Hemoglobin Electrophoresis and Hemoglobinopathies in Kuwait

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Abstract
Objectives: To analyze the results of hemoglobin electrophoresis (HE) in the routine laboratory of a tertiary hospital in Kuwait and to review the common types of hemoglobinopathies prevalent in the country. Methods: This was a prospective study of HE performed on 2,386 samples in Mubarak Al-Kabeer Hospital, which serves more than 30% of the population of Kuwait, from June 1997 to May 1998. Results: Of the 2,386 HE tests, only 561 (23.5%) had abnormal hemoglobin genotypes. The most commonly identified hemoglobinopathies were β-thalassemia minor (14%), sickle cell trait (6%), sickle cell anemia (0.9%), Sβthal (0.8%) and Sβthal (0.8%). Two rare hemoglobin variants, Hb DPunjab and Hb E, were encountered. Conclusion: HE yielded only 23.5% abnormal results, thus indicating the need to streamline requests for the test. The test should be limited to patients with hematological and clinical features suggestive of hemoglobinopathies or to individuals with a positive family history.

Introduction
Mubarak Al-Kabeer Hospital is the major teaching facility of the Faculty of Medicine, Kuwait University. Located in the largest of Kuwait’s five governorates, it covers about 30% of the overall population of 2.2 million people. Anemia is a major medical problem in Kuwait, and a recent study using WHO guidelines found more than 40% of children and adolescents to be anemic [1]. Hemoglobin electrophoresis (HE) is a specialized test that is done to diagnose certain types of hemolytic anemias (hemoglobinopathies) due to abnormal structural hemoglobins (Hb) and the thalassemia syndromes. HE also detects the carrier state of such diseases, which is important in family counseling. Since hemoglobinopathies are prevalent in the Arabian Peninsula [2–5], many physicians routinely request HE in the initial investigation of anemia based mainly on clinical suspicion.

In the Hematology Department of Mubarak Hospital, we have noticed a vast number of requests for HE test, the majority of which do not appear to have clear justification. An audit of HE has not yet been carried out. However there was a previous study of the prevalence of hemoglobinopathies in Kuwait in 1993 [2].

The present study was designed to prospectively analyze all samples sent to the Hematology Laboratory for HE to ascertain its usefulness in the diagnosis of hemoglobinopathies and to further review the common types of hemoglobinopathies that are prevalent in the country.
Materials and Methods

Routine HE was performed on all samples for which the test was requested from June 1997 to May 1998. The following were carried out on each sample, prior to the HE run: complete blood count (CBC), using a Coulter counter (model STKR), sickling test, blood smear and reticulocyte stain. A fresh hemolysate was prepared, which was used for the HE on cellulose acetate medium at an alkaline pH. The relative concentrations of the Hb bands were determined by densitometry. All abnormal results were crosschecked on citric acid gel electrophoresis. The Hb A2 levels were also checked using microcolumn chromatography, while fetal Hb (Hb F) was determined using the alkali-denaturation technique. These procedures and techniques were done as recommended by Dacie and Lewis [6]. The results were then reported by hematologists following a review of the CBC findings, blood films, reticulocyte stains, examination of the HE strips and densitometric graphs.

Results

A total of 2,386 HE tests were done during the study period at an average of 200 per month. Correspondingly an equal number of CBC, blood films, reticulocyte preparations, sickling tests, Hb A2 and Hb F estimations were performed. The vast majority, 182 (76.5%) of the HE tests had a normal Hb AA pattern, while only 561 (23.5%) were abnormal or positive for hemoglobinopathy.

Table 1 shows the distribution of the Hb genotypes identified. The most commonly identified abnormal pattern was A/A2, i.e. β-thalassemia minor (14%). Sickle cell trait (AS), sickle cell anemia with elevated Hb F (SF), Sβ-thal (S/A/A2) and Sβ0-thal (S/F/A2 and S/A2) were 6, 0.9, 0.8 and 0.8%, respectively. Two patients with F/A2 and a patient with F/A had β-thalassemia major. Most of the individuals with AF pattern were infants below 6 months of age. The two rare Hb variants detected in this study were Hb D with 6 heterozygotes (AD) and 1 compound heterozygote (A/D/A2 i.e. Dβ⁺-thal) and Hb E (EF). These rare variants were confirmed with cation-exchange high performance liquid chromatography (HPLC).

Discussion

Considerable resources (both manpower and financial) are expended on running HE in the routine hematology laboratory. For every sample, CBC is done along with blood film examination for red blood cell (RBC) morphology and staining both for reticulocyte count and H inclusions. The latter, when present, is suggestive of an α-thalassemia state. For meaningful interpretation of HE result, Hb A2 should be estimated by microcolumn chromatography, especially when there is Hb S in the sample. This is because some denatured Hb S may comigrate with Hb A2 on electrophoresis. In certain situations, when optimal separation of bands is desired, citrate gel electrophoresis may have to be done to identify the abnormal Hb variant(s). During the year under study, about 200 runs were made every month, giving a total of 2,386 with only 23.5% showing abnormal or positive results, indicating the poor yield of the test. There is, therefore, a need to streamline requests such that the yield from HE can be improved.

The commonest cause of anemia, especially when it is hypochromic and microcytic, is iron deficiency. In any situation when there is no strong family history or other circumstances to suggest an inherited hemolytic anemia, iron deficiency should be initially ruled out before further testing is carried out. This can be achieved by using appropriate tests of iron status and/or a therapeutic trial of iron. If the tests are negative and/or there is no response to iron, then HE is justified, to rule out thalassemia, bearing in mind that the test will help in identifying most cases of β-thalassemia (minor or major). α-Thalassemia trait, on the other hand, may be difficult to diagnose with this test and may require molecular techniques. In a situation where a sickling disorder is suspected on clinical grounds or because of a positive family history, an initial sickling
Fig. 1. Flowchart showing the suggested scheme for investigating hemoglobinopathies. MCV = Mean corpuscular volume; MCH = mean cell hemoglobin; RDW = red cell distribution width.

The present study showed that the most commonly identified hemoglobinopathy traits in Kuwait are β-thal and Hb S. The pattern of Hb variants and genotypes is similar to that previously reported [2], except that Hb J and Hb Lepore were not encountered. The only rare variants in the present study were Hb D and Hb E. The former, also known as Hb D_Punjab or D_Los Angeles, is a β-globin chain variant in which glycine is substituted for glutamate in the 121st amino acid position. It is most common in the Indus valley, but is quite widespread. Its mobility is similar to Hb S in an alkaline medium and it does not separate in an acidic medium. That is why its presence was confirmed by cation-exchange HPLC. Hematological parameters are usually normal in heterozygotes and homozygotes [7, 8]. However, in combination with Hb S, it presents a severe sickling disorder and in combination with β-thal (Dβ0-thal), it causes moderate hypochromic, microcytic anemia [9, 10].

Hb E is also a β-globin chain variant in which lysine substitutes for glutamate in the 26th amino acid position. It causes mild microcytosis in the heterozygote and normal to mild anemia in the homozygote. When coinherited with β-thal trait, it may cause severe anemia and the patient may be transfusion-dependent. It can be separated from Hb A at an alkaline, but not acidic pH. While it separates well from Hb A on cation-exchange HPLC, it may coelute with Hb A2 [7, 8].

It should be noted that most Hb S homozygotes in the present study have a significant elevation of Hb F, such that the genotype is reported as S/F and the compound heterozygote with β-thal is reported as S/F/A2. This is because the β-globin gene mutation in most Kuwaitis is on a chromosome with the Saudi Arabia/India haplotype [11]. This haplotype is associated with the –158 (C→T) mutation on the Gγ-globin gene and as such the patients have persistently elevated Hb F levels, even as adults.

**Conclusion**

HE is one of the most frequently requested tests in the routine hematology laboratory. Quite often, however, it may not be justified. In the present study, only 23.5% of the tests yielded a positive result. There is, therefore, a need to streamline requests to make the test cost-effective. It should be limited to individuals with clinical and/or hematological parameters suggestive of a hemoglobinopathy and in individuals with a positive family history.
References