Inheritance of Hereditary Persistence of Fetal Haemoglobin (HPFH) in a Family of Western Odisha, India

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ABSTRACT
Hereditary persistence of foetal haemoglobin (HPFH) is a rare inherited haemoglobin disorders in India. We encountered five cases of HPFH-3 in heterozygous condition in a single family of western Odisha, India. All the cases had raised % HbF (26.1±3.23%) with pancellular distribution of HbF in erythrocytes. There were no abnormalities found in the red cell indices. All the cases were asymptomatic till date with normal growth and development. Molecular confirmation of this haemoglobin disorders is important for control and prevention of haemoglobinopathies in this region.

CASE REPORT
A 18-year-Old female of caste “Chasa” with mild fever had attended to the Sickle Cell Clinic for screening of sickle cell status. All the haematological and molecular investigations were done at Sickle Cell Clinic and Molecular Biology Laboratory, Veer Surendra Sai Institute of Medical Sciences and Research, Burla, Odisha, India. On investigation, her haemoglobin (Hb) was 12.9 g/dL with other blood parameters being normal. Sickling slide test was negative with “AF” pattern in agarose gel Hb electrophoresis (pH 8.6). Complete blood count was analysed by Sysmex KX-21; Sysmex Corporation, Kobe, Japan. Reticulocytes count was estimated by flow cytometer (BD Facs Caliber) using Retic Count reagent/thiazole orange reagent by using FACS Comp software as per the guideline. Distribution of HbF in the red cells was analysed by Kleihauer-Betke’s slide test. Hb fractions was quantified by high performance liquid Chromatography (CE-HPLC) by using the VARIANT-II Hb testing system; (Bio-Rad Laboratories, Hercules, CA), resulting a high HbF level of 25.5%. Her family screening revealed a high HbF level in other four members in the family with a mean HbF level of 26.1±3.23%.

Complete pedigree was made carefully as per their genotypes [Table/Fig-1]. All the cases were asymptomatic till date and, had normal haematological reports. DNA isolation from peripheral blood was carried out by phenol-chloroform method. Molecular analysis for high HbF was done by procedures described earlier [1]. Both Asian Indian inversion-deletion GγAγδβ0 and HPFH-3 (NG_000007,3:g,5 0509..83170del32662), the most prevalent high HbF determinants in India were investigated. All the five cases with high % HbF were found to be heterozygous HPFH-3. Agarose gel electrophoresis for molecular confirmation of HPFH-3 was shown in [Table/Fig-2]. Alpha thalassemia was investigated by GAP-PCR [2]. Only -3.7kb deletional alpha thalassemia was found in heterozygous condition in three members of the family (IIB, IIIB and IIE). None of the case of heterozygous HPFH-3 had alpha thalassemia. Cases with alpha thalassemia had microcytic-hypochromic blood picture compared to others. All the HPFH-3 heterozygous cases had normal red cell indices. Pancellular distribution of HbF in red cells was found in all the HPFH-3 heterozygous cases. Reticulocytes were in normal range in all the cases. The haematological and molecular characterization of all the family members was described in [Table/Fig-3]. Written informed consent was obtained from the cases. This study was approved by Institutional ethical committee.

DISCUSSION
Hereditary persistence of foetal haemoglobin (HPFH) are the condition caused by a large deletions involving both δ- and β-globin genes in the β-globin cluster, with increased level of HbF levels in adults [3]. In HPFH-3, the deletion removes 48.5 kb of a DNA segment, starting from the 5’ end of the ψβ gene to a region 30 kb downstream of the β-globin gene in an L1 repetitive region. This genetic trait, although rare, have been reported from different parts of India [4–6]. In literature, seven various forms of deletional HPFH have been reported. Out of which, HPFH-3 causing 48.5 kb deletion in the β-globin gene cluster is the commonest form reported in India. The present patient-based diagnostic study reveals five cases of heterozygous HPFH-3 in a single family of western Odisha. Presence of high %HbF (26.1±3.23%; range 21.8 -30.5%) and normal red cell indices in all these five HPFH-3 heterozygous cases were similar to

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![Table/Fig-2]: Agarose gel electrophoresis showing deletional high persistence of foetal hemoglobin (HPFH-3). Amplification of 841 bp PCR product is for normal and 607 bp for deletional HPFH-3. Lane A1, A2, A3 and A4 are normal and Lane A5, A6, A7 and A8 are heterozygous for HPFH-3, lane L1 and L2 are 500bp and 100 bp DNA ladder

![Table/Fig-1]: Pedigree showing inheritance of hereditary persistence of foetal haemoglobin in heterozygous condition
previous studies on HPFH reported in India [4-6]. The finding of a homogeneous HbF distribution in red cells was in agreement to the earlier report [6]. Deletional HPFH syndromes with a pancellular distribution of HbF have been reported to present benign in heterozygous state, with or without presence of other FC loci [7]. Our study agrees to this and all five cases were asymptomatic except the index (II D), who attended to the Sickle Cell Clinic and Molecular Biology Laboratory with mild fever. On the contrary as co-inheritance of HPFH with β thalassemia or HbE usually lead to severe clinical manifestation similar to that of thalassemia intermedia [8]. It becomes essential to study the allochtonous Hb disorders like the HPFH when introduced to an indigenous populations with established high frequency of HbS and β thalassemia alleles. This unique situation is particularly applicable to the western part of the province of Odisha situated in eastern India, with a reportedly high frequency of HbS and β thalassemia carrier (21% and 3.75 % respectively) and sporadic cases of HbE [9].

We have a limitation in this study that the information provided here caters to only the population of Odisha. A guideline for population based screening of such haemoglobin disorders should be necessary to generate data on national level will be a step forward for preventing and controlling program for haemoglobinopathies.

CONCLUSION

Molecular characterization of HPFH is important for understanding the resulting patho-physiology of the phenotype and they will also provide the information on the cis-acting genetic regulatory elements present in the β-globin cluster. Besides, high HbF determinants like HPFH are vital for prenatal diagnosis to prevent the affected foetus with obvious clinical manifestation when inherited with other forms of haemoglobin disorders. Indexing various forms of HPFH based on the molecular diagnosis in countries like India with a mosaic of endogamous communities and specific cultural practices will facilitate prevention and control program for haemoglobinopathies.

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