

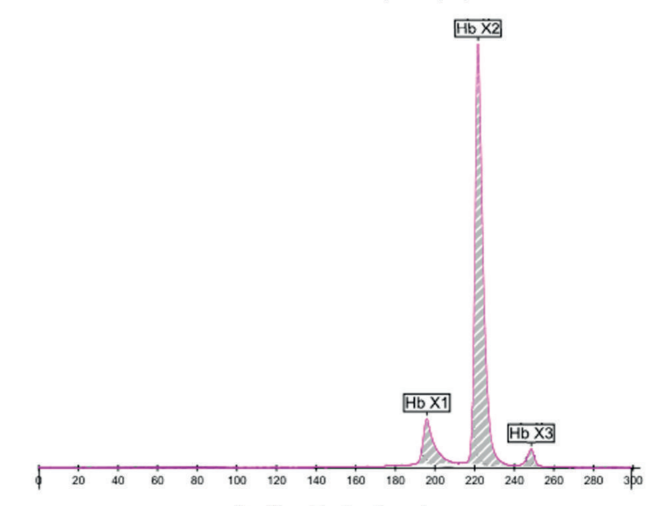


Globin chain(s) involved: Beta

Migration in zone(s): -

Status: Homozygous

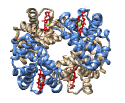
Migration in position(s): -Peak position may vary +/- 1



Capillary Electrophoresis

Fractions	Value %	
HbF (HbX1)	11,3	
Hb S (Hb X2)	85,6	
Hb A2 (Hb X3)	3,1	

Comments on this profile: In case of absence of Hb A and/or Hb A2, analyse the sample mixed with Normal Hb A2 Control to display the zones (see Package Insert). Values of each fractions must be taken on the native sample.



Mutation data

Status:	Homozygous	
Hb S		
Mutation	Beta 6 (A3) Glu>Val.	
Nomenclature	HBB:c.20A>T	
In combination with:		
Mutation		
Nomenclature		

Comments:

Hematology

Hematological parameters	Results	Hematological parameters	Results
RBC		MCH	
Hemoglobin		Blood smear	Sickle cells
Hematocrit		Serum iron and ferritin	
MCV			

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Hemolytic anemia parameters with sickle cell test positive

Other information

Clinical context:		
Clinical presentation	Severe hemolytic anemia symptoms, acute pain crises, acute chest syndromes, spleen infarctions, cerebral events	
Genetic risk	Severe risk in combination with Hb S, Hb C, Hb E, Hb D, Beta-thalassemia, Hb Lepore and other less common hemoglobin variants	
Advice	Partner and family analysis	
About this variant:		
Stability	Normal	
Oxygen affinity	Normal	
Found in	Black, Asian, Mediterranean populations	

Comments:

References: -