



**Laboratory Investigation Report**

**Name** : Mr. ORHAN AMIR ABBASI  
**DOB** : 14/03/2022  
**Age / Gender** : 2 Y / Male  
**Referred by** : DR AHSAN  
**Centre** : CITICARE MEDICAL CENTER

**Ref No.** : 37953  
**Sample No.** : 2410485351  
**Collected** : 07/10/2024 15:39  
**Registered** : 07/10/2024 21:53  
**Reported** : 07/10/2024 22:51

**BIOCHEMISTRY**

Test	Result	Flag	Unit	Reference Range	Methodology
C-REACTIVE PROTEIN (CRP)	< 0.6		mg/L	< 5.0 Please note change. Source: Roche IFU.	Particle-enhanced immunoturbidimetric assay

**INTERPRETATION NOTES :**

1. CRP measurements are used as aid in diagnosis, monitoring, prognosis, and management of suspected inflammatory disorders and associated diseases, acute infections and tissue injury.
2. C-reactive protein is the classic acute phase protein in inflammatory reactions.
3. CRP is the most sensitive of the acute phase reactants and its concentration increases rapidly during inflammatory processes. The CRP response frequently precedes clinical symptoms, including fever. After onset of an acute phase response, the serum CRP concentration rises rapidly and extensively. The increase begins within 6 to 12 hours and the peak value is reached within 24 to 48 hours. Levels above 100 mg/L are associated with severe stimuli such as major trauma and severe infection (sepsis).
4. CRP response may be less pronounced in patients suffering from liver disease.
5. CRP assays are used to detect systemic inflammatory processes (apart from certain types of inflammation such as systemic lupus erythematosus (SLE) and Colitis ulcerosa); to assess treatment of bacterial infections with antibiotics; to detect intrauterine infections with concomitant premature amniorrhexis; to differentiate between active and inactive forms of disease with concurrent infection, e.g. in patients suffering from SLE or Colitis ulcerosa; to therapeutically monitor rheumatic disease and assess anti-inflammatory therapy; to determine the presence of post-operative complications at an early stage, such as infected wounds, thrombosis and pneumonia, and to distinguish between infection and bone marrow transplant rejection.

Sample Type : Serum

End of Report

**Dr. Adley Mark Fernandes**  
M.D (Pathology)  
Pathologist

*Vyoma V. Shah*  
**Dr. Vyoma V Shah**  
M.D (Pathology)  
Clinical Pathologist

*Harshad Manikandan*  
**HARSHAD MANIKANDAN**  
Laboratory Technician

This is an electronically authenticated report

Printed on: 07/10/2024 22:53

Test result pertains only to the sample tested and to be interpreted in the light of clinical history. These tests are accredited under ISO 15189:2012 unless specified by (^). Test marked with # is performed in an accredited referral laboratory.





**Laboratory Investigation Report**

**Name** : Mr. ORHAN AMIR ABBASI  
**DOB** : 14/03/2022  
**Age / Gender** : 2 Y / Male  
**Referred by** : DR AHSAN  
**Centre** : CITICARE MEDICAL CENTER

**Ref No.** : 37953  
**Sample No.** : 2410485351  
**Collected** : 07/10/2024 15:39  
**Registered** : 07/10/2024 21:53  
**Reported** : 07/10/2024 22:19

**HEMATOLOGY**

Test	Result	Flag	Unit	Reference Range	Methodology
<b>COMPLETE BLOOD COUNT (CBC)</b>					
HEMOGLOBIN	12.1		g/dL	11 - 16	Photometric
RBC COUNT	5.2		10 <sup>6</sup> /μL	4.1 - 5.5	Electrical Impedance
HEMATOCRIT	35.9		%	33 - 47	Calculation
MCV	69.6	L	fL	74 - 89	Calculation
MCH	23.5	L	pg	27 - 32	Calculation
MCHC	33.7		g/dL	32 - 37	Calculation
RDW	14.9	H	%	12 - 14.5	Calculation
RDW-SD	36.3		fL		Calculation
MPV	6.5	L	fL	7.6 - 10.8	Calculation
PLATELET COUNT	290		10 <sup>3</sup> /uL	150 - 450	Electrical Impedance
PCT	0.2		%	0.01 - 9.99	Calculation
PDW	15.9		Not Applicable	0.1 - 99.9	Calculation
NUCLEATED RBC (NRBC) <sup>^</sup>	0.2		/100 WBC		VCS 360 Technology
ABSOLUTE NRBC COUNT <sup>^</sup>	0.02		10 <sup>3</sup> /uL		Calculation
EARLY GRANULOCYTE COUNT (EGC) <sup>^</sup>	0.2		%		VCS 360 Technology
ABSOLUTE EGC <sup>^</sup>	0.0		10 <sup>3</sup> /uL		Calculation
WBC COUNT	10.3		10 <sup>3</sup> /μL	4 - 11	Electrical Impedance
<b>DIFFERENTIAL COUNT (DC)</b>					
NEUTROPHILS	33		%	30 - 60	VCS 360 Technology
LYMPHOCYTES	62	H	%	30 - 60	VCS 360 Technology
EOSINOPHILS	0		%	0 - 6	VCS 360 Technology
MONOCYTES	5		%	1 - 6	VCS 360 Technology
BASOPHILS	0		%	0 - 1	VCS 360 Technology
<b>ABSOLUTE COUNT</b>					
ABSOLUTE NEUTROPHIL COUNT	3.4		10 <sup>3</sup> /uL	1.2 - 6.6	Calculation
ABSOLUTE LYMPHOCYTE COUNT	6.4		10 <sup>3</sup> /uL	1.2 - 6.6	Calculation
ABSOLUTE MONOCYTE COUNT	0.5		10 <sup>3</sup> /uL	0.04 - 0.66	Calculation
ABSOLUTE EOSINOPHIL COUNT	0.0		10 <sup>3</sup> /uL	0 - 0.66	Calculation
ABSOLUTE BASOPHIL COUNT	0.0		10 <sup>3</sup> /uL	0 - 0.11	Calculation

Comments : Please correlate clinically.

*Vyoma V. Shah*

**Dr. Adley Mark Fernandes**  
M.D (Pathology)  
Pathologist

**Dr. Vyoma V Shah**  
M.D (Pathology)  
Clinical Pathologist

*Thahsina Anees*

**Thahsina Anees**  
Laboratory Technologist

This is an electronically authenticated report

Page 2 of 3

Printed on: 07/10/2024 22:53

Test result pertains only to the sample tested and to be interpreted in the light of clinical history. These tests are accredited under ISO 15189:2012 unless specified by (^). Test marked with # is performed in an accredited referral laboratory.





**Laboratory Investigation Report**

**Name** : Mr. ORHAN AMIR ABBASI  
**DOB** : 14/03/2022  
**Age / Gender** : 2 Y / Male  
**Referred by** : DR AHSAN  
**Centre** : CITICARE MEDICAL CENTER

**Ref No.** : 37953  
**Sample No.** : 2410485351  
**Collected** : 07/10/2024 15:39  
**Registered** : 07/10/2024 21:53  
**Reported** : 07/10/2024 22:19

**HEMATOLOGY**

Test	Result	Flag	Unit	Reference Range	Methodology
------	--------	------	------	-----------------	-------------

COMPLETE BLOOD COUNT (CBC)

INTERPRETATION NOTES :

Please note update on CBC report format, reference ranges and method(Beckman Coulter).

Sample Type : EDTA Whole Blood

End of Report

**Dr. Adley Mark Fernandes**  
M.D (Pathology)  
Pathologist

*Vyoma V. Shah*  
**Dr. Vyoma V Shah**  
M.D (Pathology)  
Clinical Pathologist

*Thahsina*

**Thahsina Anees**  
Laboratory Technologist

This is an electronically authenticated report

Test result pertains only to the sample tested and to be interpreted in the light of clinical history. These tests are accredited under ISO 15189:2012 unless specified by (^). Test marked with # is performed in an accredited referral laboratory.

