A routine method for the determination of phosphogluco-

stance isomerase activity in body fluid

J. E. HORROCKS, J. WARD, AND J. KING

From North Lonsdale Hospital, Barrow-in-Furness, Lancashire

Phosphoglucone isomerase, the trivial name for the enzyme D-glucose-6-phosphate ketol-isomerase, code number 5.3.1.9 (Report of the Commission on Enzymes of the International Union of Biochemistry, 1961) catalyses the isomerization of D-glucose-6-phosphate to D-fructose-6-phosphate.

\[
\begin{align*}
\text{D-glucose-6-phosphate} & \rightleftharpoons \text{D-fructose-6-phosphate} \\
\end{align*}
\]

Phosphoglucone isomerase is a widely distributed glycolytic enzyme present in liver, muscle, bone, brain, lung, erythrocytes, and leucocytes in decreasing order of activity. Like other ubiquitous enzymes, activity is increased after myocardial infarction (Siegel and Bing, 1956; White, 1956; Bing, Castellanos, and Siegal, 1957). As with other glycolytic enzymes, such as lactate dehydrogenase and aldolase, in inflammatory effusions (Horrocks, King, Waind, and Ward, 1962) the phosphoglucone isomerase activity of the effusion fluid is often greater than that of the serum and is of no value in differentiating between malignant and benign origin (unpublished observations).

The serum activity is greatly increased in acute hepatitis (Bruns and Jacob, 1954; Bruns, 1957; Bing et al., 1957; Bodansky, Krugman, Ward, Schwartz, Giles, and Jacobs, 1959). Israels and Delory (1956), Israels, Delory, Hnatiuk, and Friesen (1958) and Blanchaer, Green, MacLean, and Hollenberg (1958) found raised activity in myelocy- cytic leukaemia but not in the leucocytosis of infection or in lymphocytic leukaemia.

Bodansky, Gershten, and Wilson (1954) and Myers and Bodansky (1957) reported the assay to be useful in following progress during palliative treatment for metastatic carcinoma of the breast and also of the prostate (Bodansky, Calitri, and Wilson, 1955). In this respect, interest in the enzyme has recently been renewed by the work of Jegatheesan and Joplin (1962) who confirm and extend the previous findings.

The equilibrium of the reaction catalysed favours glucose-6-phosphate, but because of the convenient methods of measuring fructose by the Seliwanof reaction (Roe, 1934; Roe et al., 1949) most serum enzyme assay techniques have used a substrate of glucose-6-phosphate (Bodansky and Calitri, 1954; Glock, McLean, and Whitehead, 1956; British Drug Houses, 1960). A coupled spectrophotometric method taking advantage of the favourable equilibrium given by a substrate of fructose-6-phosphate appeared possible. Purified glucose-6-phosphate...
dehydrogenase and coenzyme II (nicotinamide-adenine-dinucleotide phosphate, N.A.D.P.) included in the reaction mixture would oxidize the glucose-6-phosphate formed and enable the reaction to be monitored by following the increase in optical density at 340 m\(\mu\), at which wavelength the reduced coenzyme has a peak absorption. However, this procedure is complicated by the presence in human blood of 6-phosphogluconate dehydrogenase which also utilizes N.A.D.P. in the oxidation of 6-phosphogluconate, some quantity of which would result from the secondary reaction. 6-Phosphogluconate is also reported to be inhibitory to phosphoglucoisomerase (Parr, 1956). This approach, although used by Rose and O'Connell (1961) in isotope studies with purified enzyme preparations, appears to be too expensive and too complex for routine purposes.

The method of Bodansky and Caliti (1954) only requires incubation for 30 minutes at 37°C. using an acetate-veronal buffer, pH 7-4, but includes protein precipitation. The modified procedure (British Drug Houses, 1960) of Glock et al. (1956) omits protein precipitation but requires 60 minutes' incubation at 37°C. in a glycylglycine buffer, pH 7-6. Further, this latter method only measures activity in the normal range of human serum and tests must be repeated using smaller quantities of specimen or shorter periods of incubation if raised serum activity is encountered. Grave miscalculations of phosphoglucoisomerase activity are also possible unless a complete specimen and reagent blanks are included.

A point which makes difficult the comparison of results published by different workers and leads to much confusion is the relative chromogenicity of fructose and fructose-6-phosphate. The situation is also complicated by the fact that although the original method of Roe (1934) used colour development by heating at 80°C. for eight minutes, and the time was extended to 10 minutes in the later procedure (Roe et al., 1949), many workers have heated for 15 minutes to develop the cherry red colour. Roe and his colleagues do not mention any difference in colour intensity between fructose and its phosphate esters; indeed the 1-ester and 1,6 diester are rapidly hydrolysed by heating in the strongly acid solution. Bodansky (1953), using the original Roe method, found that fructose-6-phosphate, which is stable under the conditions of colour development, only gives 61% of the colour given by fructose, and Glock et al. (1956) give a value of 65% measuring the optical density at 490 m\(\mu\). Hers, Beaufays and De Duve (1953), using a modification of the method of Roe et al. (1949), gave a value of 82% of that of fructose for the chromogenicity of fructose-6-phosphate whereas Dische (1951) found that both fructose and fructose-6-phosphate gave equal colour intensity. Kahana, Lowry, Schulz, Passonneau, and Crawford (1960), using enzymatically prepared fructose-6-phosphate, agree with Dische when using the original procedure but found that fructose-6-phosphate actually gave 5% more colour than fructose when the latter method was used, regardless of whether heating is for eight or 15 minutes.

Fructose-6-phosphate of analytical grade purity is not obtainable and therefore a secondary standard of fructose must be used. It is proposed that this fructose standard be used without any modification or consideration of the possible relative chromogenicity of fructose and fructose-6-phosphate. This is arbitrary but at least a permanent and comparable measure of activity is established rather than one which is continually varying with empirical figures given by workers using compounds of purity which are unknown and not reproducible. Further, in actual assays, if at the end of the reaction period proteins are precipitated by means of zinc sulphate followed by barium hydroxide, a procedure which results in fructose-6-phosphate, but not fructose, being adsorbed to the precipitate, a colour reaction is still obtained. This indicates that some (enzymic) hydrolysis of fructose-6-phosphate occurs and therefore in actual fact an unknown proportion of both the ketose and its 6-ester is present in the reaction mixture which further justifies the use of an arbitrary fructose standard.

A study of the kinetics of phosphoglucoisomerase activity was undertaken in the hope that a simple, rapid method of assay applicable to body fluids might be elaborated.

**REAGENTS**

M10 BORATE BUFFER (pH 7-8) Dissolve 6-184 g. boric acid (AR) and 7-456 g. potassium chloride (AR) in water, add 53 ml. N/10 NaOH and make up to 1 litre with water.

BUFFERED SUBSTRATE Dissolve 3 mg. disodium glucose-6-phosphate per millilitre of buffer. Use on the same day as prepared.

30% HYDROCHLORIC ACID Make up 850 ml. hydrochloric acid (AR) to 1 litre with water.

RESORCINOL-ThIOUREA REAGENT Resorcinol (AR), and 250 mg. thiourea dissolved in 100 ml. glacial acetic acid (AR). Store in a brown bottle and discard when the solution becomes discoloured.

COLOUR REAGENT Mix together in the proportions 7:1:1 30% hydrochloric acid, resorcinol-thiourea reagent, and water. Use on the same day as prepared.
STANDARD SOLUTION  Fructose, 54 mg., is dissolved in 1 litre of 0-25% benzoic acid. Of this 1 ml. is equivalent to 100 milli international units under the assay conditions. The solution is stable for months at room temperature.

PROCEDURE

Into two tubes labelled 'test' and 'blank' pipette 1 ml of buffered substrate and place in the water bath at 37°C. for a few minutes to reach bath temperature, then add 0.1 ml of serum to the 'test' tube and begin timing. Exactly 30 minutes after addition of the specimen remove the tubes from the bath, add 0.1 ml specimen to 'blank' tube and 9 ml of colour reagent to both. Place both tubes in the 75°C. water bath. At the same time standard and standard blank tubes containing respectively 1 ml. of standard and 1 ml. of water treated with 9 ml. of colour reagent are also placed in the bath. After 15 minutes at 75°C. the tubes are immediately cooled in cold running water and the pink-red colour read in the next few minutes using a blue-violet filter or transmission at 410 mµ.

Since 0.1 ml. of specimen is used with a 30 minute reaction period and the standard is 0.0003M this is equivalent to

\[
\frac{0.0003 \times 1000}{30} \times 1 \\mu\text{mole/min./ml., i.e., international units (i.u.)}
\]

or 100 mµ mole/min./ml. or m i.u. The calculation of activity is then given by

\[
\frac{(\text{Test-blank})}{(\text{Standard-blank})} \times 100 \text{ m i.u.}
\]

King and Campbell (1961), in introducing the international units, proposed that activity be expressed per litre of specimen. The Report of the Commission on Enzymes of the International Union of Biochemistry (1961), however, recommends that activity be expressed per millilitre of specimen. Activity expressed in m i.u. per millilitre is numerically the same as i.u. per litre.

The development of colour is sensitive to time and temperature of incubation as well as to the purity of the reagents used. It is therefore strongly recommended that where possible AR grade reagents should be used and a standard carried through the colour development with each assay or batch of assays. The use of a standard curve cannot be recommended with this test. A linear relationship exists between colour development and enzyme activity up to approximately 300 m i.u. under the conditions prescribed. Specimens with greater activity should be repeated using either a smaller quantity or shorter period of incubation and the results adjusted accordingly.

Since erythrocytes contain approximately 150 times the phosphoglucose isomerase activity of serum, which is very similar to the ratio for lactate dehydrogenase activity, visible haemolysis makes a sample useless for phosphoglucose isomerase assay (King and Waind, 1961). Heller, Weinstein, West, and Zimmerman (1960) found consistently lower activity in platelet-poor plasma, approximately 60% that of serum. Very variable results were found with plasma, irrespective of the anticoagulant used, and serum is recommended.

For assay of phosphoglucose isomerase activity in cerebrospinal fluid, the reagents are unchanged but the substrate contains 5 mg. glucose-6-phosphate (disodium salt) per millilitre of buffer. Of this buffered substrate 0.6 ml. and 0.5 ml. of cerebrospinal fluid are incubated and the assay continued as for serum. The calculation is the same as previously shown which in this case makes the standard equivalent to 20 m i.u.

NORMAL VALUES

With 100 sera from normal healthy adults, aged from 19 to 60 years, a range of 13.5 to 86 m i.u., with a mean of 46.5 m i.u. and standard deviation ±15.5, was found. Ninety-eight per cent of these values was covered by the range 13.5 to 78 m i.u., and it is suggested that activity outside this range should be considered abnormal (Wootton and King, 1953).

In the first days of life much higher levels are found. Assay of 36 cord blood sera gave values from 45 to 170 m i.u. with a mean of 90 and standard deviation ±33. In this series readings were also taken at 520 mµ which gave a range 45 to 169 m i.u., mean 91 and standard deviation ±32, thereby demonstrating the reproducibility of the method at either wavelength.

With cerebrospinal fluid from patients with no demonstrable central nervous system lesions activity from 0.5 to 6.2 m i.u. has been found and this is proposed as a provisional normal range although it is much lower than that found by Brun's, Jacob, and Weverinck (1956).

RESULTS

A fairly wide range of pH optima extending from pH 7.2 to 8.4 is observed using phosphate, tris, and borate buffers. Higher activity and a more linear relationship between activity and time and activity and enzyme concentration is found with borate buffer, but we are unable to confirm the observations of Alvarado and Sols (1957) that in the presence of M10 borate the reaction equilibrium is shifted in favour of fructose-6-phosphate.

Results for the composition of the equilibrium mixture are in agreement with those of previous authors, that is, approximately 40% isomerization with a fructose-6-phosphate chromogenicity of 61% (Bodansky, 1953) to 25% isomerization with equal fructose and fructose-6-phosphate chromogenicity (Kahana et al., 1960). Zero order kinetics are observed up to 5% isomerization on the latter assumption.

The amount of substrate that may be used is limited by spontaneous non-enzymic isomerization which, with increasing substrate concentration, gives results in very high optical densities for the assay blanks. Under the conditions finally adopted for the time of incubation and quantity of specimen the
optimal substrate concentration resulted in blank values sufficiently low to permit of tests being recorded in the most sensitive range of absorbometric apparatus, that is, the optical density range of 0-9 to 0-8.

The colour reagents of Roe (1934) and of Roe et al. (1949) were investigated and the latter found to give more reproducible results, particularly when colour development was for 15 minutes at 75°C. as recommended by Hers et al. (1953). Typical absorption spectra are shown in Fig. 1 whence it is seen that the maximum difference in optical density between test and blank is obtained at 410 mμ. Furthermore, readings at 410 mμ are more stable with respect to time than at 520 mμ as previously used. At 410 mμ the optical density of test, blank, and standard increase almost equally, about 8% per hour. At 520 mμ, however, the test optical density decreases some 14% per hour while the blank value increases slightly. This behaviour is illustrated in Fig. 2.

These studies result in an assay method which is suitable for use in routine hospital laboratories, requiring neither skill nor equipment beyond that which is accepted as essential, with the possible exception of a 75°C. water bath.

REFERENCES