Review Article


Sickle cell disease in Middle East Arab countries

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The sickle cell (HbS) gene occurs at a variable frequency in the Middle Eastern Arab countries, with characteristic distribution patterns and representing an overall picture of blood genetic disorders in the region. The origin of the gene has been debated, but studies using β-globin gene haplotypes have ascertained that there were multiple origins for HbS. In some regions the HbS gene is common and exhibits polymorphism, while the reverse is true in others. A common causative factor for the high prevalence and maintenance of HbS and thalassaemia genes is malaria endemicity. The HbS gene also co-exists with other haemoglobin variants and thalassaemia genes and the resulting clinical state is referred to as sickle cell disease (SCD). In the Middle Eastern Arab countries, the clinical picture of SCD expresses two distinct forms, the benign and the severe forms, which are related to two distinct β-globin gene haplotypes. These are referred to as the Saudi-Indian and the Benin haplotypes, respectively. In a majority of the Middle Eastern Arab countries the HbS is linked to the Saudi-Indian haplotype, while in others it is linked to the Benin haplotype. This review outlines the frequency, distribution, clinical feature, management and prevention as well as gene interactions of the HbS genes with other haemoglobin disorders in the Middle Eastern Arab countries.

Key words Malaria endemicity - Middle Eastern countries - sickle cell anaemia - sickle cell disease - The Arabs

The Middle Eastern Arab community features and genetic disorders

Of particular interest in the Middle East Arabs are a set of common factors that include the rapid increase in the population and rich historical, cultural, traditional and religious commonality. The large family size, high rate of consanguinity in conjunction with tribe/clan endogamy, make the Arabs unique from the point of view of genetic analysis. Over the years, the Arabs in the Middle East have undergone a considerable transition as regards the health status of its people. Infectious diseases and nutritional disorders have decreased in prevalence as a result of the significant advances made in immunization, the discovery of antibiotics and the overall improvement in hygiene. Thus, these earlier causes of morbidity and mortality are now being exceeded by genetic diseases, which although relatively infrequent, constitute a significant cause of
chronic health problems, morbidity and mortality and hence are a major burden on health care systems.

In the industrialized countries, community surveys show that approximately 3 per cent of all pregnancies result in the birth of a child with a significant genetic disease or birth defect which can cause mental retardation, other crippling conditions or early death. Though data on genetic and congenital defects are not handy in the Arab communities, but considering the high rate of consanguinity and other relevant factors, it is predicted that these disorders are more frequent in this population. Genetic diseases due to their chronic nature impose heavy medical, financial and emotional burdens. Therefore, the efforts to combat these problems are multifaceted and the effective control and prevention strategies gain a high priority beside care and rehabilitation of the affected in the community.

Haemoglobin disorders as genetic diseases

Normal haemoglobins are of different types in human and include Hb A, Hb A₂, and Hb F. Each type of haemoglobin is a tetramer of two different globin chains, each having its own gene. The Hb A (2α2β) is almost 95-97 per cent, Hb A₂ (2α2δ) is 2.5-3.5 per cent and Hb F (2α2γ) is <1 per cent in adults. The α-globin gene cluster is located on the chromosome 16 and includes 5'-ζ-ψα-α2-α1-3', while the non-α globin gene cluster which includes 5'-ε-Gγ-Aγ-ψβ-δ-β-3' genes, is located on the chromosome 11. The expression of α1 and α2 globin genes located on chromosome 16pter-p13.3 and the β globin gene located on chromosome 11p15.5, provide α and β globin polypeptides, and the co-ordinated production of haem, the non-protein portion of Hb chains, results in the formation of HbA, in normal individuals. An A to T transversion mutation at the sixth codon of the β globin gene produces HbS, with a substitution of glutamic acid by valine at the 6th amino acid position in the β globin polypeptide. Individuals homozygous to HbS gene have only HbS in place of Hb A, with concomitant production of Hb F and Hb A₂. In double heterozygotes, the Hbs co-exists with either other abnormal haemoglobin or with thalassaemias. These groups of disorders are together referred to as sickle cell disease (SCD). Majority of the haemoglobin variants other than Hbs, HbC, HbE and HbD are rare, and therefore, rarely give rise to homozygote states. However, thalassaemias, on their own occur more frequently giving rise to homozygous disease conditions.

Pattern of inheritance of haemoglobin disorders

The abnormal haemoglobins and the thalassaemias are inherited as autosomal recessive (AR) disorders, where carrier parents transmit the abnormal genes to the offspring. If both parents are heterozygotes for HbS, there is a 25 per cent chance of having a homozygous HbSS (Sickle cell anaemia, SCA) child. If one parent is a carrier for HbS and the other is carrier for one of the abnormal HbS or thalassaemias, it results in a double heterozygote state. Heterozygotes are generally asymptomatic carriers (traits), while the SCD is expressed in the homozygotes and the double heterozygotes for two abnormal haemoglobin genes or HbS and the thalassaemias.

Pathophysiology of sickle cell disease

The Hb S is soluble in the oxygenated state, as that encountered in the lungs, but once the haemoglobin delivers the oxygen to the tissues, the HbS in the deoxygenated form undergoes a major conformational change, which leads to the formation of long fibrous aggregates (polymers) due to hydrophobic interactions between the valines in the adjacent HbS molecules. These polymers in the erythrocyte, distort its shape from normal spherical biconcave disc to the characteristic sickle shape, leading to erythrocyte rigidity and vaso-occlusion and sickled red cells are formed in the tissues. The haemoglobin polymerisation is central mechanism to the pathophysiology of SCD. Constant sickling and desickling in the tissues and the lungs respectively, increase the fragility of the red cells leading to haemolysis and hence chronic anaemia. Vaso-occlusion results from blockage of the blood vessels by the rigid sickled red cells, leading to the development of painful crises, hand-foot syndrome, inflammation, cerebrovascular disease and cognitive impairment. Recurrent episodes of vaso-occlusion and inflammation lead to vasculopathy which further results in progressive damage to most organs, including the brain, kidneys, lungs, bones, and cardiovascular system obstructs microcirculation, and causes tissue infarction. These frequently result in hand-foot syndrome in children, fatigue, paleness, and shortness of breath, pain that occurs unpredictably in any body organ or joint, eye problems, yellowing of skin and eyes, delayed growth and puberty in children. In addition, infections, stroke, and acute chest pain are some of the major complications. These complications start in early life, but become more apparent with increasing age. Several factors such as infections, dehydration, fever,
cold weather and stress precipitate the complications. Most of the treatments are directed towards prevention
of or decreasing sickling and hence reduction in the vasculopathy and clinical complications of SCD4-8.

**Origin of sickle cell gene**

Studies on haplotypes generated using restriction endonuclease, associated with HbS have confirmed that the HbS mutation occurred as several independent events in Central Africa, Central West Africa, African West coast, Arabian Peninsula and India. In Africa the HbS gene is associated with at least three haplotypes representing independent mutations. These are the Benin haplotype, the Senegal haplotype in the Central African Republic or the Bantu haplotype found in the Central West Africa, the African West coast and the Central Africa (Bantu speaking Africa), respectively. A fourth haplotype, the Saudi-Asian haplotype, is found in the eastern province of Saudi Arabia and central India. Though the origin of HbS was mainly in Africa and Asia, as a result of population movement it spread to different areas of the World and became established in areas which were endemic to malaria. This is due to the natural resistance against development of malaria, in the HbS carriers. At present, HbS has been reported from several countries of the world and the frequency is high in areas with past or present history of malaria endemcity9,10.

**Haemoglobin disorders - occurrence and distribution**

The disorders resulting from inheritance of HbS gene are among the most frequently encountered group of disorders in several populations of the World, in particular among the sub-Saharan Africa; Middle Eastern populations; other Mediterranean countries such as Northern Greece, Sicily and Southern Italy; Spanish-speaking regions (South America, Cuba, Central America), Southern Turkey and much of Central India. Studies have confirmed that the HbS mutation is a relatively recent occurrence, which has occurred independently in several different populations and the presence of falciparum malaria has served as a selective factor in increasing its prevalence11. This is the consequence of the inborn resistance to the development of malaria, which arises in the HbS heterozygotes (carriers), who are less likely to die from malaria and so more likely to survive and pass on their genes, thus playing an important role in maintaining HbS gene frequency. Over the generations, the HbS gene has reached high frequencies in regions with past or present history of malaria endemcity. However, population migration has played a major role in distributing HbS gene even to non malaria endemic regions. Also several hundred mutations affect the globin genes, but only a few occur at a polymorphic level, and majority of the abnormal haemoglobins (Hbs) occur as rare variants, confined to specific ethnic groups or families12.

**Epidemiology of sickle cell gene**

The SCD is most common among people from Africa, India, the Caribbean, the Middle East, and the Mediterranean. In the Middle Eastern countries, the first documentation of abnormal Hbs (HbS) and thalassaemias came from Egypt13,14. Lehmann reported the presence of HbS in Eastern Saudi Arabia15. Extensive studies on different haemoglobin disorders have been reported from almost all the countries of the Middle East, though at a considerably variable frequency. Table I presents brief historical aspects related to identification of abnormal haemoglobins in the Middle Eastern population, and different abnormal variants that have been identified

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**Table I. Haemoglobinopathies in the Middle East Arab countries - Historical aspects**

<table>
<thead>
<tr>
<th>Discovery/country</th>
<th>Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>First case of SCD in Egypt</td>
<td>1951</td>
</tr>
<tr>
<td>HbS in Middle East</td>
<td>1959</td>
</tr>
<tr>
<td>Hb O-Arab in Egyptian family</td>
<td>1960</td>
</tr>
<tr>
<td>HbS in Saudi Arabia</td>
<td>1963</td>
</tr>
<tr>
<td>HbS and Hb O-Arab in Sudan</td>
<td>1966</td>
</tr>
<tr>
<td>Hb C in Egyptians</td>
<td>1967</td>
</tr>
<tr>
<td>Mild SCD in Saudi Arabia</td>
<td>1969</td>
</tr>
<tr>
<td>SCD in Kuwait</td>
<td>1969</td>
</tr>
<tr>
<td>Hb H disease in Kuwait</td>
<td>1969</td>
</tr>
<tr>
<td>HbS in Egyptian western desert</td>
<td>1974</td>
</tr>
<tr>
<td>Hb C in Libya</td>
<td>1975</td>
</tr>
<tr>
<td>HbS in Abu Dhabi</td>
<td>1980</td>
</tr>
<tr>
<td>Hb C in Saudi Arabia</td>
<td>1979</td>
</tr>
<tr>
<td>Hb E and Hb D in Abu Dhabi</td>
<td>1979</td>
</tr>
<tr>
<td>Hb O-Arab in Saudi Arabia</td>
<td>1980</td>
</tr>
<tr>
<td>Hbs, α- and β-thal in several regions of Saudi Arabia</td>
<td>1967-1982</td>
</tr>
<tr>
<td>Extensive studies in the Middle Eastern countries on distribution, frequency, natural history, histopathology, complications and management of haemoglobinopathies</td>
<td>To date</td>
</tr>
<tr>
<td>Preventive programmes, including pre-marital screening</td>
<td>To date</td>
</tr>
</tbody>
</table>

Source: Modified from Ref. 16
are listed in Table II. HbS is the major variant identified in all areas. Table III presents the range of HbS gene frequencies reported from the different Middle Eastern countries. Each country has characteristic distribution and clinical presentation of SCD.

**Frequency and distribution of sickle cell gene among Arabs**

Geographically, Middle Eastern Arabs can be looked at as follows: (i) the Arabian peninsula occupying the South West of Asia includes the Yemen, Saudi Arabia and other members of Gulf Co-operation Council, Kuwait, Qatar, Bahrain, United Arab Emirates and Oman; (ii) the Northern region of Arabian Peninsula that occupies the North West of Asia and includes Palestine, Jordan, Syria, Lebanon and Iraq; and (iii) the Arab countries of North Africa, that include Egypt, Libya, Tunis, Algeria and Morocco.

(i) The Middle Eastern Arab countries of Western Asia

**Yemen:** In the study of White and coworkers, the frequency of SCD in Yemen was reported as 0.95 per cent. Disease course and severity were similar to that in Africans and American blacks and from western Saudi Arabia. In the individuals with SCA, the prevalence of Xmn I polymorphic sites was reported to be similar to the prevalence reported in the south-western region of Saudi Arabia and α-gene deletion occurred at a higher prevalence in patients with Yemeni SCD patients.

**Saudi Arabia:** Sickle cell gene was first recognized in Saudi Arabia in 1963 by Lehmann and co-workers in the eastern province of the country. Gelpi reported the presence of HbS gene in the oasis population of Al-Qateef and Al-Hasa. A mild form of SCA was recognized in this part of Saudi Arabia. Studies conducted in different regions of Saudi Arabia (during 1970s to 1990’s) revealed the presence of HbS and other red cell genetic defects in several regions of the country. Three major foci for HbS gene were identified in the country, and the frequency was found to correlate with the history of malaria endemicity. A comprehensive National screening programme initiated in 1982, covered 36 different areas, provided detailed mapping and distribution of HbS gene and revealed variation in the frequency in different areas of the country. Extensive studies were conducted to trace the natural history of the SCD, and two major forms of the disease were identified, with symptoms ranging from mild to severe. Significant

<table>
<thead>
<tr>
<th>Country</th>
<th>Abnormal haemoglobins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Algeria, Tunisia</td>
<td>HbS, Hb C, Hb O-Arab, Hb J-Mexico, Hb Setif, Hb D-ouled Rabah, Hb G-Saskaton, Hb G-Philadelphia, Hb F-Sarchian, Hb Bab-Saadoun, Hb Boumerdes, Hb J Mexico</td>
</tr>
<tr>
<td>Sudan</td>
<td>HbS, Hb-Khartoum, Hb O-Arab, Hb C</td>
</tr>
<tr>
<td>Lebanon, Syria, Jordan</td>
<td>HbS, Hb C, Hb-Torino</td>
</tr>
<tr>
<td>Iraq</td>
<td>HbS, Hb C</td>
</tr>
<tr>
<td>Libya</td>
<td>HbS, Hb C, Hb J Benghazi</td>
</tr>
<tr>
<td>Egypt</td>
<td>HbS, Hb C, Hb O-Arab, Hb-Khartoum, Hb J-Cairo, Hb E,</td>
</tr>
<tr>
<td>Saudi Arabia</td>
<td>HbS, Hb C, Hb O-Arab, Hb D-Punjab, Hb-Riyadh, Hb F-Dammam, Hb E, Hb-Handsworth, Hb-Setif</td>
</tr>
<tr>
<td>UAE</td>
<td>HbS, Hb E, Hb C, Hb D-Punjab, Hb O-Arab, Hb G-Audhali, HbA1-Ain Abu dhabi</td>
</tr>
</tbody>
</table>

Source: Modified from Ref. 16
differences were observed in the HbF level in different patients. HbS gene was frequently shown to coexist with other abnormal Hbs, thalassaemias and glucose-6-phosphate dehydrogenase (G-6-PD) deficiency. Studies on associated β-globin gene haplotypes revealed the presence of the Saudi-Indian haplotype in majority of the SCD patients from the Eastern province with a mild form of the disease and the Benin haplotype in majority of the patients from the Western province with a severe form of the disease. Different treatment protocols were adopted and hydroxyurea and paracetam were shown to be beneficial for the treatment of SCD in majority of the patients. Control and prevention programs have been implemented and steps have been adopted to increase awareness about these frequent disorders.

**Bahrain:** Several studies have been carried out on the SCD in Bahrain. In a study conducted during the 1986 on Bahrani girls, the incidence of sickle cell haemoglobin (Hb AS), was approximately 7 per cent. In a group of 100 consecutive pregnant females, the frequency of HbSC was 17.5 per cent, and Hb AS was 32.5 per cent. Patients with SCD were shown to have elevated Hb F levels, and double heterozygous HbS/β-thalassaemia cases were also identified. In a large study on hospital population of Bahrain, which included 5,503 neonates and 50,695 non-neonates, the prevalence of SCD was reported as 2.1 per cent and HbS trait as 18.1 per cent in the neonates, and 10.44 per cent SCD in non-neonatal patients. Al-Arrayed and coworkers conducted a screening of student for inherited blood disorders in Bahrain and reported the prevalence as 1.2 per cent SCD; 13.8 per cent Hb AS; 0.09 per cent beta-thalassaemia; 2.9 per cent β-thalassaemia trait. The majority (84%) of the SCD patients had elevated HbF. The Saudi-Indian haplotype was the major haplotype (90%) in the Bahrani SCD patients. HbS was also reported to occur with other abnormal haemoglobins (e.g. HbS/D) in the population of Bahrain, and a high percentage of the SCD patients (47%) had associated G-6-PD deficiency.

**Qatar:** In 1985, Bakioglu and coworkers conducted screening of the Qatari population and reported the presence of HbS gene. In another screening study on 1,702 Qatari nationals, it was shown that 7.46 and 3.94 per cent were Hb AS and SCD patients, while 1.53 and 3.23 per cent of the patients had HbS/β-thalassaemia and HbS/α-thalassaemia, respectively. The SCD was reported as mild with elevated HbF level though some patients suffered from episodes of crises. Haplotype analysis also revealed the presence of Saudi-Indian haplotype.

**Kuwait:** In 1970, Ali reported sickle cell with a milder variant of the disease in Kuwaiti and showed that this was associated with unusually high levels of Hb F. Molecular characterization of β revealed the presence of Asian (Saudi-Indian) haplotype in 77.8 per cent and Benin haplotype in 16.7 per cent of the chromosomes. Marouf et al conducted a comprehensive electrophoretic screening of the Kuwaiti population and showed that 23.5 per cent had abnormal haemoglobin genotypes, where Hb AS was 6 per cent, SCA was 0.9 per cent, HbS/βthal was 0.8 per cent and HbSβthal was 0.8 per cent.

**United Arab Emirates (UAE):** Among the UAE nationals abnormal HbS is one of the most common disorders. In 1984, Kamel described biochemical features of Arab SCA patients diagnosed over a 4-year period in Abu Dhabi. The frequency of SCD in the UAE was reported as 1.9 per cent in a major study conducted on 5000 subjects from three major Peninsular Arab States. Miller et al conducted a haematological survey of preschool children and reported the frequency of HbS as 4.6 per cent. In a more recent survey Al Hosani et al reported the overall incidence of SCD among 22,200 screened neonates as 0.04 per cent (0.07% for UAE citizens and 0.02% for non-UAE citizens), where the incidence of Hb AS was overall 1.1 per cent (1.5% for UAE citizens and 0.8% for non-UAE citizens). Sickle cell anaemia and HbS/β thalassaemia were identified and cases of SCA with associated α-thalassaemia and G-6-PD deficiency were frequent. Wide variations were reported in the clinical features ranging from moderate to a severe disease, with elevated Hb F levels and associated α-thalassaemia. Other investigators showed the presence of Saudi-Indian haplotype in 52 per cent of the β chromosomes that was concurrent with the mild form of the disease.

**Oman:** In a study on 5000 subjects from three States of Arabian Peninsula, the frequency of SCD in Oman was reported as 3.8 per cent. In addition, cases of HbS Omani, a variant of HbS were identified in a few families. Rajab and co-workers reported the birth prevalence of symptomatic haemoglobinopathies in 23 Omani tribes through screening of a national register, as 1 in 323 live births or 3.1 per 1000 live births during 1989-1992, which included per 1000 live births of
homozygous SCD. It was calculated that each year, 118 new cases of SCD were expected to be born and HbAS frequency was 10 per cent. The regional distribution of SCD revealed that it was more prevalent (more than 70% of cases) in regions with smear-positive rates of malaria of 1 to >5 per cent (parts of Dhahira, Dakhliya, North and South Shargiya). Al-Riyami et al reported the overall prevalence of HbS as 5.8 per cent, though there were significant regional variations. Clinical variations in SCA presentation are largely related to the presence of different β-globin gene haplotypes identified during molecular studies, where Benin, Bantu and Saudi-Indian haplotypes were shown to be present in Oman.

(ii) Arab countries in the northern region of Arabian Peninsula

Palestine: A study from Palestine on SCD reported HbS/ thalassaemia in a 12-year-old Palestinian boy with hand-foot syndrome. Later studies have revealed a higher prevalence of β-thalassaemia, though a few cases of HbS and thalassaemia co-existing in the same patient have also been reported. In a more recent study, it was shown that SCA has a severe clinical presentation and is accompanied by variable levels of HbF (1.5-17%; mean= 5.14%). Haplotype analysis shows that the Benin haplotype predominates with a frequency of 88.1 per cent, followed by the Bantu haplotype at a frequency of 5.1 per cent.

Syria: The frequency of HbS is low (<1%) in Syria, though epidemiological studies are not available. Other abnormal variant that have been reported in the Syrians include the thalassaemias as also the molecular basis of the β-thalassaemic state. A study on haplotypes associated with sickle cell gene has shown the presence of the Benin haplotype.

Iraq: The first report of the presence of HbS gene in Iraq appeared in 1971 by Khutsishvili. Thereafter, reports have shown that β-thalassaemia major and SCA are important health problems in Iraq. The frequency varies in the different areas, where a study in four villages of Abu-al-Khasib in Southern Iraq, on school children in the age group of 10 to 12 yr showed an overall HbS prevalence rate of 16 per cent as compared to 2.5 per cent seen in a control population of children belonging to five urban schools in Basrah and sickle cell trait was evident in 13.3 per cent of the cases. In a recent study on population in Basra with age ranging from 14-60 yr, the HbS trait frequency was 3.24 per cent. Associated G-6-PD deficiency was reported and the influence of haemoglobinopathies on growth and development was demonstrated. Steps were adopted to implement control and prevention programs.

Jordan: In a study conducted on 6-10 yr old school children in Northern Jordan Valley, both α- and β-thalassaemias and HbS were identified, though HbS gene frequency was very low (carrier frequency= 0.44%). Co-existing HbS/β-thalassaemias were identified, some with elevated Hb F level, but this did not ameliorate the SCA clinical presentations. In a larger study in North Jordan, the overall prevalence of HbS and β-thalassaemia was 4.45 and 5.93 per cent, respectively and the incidence of Hb AS in the newborn sample was 3-6 per cent. The prevalence of both HbS and beta-thalassaemia was higher in the Al-Ghor area in comparison to Ajloun and Irbid. Variable clinical presentation of SCA has been reported and no correlation was demonstrated with Hb F level.

Lebanon: Dabbous and Firzli reported the prevalence of HbS gene in Lebanon. The disease was shown to be clustered in two geographic areas in North and South Lebanon and nearly all patients were Muslims. The disease was severe and the major haplotype was the Benin haplotype. Interestingly high levels of HbF were not shown to influence the clinical severity of SCA. As a result it was suggested that genetic factors other than haplotypes are the major determinants of increased HbF levels in SCD patients in Lebanon. Considerable interest was geared towards management of SCA and on clinical trials using new agents to ameliorate the clinical presentation.

(iii) The Arab countries of North Africa

Sudan: The first report of the presence of HbS gene in the Sudanese appeared in 1950. Later it was shown that the frequency of the gene varies significantly in different tribes. In some areas sickle cell trait was present in 24 per cent of the newborn and 29 per cent of those aged over five years. The SCA presentation was severe and it was frequently fatal in early childhood and was accompanied with major complications. Analysis of the haplotypes associated with the S gene indicated that the most abundant haplotypes are the Cameroon, Benin, Bantu and Senegal haplotypes.

Egypt: Some researchers hypothesized that HbS gene was present among the predynastic Egyptian and they showed the presence of HbS in mummies (about 3200
Abnormalities of haemoglobin were also identified. Other abnormalities of haemoglobin were also identified. Since then, several studies have been carried out and shown that, in Egypt, β-thalassaemia is the most common type with a carrier rate varying from 5.3-9 per cent and a gene frequency of 0.03. In Egypt, along the Nile Valley, the HbS gene is almost non-existent, but in the western desert near the Libyan border variable rates of 0.38 per cent in the coastal areas to 9.0 per cent in the New Valley oases have been reported. HbS carrier rates vary from 9 to 22 per cent in some regions. The SCD is severe with painful crises and other abnormalities. Most of the globin gene haplotypes reported are the African haplotype.

Algeria: In 1961, Juillan conducted a survey on the incidence of sickle-shaped erythrocytes in Algeria and reported the presence of HbS gene. In 1977, Trabuchet et al. showed the presence of genes for HbS, Hb C and thalassaemia in various regions of the country and reported that these genetic conditions were a major cause of severe congenital haemolytic anaemias. Co-existing Hbs/thalassaemia, Hbs/Hb C cases were also reported and Hb Setif, Hb D Ouled Rabah were described for the first time in Algerians. In 1987, Dahmane-Arbane et al. reported a case of Hb Boumerdes, an alpha chain variant (α237) (C2) Pro→Arg in an Algerian family. The propositus was also homozygous for the Hbs gene, though the sickle cell phenotype was benign. High Hb F level was reported in association with high Gγ/Gα ratio and a comparison of the clinical and haematological characteristics in SCA and Hbs/thalassaemia, showed that associated thalassaemias ameliorate the clinical presentation of SCD in Algerians. Homozygous cases for haemoglobin J Mexico (α54 (E3)Gln replaced by Glu) have been reported.

Tunisia: The first case of SCA was reported in a Tunisian family in 1967 by Ben Rachid et al. Later studies showed that haemoglobin abnormalities constitute a major public health problem in many areas in Tunisia, including the central, North-western, Kebily in south Tunisia and the North-Kebili region. The SCA is generally severe in Tunisians and haplotyping using nine restriction sites in the beta-globin gene cluster revealed that the most common haplotype is the Benin type which occurs at a frequency of over 94 per cent in SCD. An atypical haplotype was also identified. The first case of SCA was reported in 1951 by Abass. Other abnormalities of haemoglobin were also identified. Since then, several studies have been carried out and shown that, in Egypt, β-thalassaemia is the most common type with a carrier rate varying from 5.3-9 per cent and a gene frequency of 0.03. In Egypt, along the Nile Valley, the HbS gene is almost non-existent, but in the western desert near the Libyan border variable rates of 0.38 per cent in the coastal areas to 9.0 per cent in the New Valley oases have been reported. HbS carrier rates vary from 9 to 22 per cent in some regions. The SCD is severe with painful crises and other abnormalities. Most of the globin gene haplotypes reported are the African haplotype.

Factors influencing the frequency of SCD

Sickle cell disease is widespread in the Middle Eastern Arab countries, although significant inter- and intra-countries differences are encountered in the frequencies of the abnormal genes. The main factors which are believed to play a major role in the increased frequencies of the Hbs include:

(i) Consanguinity: The tradition of consanguineous marriage (inbreeding) goes far back in history and has been known in the Middle Eastern Arab countries from biblical times, where such marriages are not necessarily limited to geographic or religious isolates or ethnic minorities. Several investigations have been conducted and reported high rates of consanguinity in most Middle Eastern Arab countries, though significant differences are encountered within the different countries and even between different tribes, communities, and ethnic groups within the same country. An average of about 30 per cent is seen in most Arab countries, though the prevalence of consanguinity ranges from about 25 per cent in Beirut to 60 per cent in Saudi Arabia and 90 per cent in some Bedouin communities in Kuwait and Saudi Arabia. The most common form of inter-marriage is between first cousins, particularly paternal first cousins and includes double first-cousin marriage. In a study conducted on thalassaemias in Lebanon, it was reported that 49 per cent were offspring of first-cousin marriages, and it was suggested that consanguinity was responsible for the multiplication of the incidence of β-thalassaemia by a factor of 1.66. Other studies in other countries have demonstrated various aspects of reproductive behaviour, reproductive wastage, increased morbidity...
and mortality, and increased prevalence of genetic defects in the offspring of consanguineous mating. There are several contributing factors to this pattern, including the keenness of the people to keep the property within the family or tribe, and the attachment of people to their families or villages, the belief that cousin takes better care of each other, and the popular belief that consanguineous marriage offers a major advantage in terms of compatibility of the bride and her husband’s family, particularly her mother-in-law.

(ii) Environmental factors: In the Middle Eastern countries, a major role is played by malaria endemicity in influencing the HbS gene frequency, like in other parts of the World. The carriers of HbS have a natural resistance against malaria development and this is a major advantage to survival in adverse conditions. Several reports in the Middle Eastern Arab countries validated the “malaria hypothesis” by showing a close correlation between the frequencies of the abnormal gene and past and present history of malaria endemicity.

(iii) Large sibship size: In general, the sibship size is large in the Middle Eastern Arab countries, e.g. in Saudi Arabia an average of 6-7 children/family is the norm. In families with the mutant genes a possible disadvantage is that a larger number of family members may have the abnormal genotype. This is demonstrated by several family examples. In eight families in Algeria with an average of 6 children/family, Hb Jα-Mexico was found in 116 subjects.

(iv) Migration: The rate of population migration between and within the different Middle Eastern Arab countries is high. To some extent this is caused by prevailing financial opportunities and job prospects. In the Gulf States, including Saudi Arabia, a large sector of the population is formed of immigrants’ workers, who have migrated from high-frequency areas, where they have established. Some have settled in these countries for decades, and as a result of inter-marriages, genetic admixtures have been generated. This gene drift has led to the establishment of the abnormal genes in several areas.

Clinical presentation of SCD

The major symptoms of SCD are mild to severe anaemia, painful crises, frequent infections, hand and foot syndrome and stroke. Some patients require frequent blood transfusion, while others may never need a single transfusion during their lifetime. In severe form of SCD, the patients have retarded growth, bone defects, multiple organ dysfunction and other complications due to frequent transfusion requirements, while patients with a mild disease may reach average height and have no multiple organ abnormalities.

The SCD in different Middle Eastern Arab countries shows a significant variation in its clinical presentation. In Arabian Peninsula; Bahrain, UAE, Oman, Kuwait, and Qatar, the disease is generally mild with a mild to moderate anaemia and a few complications, though some patients have a severe disease requiring regular blood transfusions and hospitalization.

In Saudi Arabia, the eastern province shows similar mild pattern, while in the western part of the country, along the Red Sea, and Yemen, the severe pattern predominates. In countries of North Africa, including the Sudan, Egypt, Libya, Algeria, Tunisia and Morocco, most SCA patients have a severe disease, though cases with the mild form have also been reported.

Genetic factors contributing to the variability of SCA

The mutation in all SCD patients is the same GAG to GTG transversion in the 6th codon of the β globin gene. But clinically it is very diverse, ranging from a severe, life threatening state to a benign, almost asymptomatic form. Several genetic factors have been shown to be implicated in modulating the clinical presentation, where some ameliorate the disease while others have an augmenting influence. These are listed in the Fig.

It was suggested that co-existing genetic abnormalities, such as G-6-PD deficiency or the thalassaemias or other abnormal Hb variants, ameliorate the clinical presentation of SCD, thus producing a benign form of the disease. In addition, the presence of an elevated level of Hb F was considered as an ameliorating factor. The Saudi SCA patients in the eastern province were easily distinguishable from those of African origin by the mildness of clinical manifestations and the lower incidence of...
vano-occlusive complications, persistence of splenic functions, lower morbidity due to other complications and lower risk during pregnancy. Amelioration was attributed to elevated Hb F in the Saudi patients. However, later studies revealed mild SCD, SCA or double heterozygotes, even in the absence of elevated levels of Hb F.

Several studies confirmed the role of β- globin gene haplotypes in influencing the SCA clinical presentation. If the HbS mutation takes place on a chromosome carrying the Saudi-Indian haplotype, the HbS generally gives rise to a mild form mostly with an elevated Hb F. The same mutation, if occurs on a chromosome carrying a Benin haplotype, is generally associated with lower Hb F levels and a severe disease. Elevated Hb F levels clearly play a role in decreasing clinical severity, possibly through interfering with HbS sickling process. Associated α-thalassaemia also influences the severity of the disease and ameliorates the disease, but this depends also on the number of α-gene deleted or on the type of mutation producing the thalassaemic state. Presence of associated β-thalassaemia influences the clinical presentation, and is dictated by the nature of β-thalassaemia mutation. β+ mutations producing HbS/β+ thalassaemia state have an ameliorating effect, while β0 mutations result in HbS/β0-thalassaemia and this state may be equally severe as SCA. The role of presence of different polymorphic sites (Xmn I polymorphic site 5' to Gγ gene and HpaI polymorphic site 3' to β gene), is also generally believed to be an ameliorating factor. Studies on the effect of Hb F, and Gγ/Aγ ratio have demonstrated that patients with a mild disease generally have a high ratio, while the reverse is true in patients with a severe disease. Contradictions are frequent when it comes to associated G-6-PD deficiency, where both ameliorating effects and adverse effects have been reported in studies reported from the Middle Eastern Arab countries. There may be several other, yet unidentified genetic loci which also influence the SCD clinical presentation, since many patients who do not carry Saudi-Indian haplotype, or elevated Hb F level or the other possible ameliorating factors have a mild disease or vice versa.

**Management strategies**

There is a significant diversity in management protocols applied for the SCA and SCD patients in the different Middle Eastern countries due to diversity of the clinical presentations and risk factors and the status of health care. It is well documented that comprehensive and regular medical care plays an important role in the well being and normal survival of SCA patients. In some of the countries the care is near optimal, while the reverse is true in others. The management protocols for SCA patients have been slightly modulated to reach the most appropriate protocol. In some centres early diagnosis is emphasized and is followed by pneumococcal vaccination and penicillin prophylaxis, to prevent infections. Acute painful crisis is common sequel that can cause significant morbidity and negatively impact the patient’s quality of life. Proper nutrition and health care play an important role besides avoiding the factors that may cause crisis, such as low temperature, dehydration, high fever and infections. Patients suffering from crisis may be hospitalized and are often transfused. The most common analgesics are paracetamol, Voltaren and morphine sulphate. However, some studies reported that adults with painful conditions often receive inadequate or no analgesic treatment. Splenectomy is frequently carried out for SCA patients where indications for splenectomy are recurrent acute splenic sequestration and hypersplenism. In general, splenectomy has been beneficial in eliminating the risk of splenic sequestration in SCA patients and in improving the blood counts in SCA with hypersplenism.

Several investigators have used agents that may elevate Hb F level and hence decrease disease severity. Hydroxyurea has been successfully used and established for treatment of SCA and HbS/β-thalassaemia patients. Paracetamol has been used successfully in children as a drug that helps reduce painful crises and improve blood circulation. Bone marrow transplantation for suitable patients is carried out in specialized centres and recent reports demonstrated the usefulness of stem cell transplantation.

**Steps towards control and prevention**

Both SCA and SCD pose health problems in most of the Middle Eastern Arab countries. Beside the need for care and rehabilitation for the affected patients, effective strategies for control and prevention were recognized as an essential measure toward decreasing the birth of affected children (primary prevention). For a successful control programme, education, counselling and increasing the awareness of chronicity of SCA condition are essential approaches. Several countries have adopted effective steps directed toward prevention, such as (i) community screening and school screening programme have been implemented;
(ii) inclusion of relevant information in the school curricula has been adopted; and (iii) articles published in newspapers, talk shows on the radio and TV, workshops and symposia and special days for SCA are held to improve the awareness of the general public and the health care providers. An effective preventive programme was the premarital screening, which has been adopted in some of the countries, including Saudi Arabia. The programme was initiated in 1998 and was implemented in a step-wise manner. The programme was complemented by genetic counselling services for the carriers and the diseased and offered by trained counsellors and wedding authorities. These services are expected to increase awareness and provide equitable access to health services, improve quality of life of those affected and help achieve primary, secondary and tertiary prevention.

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