Glucose-6-Phosphate Dehydrogenase (G6PD) Deficiency, Thalassaemia, and Abnormal Haemoglobins in Taiwan*

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During 1960 blood samples were obtained from 300 male Taiwanese in Taipei, Chi-Chi, and Chung-Li. The majority of donors were normal adolescents and adults whose blood was taken in district health centres and factories. A few hospital patients with diseases necessitating operation were included in Taipei. The 98 from Taipei were urban non-tribal persons and originated in all parts of Taiwan. Chung-Li (centre of island) and Chi-Chi (N.W.) on the other hand are small communities that cannot be regarded as representative of Taiwan as a whole. All the persons whose bloods were examined are descendants of Chinese from Fukien province who had immigrated into Taiwan in the 17th century and later. Venous samples of blood were collected into ACD solution and sent to Seattle by air in ice-cooled vacuum flasks. They arrived within 24–48 hours in excellent condition.

Since Taiwan was until recently an area with a high endemicity of falciparum malaria, the chief interest of this study was to determine the prevalence of G6PD deficiency, thalassaemia, and abnormal haemoglobins. The opportunity was taken of determining blood and serum group frequencies of which relatively little is known in the Taiwanese population. These are reported in a companion paper.

G6PD Deficiency

Nine of the 300 males showed unmistakable evidence by the brilliant cresyl blue decolorization test (Motulsky and Campbell-Kraut, 1961) of G6PD deficiency. This frequency of 0·03 for the sex-linked gene for G6PD deficiency compares with 0·02 found by Vella (1959) among Chinese in Singapore but only 0·003 in another survey in Taiwan (Lee, Shih, Huang, Lin, Blackwell, Blackwell, and Hsia, 1963). A more recent survey (Lee, unpublished) detected 90 deficient individuals among 2,160 males—a gene frequency (0·041) similar to that reported in this paper. In the same study, however, the incidence of G6PD deficiency among recent immigrants from the Chinese mainland to Taiwan was 0·018 and that among Hakka Chinese who came to Taiwan in the 16th and 17th century from Kwangtung and not Fukien province was no less than 0·055.

It is likely that G6PD deficiency is of clinical significance in Taiwan. Among 41 children with acute haemolytic anaemia admitted to the National Taiwan University Hospital in recent years, 37 were boys and 4 were girls. The low ratio of girls is entirely compatible with similar findings in Favism and drug haemolytic anaemias caused by the sex-linked G6PD deficiency in Greece and Sardinia. The aetiological agents in Taiwan were not quite clear but included sulphur drugs in some cases.

In order to define the nature of the gene determining G6PD deficiency in Chinese, a family originating from Canton, in Kwangtong province, and living in Seattle was studied more thoroughly. The proband was an enzyme-deficient boy ascertained during a screening survey among Seattle schoolchildren. He showed an enzyme level of zero when measured by the method of Zinkham, Lenhard, and Childs (1958). Furthermore the decolorization time, using the brilliant cresyl blue test, was more than six hours. Such results are not normally found in Negroes, among whom enzyme-deficient males have a decolorization time of 2–4 hours and 10–15% of G6PD activity on quantitative testing. When these studies were done, electrophoretic and enzymological characterization of G6PD was not practicable. Further studies will have to decide whether the Chinese variant of the
gene for G6PD deficiency like that in the Philippines (Motulsky, Stransky, and Fraser, 1964) and in the Melanesians of New Guinea (Kidson and Gorman, 1962) is identical or different from that seen in Mediterranean (Greek, Italian) rather than Negro subjects. It is of interest that this boy's heterozygous mother and sister had enzyme levels well below the lower limits usually seen in Negro heterozygous females in the same laboratory.

Abnormal Haemoglobins

No major haemoglobin component other than normal haemoglobin A was found by paper electrophoresis of haemolysates. It is of interest that Blackwell and Huang (1963) studied 655 Taiwanese aborigines also without finding any abnormal haemoglobins.

Thalassaemia

Osmotic fragility of red blood cells in 0.45% NaCl was used as a screening test. The results expressed as percentage of lysis followed a normal distribution on the whole. 28 persons, however, with an increased osmotic resistance (percentage of lysis less than 75%) lay outside this main distribution. These were further examined for foetal (F) haemoglobin levels by the alkaline denaturation technique (Singer, Chernoff, and Singer, 1951a, b) and screened for haemoglobin A_2 level by cyano-gum electrophoresis (Raymond and Weintraub, 1959). In 12 of these 28 subjects visual examination of the cyanogum gel suggested a possible increase above normal in haemoglobin A_2 levels. An exact estimate of the proportion of haemoglobin present as A_2 was obtained by elution following starch block electrophoresis in these 12 specimens (Gerald and Diamond, 1958). In 9 instances the presence of high levels of haemoglobin A_2 was confirmed (Table I), and it may be assumed that these represent cases of β thalassaemia trait. In all 9 cases examination of the blood film treated with Wright's stain revealed morphological abnormalities of the erythrocytes typical of this condition. A high level of haemoglobin F (> 2%) was seen in only one of these cases and HbF was not raised in any of the 19 remaining cases with lysis of less than 75%. An additional 100 subjects with normal osmotic fragility were tested for increases of foetal haemoglobin by the alkaline denaturation technique. No abnormalities were found.

Thus an estimate of 3% as the prevalence of β thalassaemia in Taiwan is obtained, but there is a possibility that this represents a minimum figure and that other types of thalassaemia exist among the remaining 18 cases with increased osmotic resistance. Though the usual type of β thalassaemia trait is accompanied by an increased haemoglobin A_2 level, it has been suggested that another type exists in which A_2 levels are normal or even low due to interference with δ as well as β chain synthesis (Fessas, 1963). The screening techniques employed here could not have definitely identified cases of this condition, though the absence of high levels of HbF with which it is usually associated suggests that it is not present in this population.

It seems more likely in view of the work of Lie-Injo, Lie, Ager, and Lehmann (1962) on the prevalence of α thalassaemia among persons of Chinese ancestry that some cases of this condition were not identified. Though paper electrophoresis revealed no examples of haemoglobin H or haemoglobin Bart's, this is not an adequate screening technique (Fessas, 1963). Again, though haemoglobin H inclusion bodies were searched for routinely following incubation with Coleman-Bell brilliant cresyl blue (Gouttas, Fessas, Tsvevrenis, and Xeferi, 1955), the inevitable delay of 48 hours that occurred between taking the blood and this examination much reduced the chances of positive findings even in α thalassaemia trait carriers. It may well be, therefore, that several of the 18 persons with increased osmotic resistance but normal haemoglobin A2 levels were such carriers. However, the extent of iron-deficiency in this male, largely adolescent, and adult population is not known, and anaemia due to this cause could give similar findings.

In view of suggestions that malaria plays a role in the maintenance of G6PD deficiency and thalas-

<table>
<thead>
<tr>
<th>TABLE I</th>
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<tbody>
<tr>
<td>FINDINGS IN 9 SUBJECTS WITH PROBABLY β THALASSAEMIA TRAIT</td>
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<table>
<thead>
<tr>
<th>Age</th>
<th>Osmotic Fragility (%)</th>
<th>HbF%*</th>
<th>HbA2%†</th>
</tr>
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<tbody>
<tr>
<td>21</td>
<td>23</td>
<td>0.4</td>
<td>5.3</td>
</tr>
<tr>
<td>16</td>
<td>67</td>
<td>0.4</td>
<td>3.8</td>
</tr>
<tr>
<td>28</td>
<td>26</td>
<td>0.3</td>
<td>3.5</td>
</tr>
<tr>
<td>24</td>
<td>24</td>
<td>0.3</td>
<td>4.6</td>
</tr>
<tr>
<td>52</td>
<td>69</td>
<td>0.3</td>
<td>4.5</td>
</tr>
<tr>
<td>53</td>
<td>60</td>
<td>1.0</td>
<td>3.3</td>
</tr>
<tr>
<td>33</td>
<td>10</td>
<td>4.0</td>
<td>4.9</td>
</tr>
<tr>
<td>49</td>
<td>30</td>
<td>1.7</td>
<td>5.9</td>
</tr>
<tr>
<td>19</td>
<td>41</td>
<td>1.8</td>
<td>4.5</td>
</tr>
</tbody>
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* HbF levels above 2% are abnormal.
† HbA2 levels above 3.2% are abnormal under these conditions of testing.

Note: In all cases red blood cells showed morphological abnormalities typical of this condition.
thalsassemia by increased resistance of gene carriers, it might be expected that a greater than chance proportion of persons affected simultaneously with both conditions might be found in the population. In fact, of the 9 thalsassemic subjects in Table I, one only is G6PD deficient. Though the expected frequency of such persons is only 1 in 1,000, the observed incidence of 1 in 300 clearly cannot be taken, in the absence of further evidence, to be significant. A similar slight excess was found by Bernini, Carcassi, Latte, Motulsky, Romei, and Siniscalco (1960).

Summary

G6PD deficiency and β-thalassemia trait were found in equal frequencies in a population of 300 Taiwanese Chinese males (approximately 3% of each). No abnormal major haemoglobin components were found.

We should like to acknowledge the expert technical assistance of Mrs Jackie Keil.

REFERENCES


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