Screening of β-Globin gene (HBB) for rare mutations in β-thalassemia patients included haemoglobin S D-Punjab using Sanger sequencing

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Abstract

Objective: β-thalassemia is an autosomal recessive disorder which results in the formation of abnormal haemoglobin due to a variety of different mutations found in the HBB gene. These mutations render patients incapable of producing correct form of haemoglobin. The aim of this study was to identify HBB gene mutations in β-thalassemic patients, not included in the common-mutation panel of ARMS PCR, by sequencing HBB coding, intronic and promoter.

Method: A total of 10 samples previously tested for HBB gene mutations by ARMS PCR common-panel (i.e. IVS 1-1, IVS 1-5, Codon 8/9, Codon 41/42 and 619bp deletion) were analyzed by Sanger sequencing. Two healthy subjects were included as negative controls. Genomic DNA was isolated and HBB gene was amplified. Column purified amplified products were utilized for bidirectional cycle sequencing (Big Dye Terminator, ABI, USA).

Results: In the present study, a total of 10 samples were analyzed. Four were males and six females. The Mean of the patients was three years. All patients were diagnosed as β-thalassemia major based on their family history, clinical and laboratory findings. On average, patients were receiving transfusions every second week. Seven rare mutations in HBB gene were detected including point mutations. The mutations spanned in the promoter region HBBc.138C>A (-88 C>A), exon1 HBBc.17_18delCT (Codon5 –CT), HBBc.47G>A (Codon15 G>A), HBBc.92G>C (Codon30 G>C), HBBc.50A>C (CAP+1 A>C), exon2 HBBc.118C>T (Codon39) and intron2 HBBb.c.315+1G>A (IVS II-I G>A) and a heterozygous change at codon 6 (GAG→GTG) and also a heterozygous mutation at codon 121 (GAA→CAA) . All control subjects showed normal HBB gene sequence. In addition, a polymorphism T>C in codon3 at position HBB: c.59 was detected in majority of the patients and controls.

Conclusion: Although ARMS PCR is a fast and convenient method for detection of common mutations in the HBB gene, a small subset of patients may be missed because of rare mutations, which would require other means for diagnosis. Sanger sequencing is an accurate and robust technique to manage such patients.

Biography

Zeeshan Ansar, is the assistant Professor, Department of Pathology and Laboratory Medicine, The Aga Khan University Hospital, Stadium Road, Karachi. His learning outcomes explain the principles and evaluate the strengths and limitations of the Sanger sequencing used in molecular biomarker based diagnosis. Analyze the application of the different types of molecular biomarkers in the treatment stratification of malignant conditions. Understand impact of Sequencing based strategies in expansion and capacity building of the section of molecular pathology. He is proficient in Cancer Cell Biology, Heam-oncology and Clinical Genetics.