Haemoglobin D Trait-An Uncommon Haemoglobinopathy in Chattogram: A Case Report

Abstract
This case presents haemoglobin D Trait (Hb-D) which is a rare disease in Bangladesh. It is a heterozygous state for haemoglobin D who came to Laboratory Medicine Department of Chattogram Maa Shishu-0-General Hospital (CMSOGH) for haemoglobin electrophoresis. Our patient was 9 years old girl from Noakhali, Bangladesh. She was clinically silent with no splenomegaly. Her peripheral blood film was microcytic hypochromic anaemia and Capillary Electrophoresis (CE) of Hb revealed Hb D Trait. On family screening her father was Hb E trait, mother was Hb D trait and younger sister turned out normal Hb study. Compound heterozygosity for haemoglobin D and haemoglobin E may be present in subsequent pregnancy of her mother.

Key words: Heterozygosity; Hb D trait; Haemoglobinopathy.

INTRODUCTION
Thalassaemia is one of the most common hereditary disorders in the world including Bangladesh. This is a major health problem particularly in the developing countries where the resources are limited. World Health Organization (WHO) estimates that at least 6.5% of the world populations are carriers of different inherited disorders of haemoglobin. In Bangladesh the commonest varieties of Thalassaemia are Beta Thalassaemia trait (4.1%) and Haemoglobin E trait (6.1%)\(^1\). Haemoglobin D and Haemoglobin S are though rare but can be prevalent in several populations in the different parts of Southern Asia.

Hb D, a Haemoglobin variant, occurs mainly in North West India, Pakistan and Iran. It is also found sporadically in Blacks and Europeans, the latter usually coming from countries that have had close associations with India in the past\(^2\). Hb D-Punjab and Hb D-Los Angeles is identical haemoglobin. Hb D-Punjab occurs in about 3% of the population in northwest India and Hb D-Los Angeles is seen in fewer than 2% of African Americans\(^3\). HbD differs structurally from normal Hb A at 121 position on chain were glutamine is replaces glutamic acid ($\alpha^2\beta^2$,\(^{121}\text{Glu} \rightarrow \text{Gln}$). Hb D gene can be detected by DNA amplification and globin chain analysis\(^4,5\). HbD occurs in four forms: heterozygous Hb D trait, Hb D- thalassaemia, HbS-D disease and rare homozygous Hb D disease, which is usually associated with mild haemolytic anaemia and mild to moderate splenomegaly\(^6,7\). Hence we report a case of Hb D trait, an uncommon haemoglobinopathies in Bangladesh. The aim of reporting this case is that the treatment is simple but genetic counseling is important for identifying the reproductive risks of the family.

CASE REPORT
A 9 year old girl from Noakhali attended on 20th September 2020 to the Child Outpatient Department of CMSOGH for having generalized weakness and anorexia. There were no other significant complaints. Her family history was not contributory and her mother was 3 months pregnant. On examination she had mild pallor, spleen was not palpable and no other organomegaly and lymphadenopathy were present.
Laboratory examination revealed Haemoglobin 10.9 g/dl, RBC 5.36 million/cmm, MCV 63.1 fl, MCH 20.3 pg, MCHC 32.2 g/dl, RDW-CV 15.4%, Microcytic hypochromic anaemic blood film, Capillary Electrophoresis (CE) of Hb showed HbA 85.5%, HbF 1.0%, Hb D 11%, HbA₂ 2.6% indicating diagnosis of HbD trait (Figure-1). After getting such report of the patient the electrophoresis of her parents and sibling were done. Her mother aged 28 years, 3rd gravida, found moderately anaemic, Hb 9.4 g/dl, RBC 4.52 million/cmm, MCV 64.4 fl, MCHC 32.3 g/dl, RDW-CV 16.4%, Microcytic hypochromic anaemic blood film, CE showed HbA 86.0%, HbD 11.5%, HbA₂ 2.5% diagnosed as HbD Trait (Figure-2). Her father aged 38 years old had no remarkable physical finding. Fathers Hb 14.9 g/dl, RBC 6.38 million/cmm, MCV 71.0 fl, MCH 23.4 pg, MCHC 32.9 g/dl, RDW-CV 14.2%, non specific blood film, Capillary electrophoresis showed HbA 72.5%, HbE 23.8%, HbA₂ 3.7% diagnosed as HbE trait (Figure-2). Her sister aged 7 years had normal pattern of CBC, blood film and CE (Table I).

After family screening of the girl conforms she is a case of heterozygous state for Hb D. Her family pedigree is shown in Figure 2 and blood parameters of the family shown in Table I.

**DISCUSSION**

Hb D is uncommon in Bangladesh but few cases have been reported. In our case as father is HbE trait and mother is HbD trait possible siblings will be Hb D trait, which is the patient, may be normal who her sister is, other possibility will be HbE trait or compound heterozygosity for Haemoglobin D and haemoglobin E. Population demographics are changing with intermixing so Hb D trait should no longer confined to north west India & Pakistan. Haematological parameters of our case and her family are interesting but clinically not significant. Genetic counseling should be done for determining any possible reproductive risk factors. We suggest red cell indices and CE are sufficient for screening of Hb D trait in resource poor setting.

**CONCLUSION**

Patient with Hb D trait clinically asymptomatic, usually detected during investigation.

**DISCLOSURE**

All the authors declared no competing interest.
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