Automated technique for the estimation of fetal haemoglobin


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SYNOPSIS An automated alkali denaturation technique which measures fetal haemoglobin is described. This method offers greater speed and a lower standard deviation than comparable manual methods.

In 1951 Singer, Chernoff, and Singer devised the 'one-minute alkali denaturation test' for fetal haemoglobin (HbF) estimation. This test was later modified by prior conversion of haemoglobin to cyanmethaemoglobin (Betke, Marti, and Schlicht, 1959). Kristoffersen (1961) suggested that a greater sensitivity would be obtainable if the haemoglobin was estimated at the Soret band instead of at 540 nm. Cabannes and Schmidt-Beurrier (1965) have described an automated technique in which the de-natured adult haemoglobin was filtered off by material inserted into the tubing of the manifold. The automated method described here is based on the test of Singer et al (1951) and utilizes an improved filtration unit which was originally designed to automate the recognition of sickle-cell haemoglobin (Canning, Crane, Huntsman, and Yawson, 1972).

The alkali-resistant fraction of all blood samples was estimated manually by both the Singer and Betke techniques using, in each case, 540 and 420 nm. When the automated technique was used, samples were examined both as a haemolysate and as a rapidly prepared lysed red cell suspension, all tests again being read at 540 and 420 nm.

Material and Methods

SAMPLES
Venous blood was collected into sequestrene bottles from normal healthy individuals (12 samples); B thalassaemia minor patients (15 samples), including parents of children with B thalassaemia major and patients diagnosed on abnormal blood films and raised haemoglobin A₂; patients with sickle-cell haemoglobin C disease (three samples). Standards were made using different mixtures of normal adult and cord bloods to give the requisite Hb F concentrations.

PREPARATION OF SAMPLES
Standard lysates (10 g/100 ml Hb concentration) were prepared (Lehmann and Ager, 1961).

'QUICK' LYSATES
Two ml of blood of known haemoglobin concentration was centrifuged for five min at 3000 rpm (1800 g) in a graduated centrifuge tube. The plasma was removed with a Pasteur pipette, and distilled water was added carefully to the volume calculated to give a final haemoglobin concentration of 10 g/100 ml. All samples were stored in liquid nitrogen after preparation.

MANUAL METHODS
As described by Singer et al (1951) and Betke et al (1959), reading all samples in both methods at 540 and 420 nm, ie, each sample was estimated by four different techniques.

AUTOANALYZER

Reagents
1 0.083 molar sodium hydroxide.
2 Saturated ammonium sulphate containing concentrated (SG 1.18) hydrochloric acid 5 ml per 2 l.

Filter paper reels
Whatman grade 50 width 2.5 cm.

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Apparatus
Technicon Autoanalyzer mark I consisting of (1) Technicon sampler II with 20 per hr 1 to 6 sample to wash ratio; (2) Technicon proportioning pump; (3) Technicon continuous filter but a modification of the standard filtering unit constructed in the manner described by Canning et al (1972); (4) Technicon colorimeter with 15 mm tubular flow-cell and two sets of standard interference filters 420 nm and 540 nm, both of band width ± 9 nm. The Soret band maxima of the filtrates by the Betke and Singer techniques were found to be 416 and 419 nm, respectively; (5) manifold of which the flow diagram is shown in the figure. Standard tygon pump tubing is used. The polypropylene sample tube should be as short as possible. A short length of polypropylene sample tubing was also used for the filtrate line.

EXPERIMENTAL DESIGN

Hand analyses
The five standard samples were analysed six times and the 30 test samples once, each by four different techniques over 10 days, according to a statistical design which followed plan II, 1a (Cochran and Cox, 1957) with days as blocks. Estimates of error variance were calculated on the standards, correcting for timing effects, and were used to derive standard deviations.

Machine analyses
The samples were analysed in five batches: in each run, test samples (both standard and ‘quick’ lysates) and the five standard samples were analysed in duplicate, the batch order being at random. The machine was calibrated using values derived for the standards by Singer hand analysis at the appropriate wavelength.

Results
The ranges and means of results for the HbF concentration obtained in the three groups of samples by manual and automated techniques are shown in

![Flow diagram for the estimation of fetal haemoglobin](http://jcp.bmj.com/)

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**Fig**  Flow diagram for the estimation of fetal haemoglobin
Table I  Ranges and means of results of Hb F estimation

<table>
<thead>
<tr>
<th>Method</th>
<th>420 nm</th>
<th>540 nm</th>
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<tbody>
<tr>
<td>Manual Methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Singer</td>
<td>0.273</td>
<td>0.256</td>
</tr>
<tr>
<td>Betke</td>
<td>0.194</td>
<td>0.199</td>
</tr>
<tr>
<td>Machine Methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Standard Lysate</td>
<td>0.133</td>
<td>0.192</td>
</tr>
<tr>
<td>'Quick' Lysate</td>
<td>0.115</td>
<td>0.288</td>
</tr>
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</table>

Table II  Estimated standard deviations for each of the eight methods

Table III  Correlation coefficients between pairs of estimates of the 30 test samples

Table IV  Correlation coefficients between differences and means of pairs of estimates of the 30 test samples

*Significant values (p < 0.005) in bold type
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The automated technique not only offered greater speed (an output of approximately 20 per hour) but also gave standard deviations lower than those attained by using comparable manual techniques (table II). Machine readings at the longer wavelength (540 nm) tended to rise less rapidly with increase in percentage of HbF than do readings by manual methods; this phenomenon is not seen at 420 nm. The automated technique gave raised fetal haemoglobin levels in only four out of 15 samples of B thalassaemia minor and none of the samples of Hb SC disease. These four raised samples all gave recognizably high HbF levels by the manual techniques. Likewise the 11 samples of B thalassaemia minor and three samples of Hb SC disease with normal results by the automated technique gave manual results within the normal range.

Plasma absorbs light at 420 nm and thus increases the reading given by the alkali-resistant Hb fraction: this is mainly due to the protein and carotenoid content and is not significantly affected by high levels of bilirubin. A rapidly prepared lysate was therefore designed so that, after a single centrifugation, most of the plasma was removed before adjusting the haemoglobin concentration to 10 g/100 ml with distilled water. This preparation is also suitable for cellulose acetate electrophoresis as well as for the automated detection of Hb S (Canning et al., 1972). It was therefore of interest that the automated Hb F technique described here apparently gave more satisfactory results with this preparation than with the time-consuming carefully prepared haemolysate. The results of Hb F obtained with the ‘quick’ lysate method were a mean of 0-31% higher (see table V); this could be allowed for with a correction factor or by accepting a higher normal range.

References


Canning, D. M., Crane, R. S., Huntsman, R. G., and Yawson, G. I.

Table V  Test statistical values (t at 29 df) for differences in pairs of estimates of the 30 test samples

<table>
<thead>
<tr>
<th></th>
<th>a</th>
<th>b</th>
<th>c</th>
<th>d</th>
<th>e</th>
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<td></td>
<td></td>
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<td></td>
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<tr>
<td>Singer</td>
<td>a</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Betke</td>
<td>b</td>
<td>-4·76</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>Machine (standard)</td>
<td>c</td>
<td>-0·52</td>
<td>2·48</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Machine (quick)</td>
<td>d</td>
<td>-5·03</td>
<td>-2·34</td>
<td>-8·02</td>
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<td></td>
</tr>
<tr>
<td>540 nm</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Singer</td>
<td>e</td>
<td>5·40</td>
<td>6·77</td>
<td>2·56</td>
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<tr>
<td>Betke</td>
<td>f</td>
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<td>3·70</td>
<td>-1·08</td>
<td>3·84</td>
<td>-4·70</td>
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</table>

Significant values (*p < 0·005) in bold type

each pair of techniques. These show a high level of agreement.

The extent of this agreement is examined further in tables IV and V. In table IV correlation coefficients of the difference between the two estimates and of the mean of the two are given for each pair of techniques. These will attain significance if the difference between estimates by two methods varies noticeably as the Hb F level increases. This happens only with the machine at 540 nm compared with other methods. An automated technique at this wavelength would not in any case have been chosen because of low sensitivity. In this respect, however, machine methods at 420 nm appear to be good.

In table V results of a statistical test, the paired t test, on the mean difference in estimates by two techniques are given, again for each pair of techniques. The automated technique at 540 nm is excluded because of the variation shown in table IV. The average separation of estimates by two methods is thereby assessed.

Discussion

The values of HbF in the 12 normal samples by the Singer manual method (see table I) at 540 nm correlates with the findings of Singer et al. (1951) of a normal range of 0·5% to 1·7%. The cyanometahemoglobin method (Betke et al., 1959) at this wavelength gave normal values considerably greater than those quoted by these authors who regarded readings above 1·0% as clearly abnormal.

From table V, it can be seen that Singer results are lower than Betke results, at both wavelengths; either hand method gives higher readings at 420 nm compared with the same method at 540 nm. The standard deviations attained by Pembrey, McWade, and Weatherall (1972), who used a modification of the Betke technique, reading at 415 nm, are much lower than those obtained by our manual methods at either wavelength (table II). However, their standard deviation of 0·041 was obtained on 16 analyses of a single sample with a low HbF concentration (mean 0·465%).


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