Abstract
The frequencies of four malaria associated erythrocyte genetic abnormalities have been established in 1000 Omani subjects. They are: homozygous α-thalassaemia (−α/−α) 0.45; high Haemoglobin A, β thalassaemia trait 0.015; sickle trait (Hb A/S) 0.061; and glucose 6 phosphate dehydrogenase deficiency (Gd−): males 0.27, females 0.11.

From our data the α− (−α/) thal gene (confirmed by Southern blotting) is pandemic in this population. Moreover, in spite of the very high frequency of Gd−, oxidative haemolytic syndromes are very uncommon. Also preliminary data indicate that among the Omani population with sickle cell disease, homozygosity of the α− gene markedly modifies the clinical picture.

Methods
The routine blood counts were measured on a Coulter S Plus VI cell analyser. Haemoglobin A2 was measured using Helena Beta-thal and Sickle-thal ion exchange chromatography columns. The normal range for the Omani phenotype Hb AB/AB for this laboratory was 1.8 to 3.5%. Values greater than 4.5% were accepted as being diagnostic of high Haemoglobin A2, β thalassaemia trait.

Blood was screened for haemoglobin S using Ortho Sicklelex kits. Haemoglobin electrophoresis was performed on all positive samples on agar gel at pH 8.6 and pH 6.0 using the Beckman Paragon system.

Cord bloods were examined for the presence of Haemoglobin A, Haemoglobin S, and Haemoglobin Barten's by electrophoresis on agar gel at pH 8.6 and pH 6.0. The Haemoglobin Barten's was quantitated by scanning densitometry.

Haemoglobin F levels were measured by high performance liquid chromatography (HPLC) using the Biorad 'Diamat' instrument. The normal range for Omanis was <1%. Glucose 6 phosphate dehydrogenase (G6PD) activity was estimated using the Boehringer screening and quantitative kits. On deficient samples the activity was quantitated. Values less than 10 mU/10⁶ RBCs were accepted as being deficient. The quoted normal range for this method was 131 ± 13 mU/10⁶ RBCs.

The iron status of the patients was determined by measuring the serum ferritin levels using the Abbott IMX microparticle enzyme immunoassay method. A value of less than 15 ng/ml was indicative of iron deficiency.

Haemoglobin gene analysis was carried out as follows: α globin gene analysis on a subset of eight adult Omani subjects with haematological and biochemical parameters indicative of α thalassaemia was carried out by blot hybridisation using standard methods. The probes

Subjects
Nine hundred and fifty-two Omani subjects, 435 males and 517 females, identified by tribal names, were studied. These comprised hospital patients (73%) and university students (27%). Of the hospital patients 83% were adults and the remainder children. To determine the frequencies of Haemoglobin S, Gd−, and β thalassaemia the whole population was studied. However, with regard to α thalassaemia only the male student population was studied in detail and the data extrapolated to predict the frequency in the population as a whole.

One hundred and forty-two ‘non-Arab’ subjects, 82 males and 60 females, were healthy blood donors, were used as a control population for comparing the haematological parameters. They came from North America, Great Britain, Northern Europe, India, and the Philippines.

Frequency and clinical significance of erythrocyte genetic abnormalities in Omanis

J M White, B S Christie, D Nam, S Daar, D R Higgs
used in this study were $\alpha$-globin/HBA1$^{a}$ and $\Psi$C1 globin/HBZP.$^{6}$ In addition the $\beta$ globin genes of 29 patients with suspected homozygous $\beta$ thalassaemia were analysed using the amplified refractory mutation system.$^{7}$

**Results**

The distribution of the haemoglobin values of the 952 subjects is shown in fig 1. The curve was Gaussian but the geometric mean (GM) of 12.9 (SD 2.95) g/dl was significantly lower than that of our non-Arab population, 14.2 (SD 2.6) g/dl ($p=0.001$) (table 1). The main factor responsible for this ‘anaemia’ was the low MCH values of Omani, GM = 24.8 (SD 6.5) pg, when compared to the western value 30.4 (SD 3.24) pg. That the cause of this hypochromia was largely the result of thalassaemia genes was indicated by the higher red cell counts of the Omani population 5.2 (SD 1.52) $\times 10^{12}$/l compared to the non-Arab value of 4.7 (SD 1.2) $\times 10^{12}$/l ($p=0.001$), and the strong negative correlation between the MCH and the RBC ($r=-0.6$). Moreover, the distribution of the MCH showed two significantly distinct populations (fig 2). The two groups were of nearly equal size. Group A consisted of approximately 50% of the subjects in whom the MCH ranged from 14 to 25 pg (GM = 22.8 (SD 2.6 pg)). Group B consisted of the other 50% of subjects with an MCH between 25 and 32 pg (GM 27.3 (SD 4.3) pg). The difference between the geometric means of A and B was highly significant ($p<0.001$). The bimodal pattern of the MCH was constant to each subgroup of subjects when the whole population was subdivided into males or females, adults or children, and hospital patients or students. This is illustrated in fig 2 where only the male subgroup populations are shown as an example. It is assumed that the cause of the severe hypochromia in group A is $\beta$ or $\alpha$ thalassaemia genes or iron deficiency. The cause of the hypochromia of the less severe group B is $\alpha$ thalassaemia or iron deficiency.

**HIGH Hb A, $\beta$ THALASSAEMIA**

The levels of Hb A were measured on 400 blood samples from the total population of 952 who had an MCH of less than 24 pg (group A). It was increased in 14 with values ranging from 4.5 to 5.6%. This gives an overall frequency of high Hb A, $\beta$ thalassaemia genes in the total population of 0.015.

**$\alpha$ THALASSAEMIA**

Because there is no simple diagnostic marker for $\alpha$ thalassaemia, and it is impractical to carry out DNA analysis on 1000 people, it was decided to select eight samples at random from patients whose haematological indices indicated $\alpha$ thalassaemia trait; namely an MCH of $<25$ pg, a normal Hb A$_2$, and a normal ferritin. These were then sent blind to another laboratory for $\alpha$ gene analysis. The results showed that all eight patients were homozygous for the $\alpha^{-}$ deletional ($-\alpha$) thalassaemia gene (table 2).
Once confirmed, the frequency of the gene in Omani could now be estimated with some degree of certainty by only measuring the MCH, Hb A2, and ferritin levels. For this we examined 127 of the male student population.

The MCH distribution pattern of this group is shown in fig 2. There were 63 students with a severe hypochromia with an MCH of 20 to 25 pg. Of these, 58 had normal ferritins of greater than 25 ng/ml and only one had a raised Hb A2. It was therefore assumed that the only cause of the hypochromia of the remaining 57 was homozygosity for \( \alpha^+ \) thalassaemia (\( \alpha/\alpha \)) which gives a frequency among the male students of 57/127 = 0.448. Since the relative size and MCH distribution of this student group ran true throughout the whole population it seems realistic to assume that the frequency of this type of \( \alpha \) thalassaemia is also true for the whole population.

To test this further the frequency was calculated from the number of cord bloods which contained 3 to 5% Hb Bart’s (\( \gamma_\lambda \)) and also from the levels of Hb S in A/S carriers.

Of the 247 cord bloods analysed, 104 (42%) contained visible (3 to 5%) Hb Bart’s (fig 3). This is well recognised to be associated with a two deletional type of \( \alpha \) thalassaemia. This value gives a frequency of two deletional \( \alpha \) thalassaemia in the population of 0.42. Finally, among the 93 sickle cell trait patients, 38 had levels of Hb S less than 30% (fig 4). Again such levels are thought to be the result of association with two \( \alpha \) gene deletional \( \alpha \) thalassaemia and this value would give a frequency of 0.41.

The major cause of the hypochromia in group B is also probably the result of a milder form of \( \alpha \) thalassaemia most likely \( \alpha/\alpha \alpha \) since only three of the 64 students in group B had low ferritin levels and all had normal Hb A2 levels. From the frequency of the homozygosity of the \( \alpha \) haplotype of 45% one can compute that the frequency of the \( \alpha/\alpha \alpha \) genotype is 0.44 and \( \alpha/\alpha \alpha \) in this population. It is worth stressing that the frequency of the \( \alpha \) haplotype is remarkably high at 0.67.

SICKLE Hb

Of the 952 samples tested, 58 gave a positive sickling test (6.1%) and all were shown to be heterozygotes for Hb A and Hb S. This gives an Hb S frequency of 0.06. Of the 247 cord bloods analysed by acid electrophoresis, 15 were found to contain sickle haemoglobin (Hb F, A, and S) which would give a frequency of 0.060. If correct, there should be 37 homozygotes per 1000 live births and approximately 5000 to 6000 in the country.

Of interest is the distribution of Hb S in heterozygotes which is shown in fig 4, where the percentage of Hb S (x axis) is correlated with the patients’ MCH (y axis). Two distinct but equal populations are seen. None of these carriers was iron deficient. Therefore the strong correlation with the MCH (\( r=0.98 \)) would indicate that the amount of Hb S found is mainly a function of red cell hypochromia and in the Omani population is largely dictated by the number of functional \( \alpha \) chain genes.

S-GLUT

Among the 435 males examined, 119 (27.3%) were found to be G6PD deficient on screening and in all these the activity of the enzyme was less than 10 mU/10¹⁰ RBCs. Of the 517 females tested, 64 were deficient; 57 of these (11.0%) had values of less than 10 mU/10¹⁰ RBCs and
are thus probably homozygous for the deficiency. These data give a frequency for male hemizygotes of 0.27 and for female homozygotes of 0.11. The DNA from a subset of samples from deficient subjects is now being examined by restriction enzyme analysis (S Daar, L Luzzatto, unpublished data).

**Discussion**

As stated, Oman is a nation of some 1.5 million people and is geographically isolated on the south-east Arabian Peninsula by the high western mountain range and the south-eastern desert quarter of Saudi Arabia. Falciparum malaria is endemic in this country with 27,000 cases notified to the Ministry of Health in 1991. It is therefore not surprising that malaria associated erythrocyte genetic defects are found. What is surprising is the high frequency they have reached.

The findings from this study support the early suspicion that the frequency of \( \alpha \) thalassaemia and \( \text{Gd}^- \) were the highest of the Peninsular Arab races, and indeed are among the highest in any race so far reported. The frequencies are summarised in table 3. There are three aspects of these abnormal genetics which merit further discussion.

**HIGH Hb A, \( \beta \) THALASSAEMIA GENE**

The frequency of 0.015 is relatively low but in keeping with the frequencies found in Saudis, Yemenis, and UAE nationals. Using this frequency it was calculated that there would be 300 homozygotes among Oman’s population of 1.5 million and indeed 137 patients are registered at this hospital. However, after examining the tribal names of homozygote patients, 75% were found to be non-Arab and had originated from Beluchistan two centuries ago. This was supported by DNA analysis of the \( \beta \) globin genes of 29 patients, 19 of whom (70%) showed the common Indian allele IVS1-5 G\( \rightarrow \)C. This means that the true frequency for the Omani Arab is much lower and of the order of 0.0038.

**\( \alpha \) THALASSAEMIA**

We conclude from the data derived from examining the male student population that the \( \alpha \) thalassaemia gene is endemic among the Omani and that it is the result of the presence of the \( \alpha^+ \) (\(-\alpha/\)) gene. This is not only based on the DNA analysis but is supported by the fact that non-mutational HbH disease (\(-/-\)) and Bart’s hydrops fetalis (\(-/-\)) have to our knowledge never been reported in Peninsular Arabs. From the red cell parameters 45% of the population are homozygous for this gene (\(-\alpha/-\)). This is supported by the frequency of Hb Bart’s in cord bloods and the levels of Hb S in carriers.

Furthermore, it is likely that the major cause of the hypochromia among the other half of the population, with MCH values between 25 to 28 pg, is heterozygosity for the \( \alpha^+ \) gene (\(-\alpha/\alpha\)). If correct, it means that the \(-\alpha/\) haplotype is pandemic. The very high frequency is probably because of Oman’s geographical isolation. However, it also implies that this genetic defect is harmless and, moreover, it probably affords some biological advantage. There is no evidence that a single \( \alpha \) gene deletion affords protection against malaria. Of clinical importance is that \( \alpha/\beta \) thalassaemia must be common among our 117 homozygous \( \beta \) thalassaemia patients who are already in a hypertransfusion programme. So far five out of 10 of our patients have now been shown by DNA analysis to be heterozygous for \( \alpha^+ \) and homozygous for \( \beta^+ \) thalassaemia genes (\(-\alpha/-\alpha \beta^+/-\beta^+\)) and may well have shown a thalassaemia intermedia-like picture. Therefore, where \( \alpha \) thalassaemia is endemic complete \( \alpha \) and \( \beta \) globin gene analysis is essential when these patients first present before a long term hypertransfusion policy is adopted. We have found that to discontinue transfusion after years is clinically difficult because of cardiac siderosis and decompensation, and psychologically disturbing for the patients and parents.

**GLUCOSE 6 PHOSPHATE DEHYDROGENASE DEFICIENCY**

The frequency of \( \text{Gd}^- \) in males (0.27) and in females (0.11) is the second highest reported. From these frequencies it is calculated that there are nearly 400,000 people who are deficient in this country. However, as stated previously, clinically there are relatively few oxidative haemolytic crises.
Varieties of fava bean are grown and commonly eaten but very few patients suffer haemolysis. However, there are two other quantitatively more reliable stress factors, namely the use of antimalarials and oxidative antibiotics. In Oman in 1990/91 there were 27 000 cases reported of falciparum malaria which it is presumed were treated. Moreover, during 1991/92, 78 cases of falciparum malaria were admitted to this hospital and all received full doses of chloroquine therapy. Of these patients, 36 were G6PD deficient and none of them suffered from haemolysis. Similarly according to the country’s main drug distributor, 34 000 patient doses of Septrin® were issued in 1990/91. From our frequencies of the ‘at risk population’, approximately one fifth of the patients who received antimalarials or Septrin® would have been G6PD deficient, that is, 12 200 patients. In spite of this, the University Hospital has only admitted five cases of oxidative haemolysis during 1990 to 1992 from a catchment area of one quarter of the country’s population and in three of these patients there were exceptional circumstances. One had severe chronic renal failure and had been given Septrin® and chloroquine. Another was given simultaneously chloroquine, primaquine, and Septrin®. The third was a case of leprosy given high dose Dapsone®. The other two patients were mango growers living in the same area and had been harvesting the fruit a few days before admission, but the cause of their haemolysis remains obscure. However, it is clear that oxidative haemolysis is rare. This has been recently reviewed and stressed by Beutler.10

SICKLE CELL ANAEMIA
Finally the very high frequency of α thalassaemia has been found to modify the clinical and haematological picture markedly in patients homozygous for the sickle gene. At present 177 patients homozygous for α are registered at the hospital. The haematological data are summarised in table 4. Twelve percent (12%) have a normal MCH and presumably a normal complement of α genes, 28% had MCH values of 27 to 25 pg and are probably −/αα, and 60% had an MCH of 25 to 20 pg and are probably −α/α. Among this group the most obvious anomaly is that the 1:1 ratio of −/αα:−α/α seen in the overall population (and the Hb A/S ratio) has now fallen to 0.5:1 (table 4). There is no clear reason for this. The Hb levels and Hb F levels are the same in both groups. The only difference is that the mean age of the −/αα group is somewhat lower but this is not significant. One could postulate either early death of the −/αα patients or that this genotype is much more benign than −α/−α and that they do not attend hospital and therefore remain undiagnosed. However, it may be because the determination of the MCH of the red cells in sickle cell disease using automated machines is unreliable owing to poor lysis of sickled cells. The answer will only come when the α gene status of each patient is determined.

The clinical picture that is emerging in the 103 patients with an MCH of < 25 pg (−/α−/α) is that they have an increased frequency of painful bone crises especially lower vertebral (average 6:3/patient/year) than the non −α thalassaemia β homozygotes (2:1/patient/year). Moreover, persistent splenomegaly is usual into adolescence, by which time all are hypersplenmic, with low white cell and platelet counts. Twenty-six of our patients have had to be splenectomised since they required regular transfusion to maintain their haemoglobin level above 6 g/dl. The lack of splenic inflation may be because of a combination of the lower MCV of the sickle cells and the differences in the anatomy of splenic microcirculation as compared with that of the bones. However, the gross splenomegaly found in a proportion of our patients might be the result of chronic malaria.

Finally it must be stressed that of these 103 patients only five are aged over 20 years and only one over 30. Early death therefore would appear to be common. Although this may be the result of environmental factors rather than a cellular event, it is clear that loss of two α genes does not ameliorate the sickle cell disease.

We would like to thank Professor Lucio Luzzatto for his helpful comments and Mr David Gravel for providing some of the data.

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Table 4  The mean of some haematological values of 177 Hb S/S patients.

<table>
<thead>
<tr>
<th>Probable genotype</th>
<th>No.</th>
<th>Age (y)</th>
<th>Hb (g/dl)</th>
<th>MCH (pg)</th>
<th>RBC (10^12/l)</th>
<th>MCV (fl)</th>
<th>Retic (%)</th>
<th>Hb Aα (%)</th>
<th>Hb F (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>α2/α2</td>
<td>24</td>
<td>13-1</td>
<td>81</td>
<td>29.7</td>
<td>2.64</td>
<td>88.6</td>
<td>19.2</td>
<td>2.99</td>
<td>9.4</td>
</tr>
<tr>
<td>−α/α</td>
<td>50</td>
<td>9-9</td>
<td>8.2</td>
<td>26.5</td>
<td>3.14</td>
<td>81.7</td>
<td>13.9</td>
<td>2.97</td>
<td>12.0</td>
</tr>
<tr>
<td>α2/−α</td>
<td>103</td>
<td>11-0</td>
<td>8.7</td>
<td>21.8</td>
<td>3.99</td>
<td>68.8</td>
<td>10.5</td>
<td>3.39</td>
<td>12.2</td>
</tr>
</tbody>
</table>

A comparison of the means of some haematological parameters among 177 patients homozygous for the β gene. This was confirmed by screening of parents. The three groups have been separated according to the MCH: MCH > 28 pg, 27-25 pg, and < 25 pg. The α gene complement shown is speculative. See text for discussion.

The significant differences seen are the decreases in the ratio of −α/αα:−α/−α from 1:1 in the normal Omani population to 0.5:1. Also, as assessed by the reticulocyte count the rate of haemolysis would appear to be less as the MCH falls but the numbers are too few at present to provide reliable statistical data.

* The higher Hb Aα value is probably because when δ chains are rate limiting the αδ affinity is greater than αβ (to be published).