NINETEEN YEARS STUDY OF BETA-THALASSAEMIA IN SLOVAKIA

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INTRODUCTION

Little attention is paid to the congenital disorders of red blood cells and other emergent haematological problems due to their rare occurrence and the fact that they do not cause significant trouble to the carrier. Nonetheless, haematologists encounter a small incidence of anaemia, which remain outstanding and present an unnecessary burden on the patient’s diagnostic and therapeutic interventions. Haemoglobinopathies prevail, beta-thalassaemia in particular.

Although the disease can be most commonly found in the Mediterranean region and in South-East Asia due to migration, we can come across it anywhere in the world. Given the raids from South-East Asia in the past centuries and migration, Slovakia is no exception. Similarly, since opening the borders in the past decades, we can encounter the aforementioned mutations in foreign nationals visiting Slovakia and our medical personnel working abroad encounter these mutations abroad. At the same time, they occur in the offsprings of mixed marriages where a spouse comes from the most seriously affected area.

Beta-thalassaemia results from mutations in the beta-globin molecule. In the clinical presentation, it varies from quite subtle changes to severe anaemia. Commonly, the heterozygotes present a noticeable microcytosis of erythrocytes, normal values of iron parameters, and elevated haemoglobin electrophoresis fractions $A_2$ (HbA2), possibly also haemoglobin F (HbF).

In the past, diagnosis of beta-thalassaemia was established rather sporadically, therefore, in 1993, we decided to conduct an epidemiological study with the aim to actively seek out carriers of beta-thalassaemic genes in Slovakia. The study was aimed at:

- simplifying diagnosis of beta-thalassaemia and avoiding unnecessary treatment by iron preparations;
- determining the prevalence and type of mutations found in our territory;
- according to the type of prevailing mutations, selecting the appropriate genetical methods for their screening;
- introducing genetic counselling (particularly before entering into a marriage with a partner from the affected area);
- entering the results into the EU-wide screening programme for haemoglobinopathies.
The first part of the study lasted 12 years (1993–2004) with 70 doctors participating in the study. The centre of the study were 3 haematology centres in Bratislava. Diagnosis was based on a clinical examination with thorough family history, particularly with the finding of microcytosis of erythrocytes in the blood test, normal values of iron parameters, and elevated levels of HbA₂, possibly also HbF in the electrophoresis of haemoglobin.

Overall, we managed to record 346 patients with suspected heterozygous beta-thalassaemia, namely heterozygotes with beta-thalassaemia minor or intermedia from the clinical view. In addition, we conducted molecular genetic testing in 37 patients (10.7% of the studied group). The prevailing mutation was IVS 1.110 and IVS 2.1, which are common Mediterranean mutations. It reflects the historical gene transfer onto our territory from the Mediterranean area. Results of the study were published (1, 2).

After evaluation of the first part of the study, it was clear that the recorded cases represented only a fraction of potentially affected population. Since we kept receiving requests for diagnostics of haemoglobinopathies from other centres, the need for diagnosing haemoglobinopathies, centralisation of patients as well as genetic counselling became obvious. We decided to establish a Study Group for Beta-thalassaemia and Other Haemoglobinopathies in 2005 and continued in the haemoglobinopathy search in Slovakia. Molecular genetic diagnosing of beta-thalassaemia conducted in Slovakia significantly contributes to this field. In this paper, we present outcomes of the second phase of the study which lasted 7 years (2005–2011).

MATERIALS AND METHODS

Patients with suspected beta-thalassaemia were addressed in the local haematological surgeries or were sent directly to the aforementioned haematological centres. Criteria for participating in the study:

- history (family history);
- objective testing (e.g. splenomegaly);
- microcytosis of erythrocytes in the blood count: MCV (medium volume of erythrocytes) below 78 fl, possibly MCH (medium size of haemoglobin in the erythrocyte) below 25 pg, anæmia was not necessarily present;
- finding in the peripheral blood smear: microcytes, target erythrocytes, possibly further morphological changes;
- normal or elevated levels of serum iron, particularly ferritine or persistent microcytosis of erythrocytes following treatment by iron preparations;
- positive test for haemolysis (increased reticulocyte count, reduced levels of haptoglobin, elevated indirect bilirubin, etc.).

Methods:

- parameters of blood counts (reticulocyte count, MCV, MCH) were obtained from normal high-parametre counters of blood cells;
- peripheral blood smears were evaluated microscopically (Pappenheim’s staining);
- examination of parameters of iron, bilirubin or other biochemical parameters was conducted through standard biochemical analysers; haptoglobin test was performed by laser nephelometry;
- electrophoresis of haemoglobin tests were carried out by the acetate cellulose sheets method in the aforementioned haematological centres. The values of HbA₂ fraction below 3.5% and HbF below 1.1% (adults) were considered normal. Only patients meeting the above mentioned criteria were able to participate in the study.

If the results came back positive, we would ask the proband’s relatives to be referred to us; this included ascending, descending and horizontal line of the family.

At the same time, we commenced cooperation with the ProGen Laboratory (Molecular genetics laboratory) in Nitra, which performed diagnostics of beta-thalassaemia and other genetic haematological disfunctions. In rare cases, we consulted with the Department of Biology, Faculty of Medicine, Palacký University Olomouc.

RESULTS

Results can be found in the following Tables:

Table 1: Results of examinations conducted by the haematological laboratory of the Synlab Company, University Hospital, Kramáre, Bratislava.

Table 2: Results of examinations conducted by the haematological laboratory of the Paediatric Faculty Hospital and Clinic, Bratislava.

Table 3: Results of examinations conducted by the haematological laboratory of the L. Pasteur Faculty Hospital, Košice.

The first and the second column of the Tables show the number of examinations of haemoglobin electrophoresis in each year as well as the total number of examinations in six years. The third column shows the number of positive results of tests, i.e. HbA₂ above 3.5%. The fourth column states the number of positive results where, apart from the elevated levels of haemoglobin A₂, haemoglobin F was above 1.1 percent. In evaluating the outcome of haemoglobin, we excluded children under one year of age. Overall, we recorded 23 foreign nationals of Asian origin and their children.

In Table 1 (the Synlab Company), we find that the total number of positive results (increase in HbA₂ above 3.5%) accounted for 189 cases (52.3%) out of the total number of tests. The portion of positive results recorded by the Paediatric Faculty Hospital and Clinic in Bratislava represented 158 cases (16.2%) in Table 2 and the Faculty Hospital in Košice recorded 55 cases (9.16%) in Table 3. In the last laboratory, haemoglobin S (sickle cell anaemia) was discovered in a family of a foreign national. Out of a total of 1,834 electrophoreses of haemoglobin examinations in three

Table 1. Number of examined patients, results with higher level of HbA₂ and results with higher HbA₂ + HbF: University Hospital Kramáre, Bratislava, Synlab Comp., haematology laboratory

<table>
<thead>
<tr>
<th>Year</th>
<th>Number of patients</th>
<th>Number of HbA₂ &gt; 3.5%</th>
<th>Number of HbF &gt; 1.1%</th>
</tr>
</thead>
<tbody>
<tr>
<td>2005</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>2006</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>2007</td>
<td>45</td>
<td>24</td>
<td>24</td>
</tr>
<tr>
<td>2008</td>
<td>78</td>
<td>49</td>
<td>49</td>
</tr>
<tr>
<td>2009</td>
<td>144</td>
<td>85</td>
<td>83</td>
</tr>
<tr>
<td>2010</td>
<td>94</td>
<td>31</td>
<td>22</td>
</tr>
<tr>
<td>Total</td>
<td>361</td>
<td>189</td>
<td>178</td>
</tr>
</tbody>
</table>
aforementioned centres the number of suspected heterozygous beta-thalassaemia cases was 402 (21.9%). It is noteworthy that in 23 cases, representing 5 families, an unusually high level of F haemoglobin was detected in adults (6–21%) without elevated levels of HbA2; we contemplated an unknown form of haemoglobinopathy or hereditary persistence of haemoglobin F. We carried out molecular-genetic testing for beta-thalassaemia in 87 patients and found mutations of beta-globin chain in 70 of them. Table 4 illustrates analysis results. The most frequent mutations included IVS 2.1 (G→A) in 28.5%, IVS 1.110 (G→A) in 25.6% and IVS 1.1 (G→A) 11.3%, which represent typical Mediterranean mutations spread on our territory by immigrants from this area. Particular mutation – cd 17 (AAG→TAG) (Lys STOP) has been demonstrated in the Department of Biology in Olomouc. In the remaining cases, beta-thalassaemia was either not detected or it was a case of mutations which were impossible to detect by the genetic testing available. Table 5 illustrates all Mediterranean mutations examined in this study.

**DISCUSSION**

Every year, 300,000 children are born with serious congenital hemoglobinopathies in the world; approximately two thirds of them in less or average developed countries. Eighty-three % of them suffers from sickle cell anaemia, 17% from thalassaemia.

Haemoglobinopathies account for 3.4% of children mortality under 5 years of age in the world. Approximately 7% of pregnant women worldwide are carriers of beta or alpha-thalassaemia genes, haemoglobin S, C, D or E genes. With the continually improving hygiene standards in third world countries, the percentage of neonatal and child mortality decreases, which results not only in an increase in surviving severe forms of haemoglobinopathies, but also in the number of mutated genes carriers.

In Sout-East Asian countries with high prevalence of haemoglobinopathies, there is a long-term effort to map the genetic region and introduce effective measures to reduce their occurrence. For example, in Japan, beta-thalassaemia is mapped very well, the incidence of beta-thalassaemic genes is 1 in 1000 inhabitants (3). In India, they discovered that 4.05% carriers of beta-thalassaemic genes (from 1 to 17% depending on the area) (4) can be found.
among the population. In Bahrain, in a 10-year study involving 60,000 healthy individuals, 3.5% carriers of the gene were found; from among 13 types of known mutations, 4 of them represented 80% (5). In the US, a study conducted in California between 1990–1996 revealed 0.17% incidence of sickle cell anaemia or thalassaemia in subjects of white non-Hispanic origin. In the Afro-American population, the incidence was significantly higher; haemoglobin S was found in 4%, in 1.5% haemoglobin C and in 4% beta-thalassaemia.

A large number of studies from the 80s and 90s was mapping the situation in the most affected areas of the Middle East and Southern Europe (6–8). In central Europe, in countries with a large number of immigrants, e.g. in Germany, 300 patients with beta-thalassaemia major were recorded (9) in the year 2000, and in Switzerland, 2,672 heterozygotes with beta-thalassaemia were recorded in an epidemiological study conducted between 1968–1974. Only 2.6% were native people, the remaining were immigrants from Turkey or Southern Europe. Overall, 56 haemoglobinopathies with severe form of the disease were recorded (10). Since the establishment of the European Union, post-communist countries (Romania, Bulgaria, former Yugoslavia) have started to pay attention to this issue. For example, in a study in Romania, the incidence of the beta-thalassaemic gene was found in 0.49% (11). Great Britain has become exemplary in diagnosing and treating haemoglobinopathies. As a multi-cultural country, it had to start tackling the high prevalence of haemoglobinopathies in some minority groups of the population several decades ago. In 1999, all types of recorded haemoglobinopathies were evaluated in the Thalassaemia Register and in the Register for Prenatal Diagnosis for Haemoglobin Disorders (12, 13).

Due to the large influx of immigrants to the whole of Europe in the last decade, the problem of haemoglobinopathies has become more prominent. A necessity to monitor their prevalence in immigrants as well as in the native population has emerged given the increasing number of mixed marriages. It is interesting to notice that in the recent years the prevalence of haemoglobinopathies has been higher in Western European countries than in the “classic countries” like Greece, Italy, Sardinia, etc. The structure of haemoglobinopathies has changed; the number of carriers of sickle cell anaemia increases which is caused particularly by the inflow of immigrants from Africa (14, 15). The situation induced the need to centralise and analyse the statistical data on the incidence of diseases for better diagnosis, treatment and prevention. For this purpose, WHO established a Centre for Clinical and Applied Bioinformatics and CHIME’s WHO Collaborating Centre for the Community Control of Hereditary Disorders (Modells’ almanach). It monitors the epidemiology of haemoglobinopathies, provides information, trains doctors and analyses the financial costs. The Centre monitors the situation virtually in the entire world and maintains a database which is regularly updated. WHO Bulletin of 2008 states that 39,303 children affected by beta-thalassaemia are born every year and that there are 25,866 transplants and a number of those dependent on blood transfusions and the chelating therapy. In Europe, 992 children suffering from a severe form of the disease are born every year (16).

In developed countries, diagnostics are performed in centres attached to hospitals or on an outpatient basis. The centres have their own protocols which are derived from the patient population, type of the medical centre, laboratory and the testing method. Particular emphasis is placed on prenatal examination when both parents are carriers of severe mutations and are considering premature termination of pregnancy. Prenatal diagnosis can be avoided by conducting screening, e.g. in Cyprus and in Greece. In countries where marriages between relatives are common and the family is a carrier of the known form of haemoglobinopathy mutations, it is recommended to examine the whole family (Pakistan) (17). The rules of prevention (10) are constantly being improved and simpler screening methods are introduced, e.g. bloodshot screening (18).

In the year 1993, in the former Czechoslovak Republic, an extensive study to search the most common mutations of beta-thalassaemia was conducted (19). The study found 12 different mutations in the population. The most often were IVS 1.1 (G→A) 46.3%, CD 39 (C→T) 11.1% and IVS 2.1 (G→A) 9.3%. The results of our study were very similar. We acknowledged the occurrence of 10 mutations, which were also of the Mediterranean origin. The most common were IVS 2.1 (G→A) 28.5%, IVS 1.10 (G→A) 25.6% and IVS 1.1 (G→A) 11.3%.

Our patients are only heterozygotes with beta-thalassaemia minima, minor or intermediate in good clinical conditions. They do not require any transfusion or chelating therapy. However, in the past, beta-thalassaemia major occurred in Slovakian descendants of Croatian immigrants.

Although Slovakia has been spared of the aforementioned problems so far, we cannot assume that they will not appear in Slovakia in the future. Due to globalisation, migration of the population and hereditary beta-thalassaemic genes in our population, the incidence of homozygous forms could be a surprise to us.

We have made the first step in this area. It is comforting that haemoglobinopathies have started to be considered in the diagnostics and patients are not treated by iron preparations unnecessarily. The awareness has increased also among the population, the patients actively seek professional medical advice. The possibilities of genetic testing are expanding, the affected families are sought out and offered counselling.

Haemoglobinopathies are a global issue and the situation can be addressed only through concentrated efforts of national haematological societies with the contribution of major international health organisations and support funds (19–21).

Acknowledgements
We would like to thank Prof. MUDr. Karel Indrák, DrSc., Doc. RNDr. Vladimír Divoký, PhD and RNDr. Martina Divoká (Palacký University Olomouc) for valuable advice and assistance and all our colleagues who have participated in the work of the study group for haemoglobinopathies in Slovakia. The study group operates without any financial support or personnel.

REFERENCES


Received December 16, 2011
Accepted in revised form July 2, 2012