

Shinde Diagnostics-Ahmednagar

Balikashram Road, Savedi, Ahmednagar

Maharashtra

Ahmednagar - 414003

Contact:

Report Of: Mrs. SUSHMA AKSHAY MANE



Sample ID 2300131049 Patient ID 1002352845 Received on 13/07/2023 12:00 Registered on 12/07/2023 19:16 Reported on 13/07/2023 17:33 Referred by DR.ASHOK NARWADE

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Patient Name: Mrs. SUSHMA AKSHAY MANE	Sample Type: Whole Blood EDTA	4		
Date of Birth/Age: 26/07/1998	Gender: FEMALE	City:		
Method: High Performance Liquid Chromatography (HPLC)	Blood Transfusion History: No			
Referral Reason or Clinical History:				

Hemoglobinonathy Screening

#### About the test

Hemoglobinopathy screening by high performance liquid chromatography is a blood test that is used for detecting quantitative and qualitative abnormalities of hemoglobin (Hb), namely, Thalassemia and Structural Hb variants (e.g. HbS) respectively. The test helps identify individuals with these disorders so that they can receive timely and appropriate treatment and care. Antenatal diagnosis of these disorders allows measures to reduce the chances of the birth of an affected baby. It is also possible to screen the newborns for hemoglobinopathies using this approach, thereby decreasing the mortality & morbidity associated with conditions like Sickle cell disorder.

Test findings				
Hb Fraction	Observed Value (%)	Expected Value (%)		
HbF	2.4*% 🚹	<2%		
P2*	3.7%	<4.6%		
HbA0	82.4% 🛕	85 - 95%		
HbA2/HbE	<b>6.1</b> *% <b>⚠</b>	1.8 - 3.5%		
HbD	ABSENT	Absent		
HbS	ABSENT	Absent		

<sup>\*</sup>The mentioned P2 value from BioRad Variant-II HPLC system is equivalent of HbA1c value in BioRad D10 system

Indicates that the individual requires further evaluation and opinion from the clinician.

#### Interpretation

Chromatogram shows elevated HbA2 (6.1%) suggesting the diagnosis of Beta Thalassemia trait. Hemoglobin and PCV are reduced, RBC count is normal and red cell indices are microcytic with high RDW suggesting co-existent iron deficiency in a case of beta thalassemia trait.

### **Suggestions**

- 1) Molecular studies to confirm the above diagnosis.
- 2) Serum iron studies.3) Hemoglobin analysis of the parents and the partner by HPLC.

Verified by Mr. Pradip Kadam Incharge Biochemistry Dr. A. Dasgupta MD, PhD,

Consultant Hematopathologist

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Patient Name: Mrs. SUSHMA AKSHAY MANE

Calibrated

Sample ID: 2300131049

## **HPLC Findings**

Patient Data 2300131049 Patient ID: Name: Physician: Sex: DOB: Comments:

Analysis Data Analysis Performed: Injection Number: Run Number: Rack ID: Tube Number: Report Generated:

Retention Time (min)

07/12/2023 19:48:48 308 0010 07/13/2023 10:03:54

Operator ID:

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	30.0				
96	22.5				
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	0.0	_~~~_	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	$\triangle$	
	7	<b></b>			

Time (min.)

Analysis comments:

45.0<sub></sub>

\*Values outside of expected ranges

1615 Unknown 1.00 2.4 36984 1.11 Unknown 0.7 1.31 59831 Р3 4.6 1.72 74054 2.38 1336843 82.4 Ao 6.1\* 3.66 100540

> Total Area: 1,621,461

F Concentration = 2.4\* % A2 Concentration = 6 1 \* %

A2

# Important Blood Indices (from CBC Analysis)

Parameters	Result	Reference Range	Units
Hemoglobin (Hb)	9.24 ⚠	12 - 15	g/dL
RBC Count	4.48	3.8 - 4.8	x 10 <sup>6</sup> /μL
Hematocrit	27.00 🗥	36 - 46	%
Mean Corpuscular Volume (MCV)	60.30 🛕	83 - 101	fL
Mean Corpuscular Hb (MCH)	20.60 🛕	27 - 32	pg
Mean Corpuscular Hb Conc. (MCHC)	34.20	31.5 - 34.5	g/dL
RBC Distribution Width (RDW) (CV)	23.30 🛕	11.6 - 14	%
RBC Distribution Width (RDW) (SD)	39.80	39 - 46	fL

### Notes:

- 1. Recent blood transfusions and iron deficiency can interfere with the results, repeat testing is recommended three months after the last blood transfusion. In case of iron deficiency, it is recommended to evaluate the result post-correction of iron deficiency.
- 2. Megaloblastic anemia can cause elevated HbA2 levels. A repeat assay is recommended after correction of VitB12 deficiency.
- Mild to moderately elevated fetal hemoglobin (HbF) values are observed during pregnancy, hypoxia, chronic kidney disease, use of certain 3. drugs, myelodysplastic syndromes (MDS), aplastic anemia and conditions of stress hemopoiesis.
- Cases with borderline HbA2 levels (3.1-3.9%) could represent Silent Beta-thalassemia trait, or co-existent iron deficiency or Alpha-thalassemia 4. in a case of Beta-thalassemia trait. They need to be investigated further by appropriate tests.
- Confirmatory molecular tests for Beta-thalassemia traits and abnormal hemoglobin disorders (e.g. HbS, HbE, and HbD), followed by subsequent 5. prenatal diagnosis (If required) are available at our centre.
- The mentioned P2 value from BioRad Variant-II HPLC system is equivalent of HbA1c value in BioRad D10 system

### **Disclaimers:**

- The Hb-HPLC is a screening test that detects Beta-thalassemia and other hemoglobin variants. It does not identify Alpha-thalassemia and 1. Silent Beta-thal-assemia carriers. DNA analysis is recommended to rule out Alpha-thalassemia and Silent Beta-thalassemia carriers.
- 2. The result must be interpreted in conjunction with the complete blood counts (CBC), VitB12 and iron profile of the individual.
- Each sample received at Lilac Insights' processing centre is handled with the utmost sensitivity and care. All samples received on Sundays and 3. National holidays are stored as per specific guidelines for the respective specimens and processed on the next day.
- P2 peak in Bio Rad's Variant II HPLC platform represents glycated hemoglobin. It is elevated in uncontrolled diabetes.

Verified by Mr. Pradip Kadam Incharge Biochemistry

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Dr. A. Dasgupta MD, PhD, Consultant Hematopathologist Page 2 of 2