evaluated by the paired t-test was highly significant ($t_0 = 10·63; p < 0·001$). Twenty-six samples were analysed fresh, and again after 7 days at 4°C. A mean fall of 5·0 per cent in HBD activity occurred, 19 samples showing a decrease, 5 an increase, and 2 unaltered activity. This change was also statistically significant ($t_0 = 2·31; p < 0·05$). It has been established that the slow LDH isoenzymes associated with molecular subgroup A are partially inactivated by freezing (Zondag, 1963; Kreutzer and Fennis, 1964), but enhancement of activity of subgroup B by freezing has not previously been demonstrated.

Clinical material

Normals Three groups of subjects were studied.

a) Healthy laboratory personnel aged 16 to 37 years sampled in the post-absorptive state between 11 and 12 a.m. (77 subjects).

b) Subjects attending a well-patient screening clinic aged 18 to 79 years sampled between 2 and 6 p.m. (407 subjects).

c) Ambulant subjects aged 65 and over attending a special screening clinic for the elderly. Many were under treatment for chronic illness but were well compensated at the time of examination. Subjects with ischaemic heart disease and megaloblastic anaemias were excluded. Blood was drawn between 2 and 6 p.m. (164 subjects).

The total number of subjects was thus 648; the distribution with respect to age and sex is given in Fig. 3. Serum was separated within an hour, divided into aliquots for each subject, and stored at −20°C. Over the next 6 weeks, in addition to SHBD activity in all subjects, the following determinations were carried out on subjects in groups a and b: urea, uric acid, cholesterol, alkaline phosphatase, inorganic phosphate, and total protein by Auto Analyser techniques; calcium and magnesium concentration by Atomic Absorption Spectroscopy; the activities of SGOT and SGPT (Henry et al., 1960)
**Serum hydroxybutyrate dehydrogenase activity in myocardial infarction**

![Diagram](image)

**FIG. 3** Change of mean SHBD activity with age. Solid line, men; broken line, women. Inset figures represent total number of men (above) and women (below).

5'-nucleotidase (Belfield and Goldberg, 1968), and creatine phosphokinase (Worthy, Whitehead, and Goldberg, 1970). The above parameters, together with social class, blood pressure, weight, and percentage overweight (Documenta Geigy, 1956) were studied for a possible influence on SHBD activity using an ICL 1907 computer with a multiple linear correlation programme giving the values and significance of the correlation coefficient R.

**Ischaemic heart disease** The material for this part of the study consisted of sera submitted to the laboratory from patients with suspected myocardial infarction, one sample being taken shortly after admission and a second 24 hours later. A third sample was taken where the results of the first two were ambiguous. The activities of SGOT and SHBD were estimated on all sera. SGPT activity was estimated on the first sample, but not subsequently, unless the initial transaminase ratio raised the possibility of hepatic involvement, i.e. SGPT:SGOT > 1. The upper normal limits for SGOT and SGPT in our laboratory are 40 and 30 mIU/ml., respectively.

After six months of operating this system, the entire material (528 cases) was reviewed and, without knowledge of the enzyme results, each subject was assigned to one of the following 4 categories.

(1) **Positive**

Serial electrocardiographic examination confirmed the diagnosis of infarction which was supported by convincing clinical evidence, such as characteristic pain, shock, hypotension, and increase in temperature, white cell count, and erythrocyte sedimentation rate.

(2) **Electrocardiogram-positive cases without clinical evidence**

Despite unequivocal evidence of recent infarction on the tracings, patients lacked the characteristic pain (silent infarct), and investigation was usually prompted by onset of cardiac failure, arrhythmia, hypotension, or embolism. The precise time of infarction could not always be ascertained.

(3) **Clinically-positive cases without confirmation on electrocardiogram**

In these patients, the electrocardiogram gave normal or ambiguous results, often because of previous infarction or the presence of arrhythmia. But the typical clinical features described under (1) above were present, and, despite careful investigation could not be explained on grounds other than infarction.

(4) **Negative**

Electrocardiograms in these patients gave normal or equivocal results. The clinical features were rarely pathognomonic of infarction and, with few exceptions, were adequately explained by a final diagnosis other than that of myocardial infarction.

**Results**

**Normals** The mean and SD of SHBD activity for men was 128 ± 35 mIU/ml, and for women 126 ± 33 mIU/ml. The distribution was typically Gaussian, 13 cases lying above 200 mIU/ml., and 12 below 60 mIU/ml. This range of 60–200 mIU/ml. thus embraced 95 per cent of the population and fulfilled acceptable criteria for a normal range. Though 53 of the 164 subjects in Group C (aged 65 and over) were under treatment for chronic illness, their inclusion had no effect upon the data.

A striking correlation between age and SHBD activity was seen in women (Fig. 3), the mean value almost doubling between the second and ninth decades (R = 0.29; p < 0.001). In men, the influence of age upon SHBD activity was less apparent (R = 0.227; p < 0.005). An inverse relation between SHBD activity and serum urea concentration was seen in men (R = -0.281; p < 0.001), and a direct relation between SHBD activity and serum magnesium concentration in women (R = 0.209; p < 0.01). There was no indication whether the former relation was due to chemical inhibition by urea of LDH isoenzymes (Withycombe, Plummer, and Wilkinson, 1965; Emery, Moores, and Hodson, 1968) or to a dependence of urea concentration and SHBD activity upon renal function.
Neither explanation clarifies the difference in behaviour seen in both sexes.

**Ischaemic heart disease** All but 18 of 528 cases investigated were placed in one of the four categories previously defined; those cases defying classification are not considered further.

For the remainder, the highest recorded values for SHBD and SGOT were compared. These are presented for positive and negative cases in Fig. 4 and 5, respectively. Values in excess of 200 and 40 mIU/ml, respectively, were considered raised, and indicative of infarction.

**Positive (179 cases)**
Raised SHBD activity was detected in 150 cases and raised SGOT in 144 cases. Since 12 of the cases with raised SGOT activity had even higher SGPT activity in the same specimen, aminotransferase estimations on these subjects ought to be classed as equivocal rather than diagnostic.

Seven cases died within 8 hours of onset of symptoms. Of these, 4 had raised SGOT and only 2 had raised SHBD, emphasizing the more rapid rise of the former after infarction. Fifteen cases were not seen until at least 5 days after onset of symptoms. All had normal SGOT activity, but 6 had raised SHBD activity; this confirms the superiority of the latter in detecting late cases of myocardial infarction. Seventeen cases with consistently normal SHBD activity and 16 with consistently normal SGOT activity had at least one sample taken during the 12- to 72-hour period after infarction when maximal enzyme increases are to be expected.

**Negative (286 cases)**
Raised SHBD activity was detected in 67 cases. Haemolysis of the sample accounted for 12 of these, Addisonian anaemia for 2, pericarditis for 3, renal failure for 3, cardiac surgery for 3, non-thoracic major surgery for 4, pulmonary embolism for 3, cardiac arrhythmias for 5, severe angina for 5, congestive cardiac failure for 4, and bronchial carcinoma for 1. The remaining 22 suffered from miscellaneous conditions, mainly abdominal, occasionally associated with raised SHBD activity (Elliott and Wilkinson, 1963).

Of 62 patients with raised SGOT activity, the ratio of aminotransferase activities (SGPT: SGOT > 1) in 25 was suggestive of hepatic disease, and the result could not be
considered as truly false-positive. This left 37 cases in which the increase could be ascribed to haemolysis of the sample in 8, pericarditis in 4, cardiac or major surgery in 4, congestive cardiac failure in 3, cardiac arrhythmia in 2, angina and Addisonian anaemia in 1 case each. A firm diagnosis was not possible in the remaining 14.

Positive cases on electrocardiogram, without clinical evidence (14 cases)
In 7, SHBD activity was raised and in 7 it was normal. Two subjects whose infarction occurred more than one week before admission showed raised activity; 2 other subjects in this category had normal activity. SGOT activity was normal in 8 cases and raised in 6, but in 3 of the latter the SGPT:SGOT ratio was greater than unity and all had congestive heart failure; on this basis alone, infarction would not therefore have been suspected. All 4 cases admitted later than one week from the date of infarction gave normal values for SGOT.

Clinically-positive cases without confirmation on electrocardiogram (31 cases)
In 23, SHBD activity was raised and in 8 it was normal: the latter included one case examined for the first time 16 days after suspected infarction and one case dying in 8 hours of onset of symptoms. SGOT activity was normal in 7 cases and raised in 24, but in 2 of the latter the SGPT:SGOT ratio was greater than unity; aminotransferase determinations in these patients thus did not provide unequivocal support for the clinical diagnosis.

Discussion

Technical aspects Previous automated determinations of SHBD activity (Schwartz et al., 1961; Strandjord and Clayson, 1966; Dube et al., 1968) utilize direct calculation of the change in absorbance arising from conversion of NADH to NAD as a function of time for the expression of results. This necessitates accurate knowledge of the incubation time, the path length of the flow cell, and the pumping rates of reagent and sample, some of which may vary from day to day. Absorbance requires to reach a steady plateau, and this may be expensive in terms of both sample and reagent. The assumption is made that there is a linear relation between enzyme activity and the change in absorbance, and this may not be true of high and low activities. The loss of zero-order kinetics is also ignored.

The present method employs as standards sera previously assayed by an accurate kinetic technique for SHBD activity. Pumping rates and incubation time need not be accurately known, and loss of zero-order measurements is compensated as between standards and unknowns. Changes in background reagent-stream absorbance are carefully monitored, and a further standard with every 9 unknowns corrects for possible instrument drift. If serum is aspirated continuously, 85 per cent of the plateau absorbance is reached in 10 seconds, 96 per cent in 30 seconds, and 100 per cent in 40 seconds. It is therefore possible to present sera at 30-second intervals, sampling for 10 seconds and washing for 20 seconds. This reduces carry-over from sample to sample to approximately 4 per cent, and permits 60 sera with blanks to be assayed per hour, with a total sample volume of only 15 µlitre for combined test and blank determinations. The speed of sampling is far greater and the volume of sample far smaller than in any previous method, and by carrying out the automated reaction at 38°C sensitivity is increased and the relative significance of the blank effectively diminished.

Clinical aspects In this series, estimation of SHBD activity was as reliable as aminotransferase determinations in confirming the diagnosis of myocardial infarction. In many clinical situations, both aminotransferases require to be estimated, especially where possible hepatic involvement is concerned; if this is not done, an unacceptably high false-positive rate ensues. The choice of a single enzyme determination would thus favour SHBD. The superiority of SHBD in detecting cases presenting several days after the onset of infarction is especially impressive, and confirms previous work (Elliott et al., 1962; Rosalki, 1963; Preston et al., 1964; Stuart et al., 1965). The fact that aminotransferase activities were determined by a kinetic spectrophotometric technique requires emphasis; these have been strongly recommended by Amador, Franey, and Mas sod (1966) in preference to colorimetric techniques for the diagnosis of infarction, though this position has been disputed by Crowley (1967). Our experience accords with that of Amador et al. (1966). Before undertaking the present work, we surveyed a series of patients classified according to the criteria already outlined in whom 4 consecutive aminotransferase determinations spanning the period of 8–72 hours after onset of symptoms were carried out by a colori-
metric technique (King, 1958). Only 30 of 46 positive cases gave raised aminotransferase values with a ratio of SGPT:SGOT less than unity; in 5 SGOT was raised but was less than SGPT activity. Seven of 57 negative cases had raised SGOT activity only, and a further 10 had increases in both aminotransferases, SGPT being the higher of the two. The results obtained by the present SHBD method have been compared in the Table with those of other authors using various other methods. It is difficult to be sure that the criteria employed by these authors for classifying a patient as positive or negative were the same as ours, or that the timing of blood samples was satisfactory in all cases.

**Table**  Percentage of cases positive and negative for myocardial infarction showing raised SHBD activity in present and previous reports

<table>
<thead>
<tr>
<th>Authors</th>
<th>Per cent positive cases</th>
<th>Per cent negative cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elliott et al.</td>
<td>100</td>
<td>37</td>
</tr>
<tr>
<td>Preston et al.</td>
<td>97</td>
<td>14</td>
</tr>
<tr>
<td>Stuart et al.</td>
<td>100</td>
<td>63</td>
</tr>
<tr>
<td>Griffiths</td>
<td>94</td>
<td>21</td>
</tr>
<tr>
<td>Smith (1967)</td>
<td>94</td>
<td>7</td>
</tr>
<tr>
<td>Present</td>
<td>84</td>
<td>23</td>
</tr>
</tbody>
</table>

The percentage figures given for the present series include every positive and negative case, irrespective of factors such as timing of the sample or haemolysis of the specimen which tend to produce unfavourable results. The most fundamental comparison is with the SGOT results in the same series, since the latter estimation was made by the most accurate technique currently available, and both estimations were carried out as part of the routine service of a hospital laboratory. It is clear that SHBD determinations are at least as satisfactory as SGOT determinations in confirming a diagnosis of myocardial infarction, and that there is no total agreement between the results in every case — in other words each of the two estimations has something to offer. It is also apparent that even when extenuating circumstances are taken into account, a significant number of patients give false-negative or false-positive results for one or both determinations. Realization by the clinician that results obtained in a routine service rarely match the enthusiastic claims based upon research studies will do much to ensure the proper assessment of serum enzyme determinations in relation to myocardial infarction.

From the standpoint of fitting SHBD determinations into the work-load of the general laboratory, the present technique has considerable advantages. The samples may be stored at −20°C or at 4°C provided that the normal range is adjusted to take account of storage conditions, and all specimens are treated similarly. Micro-tubes containing 20 to 50 μl. serum are required, thereby minimizing the need for storage space. A week's collection of sera may be analysed in the course of 60 to 90 minutes. The demands on technician time and machine time are minimal, the cost of the reagents modest, and much useful information is available to the clinician, supplementing that obtained from ward procedures, electrocardiography, and aminotransferase estimations. It is in this manner that we recommend the employment of the present technique.

We wish to thank the Medical Staff of the United Sheffield Hospitals who provided blood samples and clinical details of the patients, Dr. Eric Wilkes who provided sera from both screening clinics, and the Technical Staff of the Department of Chemical Pathology who co-operated in this study. We are also indebted to Mr. A. J. Handyside and Mr. K. Trout for help in the computer analysis of the data, the Department of Medical Photography, Royal Hospital, Sheffield, for Fig. 1-5, and Dr. Arthur Jordan in whose laboratory much of this work was carried out.

**References**


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