Haemoglobinopathy prevention program in Turkey

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Abstract

Thalassemia and abnormal haemoglobins are a serious health problem in Turkey. Very important steps for toward preventing thalassemia have been taken in Turkey by Ministry of Health (MOH), Turkish National Haemoglobinopathy Council (TNHC) and Thalassemia Federation of Turkey (TFT) since 2000. In 1993, a law was issued called Fight Against Hereditary Blood Disease especially for thalassemia and haemoglobinopathies. The law commends to prevent haemoglobinopathies and to treat all patients with haemoglobinopathy and thalassemia. A pilot project was started and centres were created in the MOH Hospitals in the southern provinces of Turkey. In 2000, TNHC was installed to combine all centres, foundations, and associations into one organization controlled by the MOH. In 2001, the MOH and the TNHC made an inventory of all recorded patients with thalassemia and abnormal haemoglobin in Turkey, registering at least 4513 patients. In 2002, written regulations for the Fight Against Hereditary Blood Disease were published. MOH and TNHC selected 33 provinces situated in the Thrace, Marmara, Aegean, Mediterranean and South Eastern regions with high birth prevalence of severe haemoglobinopathies. In 2003, the haemoglobinopathy scientific committee was set-up, a guidebook was published and a national Haemoglobinopathy Prevention Program (HPP) was started in these high risk provinces. This program is running in these provinces successfully. In 2005, TFT was established as a secular society organization instead of TNHC. In 2007, National Thalassemia Prevention Campaign (NTPC) was organized for public education by TFT. This campaign contributed very important supporting to HPP in Turkey, because totally 62,682 people such as health workers, students, teachers, demarches, religion officers and the other many people were educated for preventing thalassemia and haemoglobinopathies. In 2009, National Thalassemia Education Seminars (NTES) for health personnel have been planned in 26 cities by MOH and TFT. A total 3,600 health persons were educated on thalassemia prevention and therapy with NTES in 18 centres in 2009 and 2010. In conclusion, according to reports of MOH, 46 first level haemoglobinopathy diagnosis centres, 5 second level diagnosis and therapy centre and 5 third level prenatal diagnosis centre were set-up and licenced in 30 cities between 2003 and 2009. While premarital screening tests were 30% of all couples in 2003, it increased continuously during 6 years and it reached 81% in 2008. The number of new born with thalassemias and haemoglobinopathies was 272 in 2002, it was decreased to 23 in 2008, as a result there has been an 90% reduction in new affected births.

Background

In 1958, the first clinical and hematological studies on thalassemia and haemoglobinopathies in Turkey was published by Aksoy et al [1]. In 1971, Cavdar and Arcasoy reported that overall incidence of β-thalassemia in Turkey was 2.1%.2 In 1983, first prenatal diagnostic procedures for identifying haemoglobinopathies were performed by Altay et al.3 In 1987, Akar et al. showed that the most frequent thalassaemia allele in the Turkish population was the IVS-1 nt 110 mutation, which was the most common thalassaemia mutation in the majority of the high risk regions of the Mediterranean area.4 In 1992, Başak et al. reported the spectrum of β-thalassemia mutations occurring in the Turkish population.5 In 1993, a law was issued called Fight Against Hereditary Blood Disease a pilot Project was started and centres for Research and Treatment for Hereditary Blood Disease were created in the Ministry of Health (MOH) Hospitals in the south provinces of Turkey, Antalya, Antakya, Mersin, Mugla regions. In 2000, Turkish National Haemoglobinopathy Council (TNHC) was installed to combine all centres, foundations and associations into one organization controlled by the MOH. In 2001, the MOH and the TNHC made an inventory of all recorded patients with thalassemia and abnormal haemoglobins in Turkey.6-8 In 2002, Altay published that screening studies for abnormal haemoglobins and β-thalassemia during the last 40 years. These studies revealed most common abnormal haemoglobin was HbS followed by Hb D, HbE, and Hb O Arab. In addition 42 abnormal haemoglobins were identified in Turkish population.6 In 2002, TNHC made an inventory of all recorded patients with thalassemia and abnormal haemoglobinopathies. In 2003, the haemoglobinopathy scientific committee was set-up, a guidebook was published and a national Haemoglobinopathy Prevention Program (HPP) was started in these high risk provinces on the 8th of May, 2003.6-8

The aims of HPP are following up: i) to assess the present situation in each region by recording patients and carrier, ii) to produce a...
Materials and Methods

The General Director of the Mother and Child Health / Family Planning of the MOH was responsible for national HPP depends on the written regulations of the FAHBD. According to FAHBD plans, the project included three levels. First Level (Diagnosis and Screening Centres); new reference centres were created for public education, screening and genetic counselling and existing centres were officially licensed. Second Level (Diagnosis and Therapy Centres); these centres were created for state of the art treatment after the diagnosis is established for the diagnosis, therapy and follow-up of patients with thalassemia and haemoglobinopathies. Third Level (Prenatal diagnosis, mutation analysis or bone marrow transplant centres); the existing centres at university hospital had to be licenced according to the special regulations. The General Director of the Treatment Service of the MOH was responsible for therapy. In this way, both directors were working in a coordinated way with the TNHC. A public health manager in each city coordinate this project and organize a team for education, screening, genetic counselling and the records of all patients and carriers on regular basis.

Thalassemia Federation of Turkey (TFT) was established as a secular society organization instead of TNHC in 2005. Thalassemia Federation have defined the following objectives: i) To speak with one voice at all national and international conventions and gatherings, ii) To find solutions for the problems of haemoglobinopathy patients, their families and haemoglobinopathy centers iii) To determine standard procedures for prevention and treatment of haemoglobinopathy.

National Thalassemia Prevention Campaign (NTPC) was organized for public education and supporting of HPP was started by MOH and TFT in 2007.

National Thalassemia Education Seminars (NTES) for health personnel have been planned by MOH and TFT in 26 cities in 2009.

Results

The MOH and the TNHC picked up the results of screening from 16 different cities on the Turkish Mediterranean coastal areas and western regions around the Aegean Sea and Marmara Sea made between 1995 and 2000. In total 377.339 healthy subjects were screened, yielding an average frequency of β-thalassemia trait was 4.3%. The highest prevalence of β-thalassemia trait (13.1%) was found in Antalya region and Hbs trait (10 %) was found in the Çukurova region.

The MOH and the TNHC made a registry of all recorded patients with thalassemia and abnormal haemoglobins in Turkey, registering at least 4513 patients. The classification of these patients was 2567 β-thalassemia major, 1050 Sickle Cell Disease (SCD), 700 β thalassemia intermedia, and 196 cases in various combinations with other abnormal haemoglobins in 2001.

Thalassemia Federation authority committee arranged an assembly with Ministry of the Health, Ministry of the Internal Affairs, Ministry of Education, Ministry of Religion to take support of whole official foundations for the NTCP. After an academic committee prepared master slides for public and health workers’ education programs. A long vehicle Talotir was specially designed as an education hall and symbolically represented the National Prevention Campaign. A total 23 cities in Thrace, Western and Southern Anatolia with high incidence of thalassemia were chosen for program. Target population of education were mainly doctors, other health staff, teachers, students, demarches, religion officers, patients, their families in addition to public. This campaign contributed very important supporting to HPP in Turkey, because totally 62.682 people such as health workers, students, teachers, demarches, religion officers and the other many people were educated for preventing thalassemia and haemoglobinopathies.

A total 3600 health persons were educated on thalassemia prevention and therapy by NTES in 18 centres in 2009 and 2010.

According to reports of MOH, 46 first level haemoglobinopathy diagnostic centers, 5 second level diagnosis and therapy centre and 5 third level prenatal diagnosis centre were licenced in 30 cities between 2003 and 2009. While premarital screening tests were 30% of all couples in 2003, it was increased continuously between 2003 and 2008 and it was reached 81%. (Figure 1).

The number of new born with thalassemia and haemoglobinopathies was 272 in 2002, it was decreased to 23 in 2008, as a result there has been an 90 % reduction in new affected births. (Figure 2).
Discussion

Thalassemia and abnormal haemoglobins are a serious health problem in the all world. It is estimated that in excess of 300,000 children are born each year with a severe inherited disorder of haemoglobin and that approximately 80% of these births occur in low- or middle-income countries. World Health Organization (WHO) started first studies on prevention of thalassemia in Mediterranean countries in 1970's. WHO guidelines for the control haemoglobin disorders includes optimal management for patients, community-based prevention education, prospective heterozygote screening programs, genetic counselling and prenatal diagnosis services based on an integrated strategy definitions.

Prevention strategies for severe haemoglobinopathies changes in endemic and nonendemic immigration countries such as the Latium example. Thalassaemia carrier screening programmes provide a unique opportunity to compare the delivery of carrier screening programmes carried out in different cultural, religious and social contexts. The manner in which thalassaemia carrier screening programmes are structured among different populations varies greatly in several aspects, including whether the programmes are mandatory or voluntary, the education and counselling provided and whether screening is offered pre-pregnancy or antenatal.

Thalassemia and haemoglobinopathies are very important health problem in Turkey. After regulations for the FAHBD published, a national HPP was started in selected 33 provinces situated in the Thrace, Marmara, Aegean, Mediterranean and South Eastern regions with high incidence of severe haemoglobinopathies by MOH and TNHC on the 8th of May, 2003. New diagnostic centres were created for public education, premartial screening tests and genetic counselling. NTCP contributed very important supporting to HPP in Turkey, because totally 62.682 people were educated for prevention of thalassemia and hemoglobinopathies in 2007. In addition, a total 3600 health persons were educated on thalassemia prevention and therapy by NTES in 18 centres in 2009 and 2010.

According to reports of MOH, 46 first level haemoglobinopathy diagnostic centres, 5 second level diagnosis and therapy centre and 5 third level prenatal diagnosis centre were licenced in 30 cities between 2003 and 2009.

In conclusion, premartial screening tests were 30% of all couples in 2003, it increased continuously during HPP and it reached 81% in 2008. As a result there has been an 90 % reduction in new affected births in Turkey.

References