

Patient Name : Mr.RAHUL RAJENDRA SHARMA	Collected : 12/Sep/2024 09:01AM
Age/Gender : 41 Y 4 M 27 D/M	Received : 12/Sep/2024 12:37PM
UHID/MR No : CWAN.0000137899	Reported : 12/Sep/2024 01:45PM
Visit ID : CWANOPV239720	Status : Final Report
Ref Doctor : Self	Sponsor Name : ARCOFEMI HEALTHCARE LIMITED
Emp/Auth/TPA ID : 9920144559	

DEPARTMENT OF HAEMATOLOGY

PERIPHERAL SMEAR , WHOLE BLOOD EDTA

**RBC's are Normocytic Normochromic,
WBC's Mild Eosinophilia
Platelets are Adequate
No Abnormal cells seen.**


Dr Sneha Shah
MBBS, MD (Pathology)
Consultant Pathologist

SIN No:CWA240900106

This test has been performed at Apollo Health and Lifestyle Ltd- Sadashiv Peth Pune, Diagnostics Lab



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DEPARTMENT OF HAEMATOLOGY

ARCOFEMI - MEDIWHEEL - PMC PACK D - PAN INDIA - FY2324

Test Name	Result	Unit	Bio. Ref. Interval	Method
HEMOGRAM , WHOLE BLOOD EDTA				
HAEMOGLOBIN	14.5	g/dL	13-17	Spectrophotometer
PCV	43.50	%	40-50	Electronic pulse & Calculation
RBC COUNT	4.77	Million/cu.mm	4.5-5.5	Electrical Impedance
MCV	91.3	fL	83-101	Calculated
MCH	30.4	pg	27-32	Calculated
MCHC	33.3	g/dL	31.5-34.5	Calculated
R.D.W	14.1	%	11.6-14	Calculated
TOTAL LEUCOCYTE COUNT (TLC)	7,420	cells/cu.mm	4000-10000	Electrical Impedance
DIFFERENTIAL LEUCOCYTIC COUNT (DLC)				
NEUTROPHILS	49	%	40-80	Electrical Impedance
LYMPHOCYTES	36.1	%	20-40	Electrical Impedance
EOSINOPHILS	7.2	%	1-6	Electrical Impedance
MONOCYTES	7.2	%	2-10	Electrical Impedance
BASOPHILS	0.5	%	<1-2	Electrical Impedance
ABSOLUTE LEUCOCYTE COUNT				
NEUTROPHILS	3635.8	Cells/cu.mm	2000-7000	Calculated
LYMPHOCYTES	2678.62	Cells/cu.mm	1000-3000	Calculated
EOSINOPHILS	534.24	Cells/cu.mm	20-500	Calculated
MONOCYTES	534.24	Cells/cu.mm	200-1000	Calculated
BASOPHILS	37.1	Cells/cu.mm	0-100	Calculated
Neutrophil lymphocyte ratio (NLR)	1.36		0.78- 3.53	Calculated
PLATELET COUNT	306000	cells/cu.mm	150000-410000	Electrical impedance
ERYTHROCYTE SEDIMENTATION RATE (ESR)	4	mm at the end of 1 hour	0-15	Modified Westergren
PERIPHERAL SMEAR				

RBC's are Normocytic Normochromic,
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DEPARTMENT OF HAEMATOLOGY

ARCOFEMI - MEDIWHEEL - PMC PACK D - PAN INDIA - FY2324

Test Name	Result	Unit	Bio. Ref. Interval	Method
BLOOD GROUP ABO AND RH FACTOR , WHOLE BLOOD EDTA				
BLOOD GROUP TYPE	B			Microplate Hemagglutination
Rh TYPE	Positive			Microplate Hemagglutination

Sneha Shah

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UHID/MR No : CWAN.0000137899	Reported : 12/Sep/2024 12:46PM
Visit ID : CWANOPV239720	Status : Final Report
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DEPARTMENT OF BIOCHEMISTRY

ARCOFEMI - MEDIWHEEL - PMC PACK D - PAN INDIA - FY2324

Test Name	Result	Unit	Bio. Ref. Interval	Method
GLUCOSE, FASTING , NAF PLASMA	85	mg/dL	70-100	HEXOKINASE

Comment:

As per American Diabetes Guidelines, 2023

Fasting Glucose Values in mg/dL	Interpretation
70-100 mg/dL	Normal
100-125 mg/dL	Prediabetes
≥126 mg/dL	Diabetes
<70 mg/dL	Hypoglycemia

Note:

- 1.The diagnosis of Diabetes requires a fasting plasma glucose of > or = 126 mg/dL and/or a random / 2 hr post glucose value of > or = 200 mg/dL on at least 2 occasions.
2. Very high glucose levels (>450 mg/dL in adults) may result in Diabetic Ketoacidosis & is considered critical.



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Consultant Pathologist

SIN No:CWA240900107

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Patient Name : Mr.RAHUL RAJENDRA SHARMA	Collected : 12/Sep/2024 09:01AM
Age/Gender : 41 Y 4 M 27 D/M	Received : 12/Sep/2024 12:25PM
UHID/MR No : CWAN.0000137899	Reported : 12/Sep/2024 12:55PM
Visit ID : CWANOPV239720	Status : Final Report
Ref Doctor : Self	Sponsor Name : ARCOFEMI HEALTHCARE LIMITED
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DEPARTMENT OF BIOCHEMISTRY

ARCOFEMI - MEDIWHEEL - PMC PACK D - PAN INDIA - FY2324

Test Name	Result	Unit	Bio. Ref. Interval	Method
ALANINE AMINOTRANSFERASE (ALT/SGPT), SERUM	24.65	U/L	<50	IFCC



DR.Sanjay Ingle
M.B.B.S,M.D(Pathology)
Consultant Pathologist

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
DEPARTMENT OF BIOCHEMISTRY

ARCOFEMI - MEDIWHEEL - PMC PACK D - PAN INDIA - FY2324

Test Name	Result	Unit	Bio. Ref. Interval	Method
TOTAL CHOLESTEROL , SERUM	242	mg/dL	<200	CHO-POD

Test Name	Result	Unit	Bio. Ref. Interval	Method
UREA , SERUM	16.45	mg/dL	17-43	GLDH, Kinetic Assay

Test Name	Result	Unit	Bio. Ref. Interval	Method
CREATININE , SERUM	0.69	mg/dL	0.72 – 1.18	Modified Jaffe, Kinetic

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DEPARTMENT OF CLINICAL PATHOLOGY

ARCOFEMI - MEDIWHEEL - PMC PACK D - PAN INDIA - FY2324

Test Name	Result	Unit	Bio. Ref. Interval	Method
COMPLETE URINE EXAMINATION (CUE) , URINE				
PHYSICAL EXAMINATION				
COLOUR	YELLOW		PALE YELLOW	Scattering of light
TRANSPARENCY	CLEAR		CLEAR	Scattering of light
pH	6.0		5-7.5	Bromothymol Blue
SP. GRAVITY	1.013		1.002-1.030	Bromothymol Blue
BIOCHEMICAL EXAMINATION				
URINE PROTEIN	NEGATIVE		NEGATIVE	PROTEIN ERROR OF INDICATOR
GLUCOSE	NORMAL		NEGATIVE	GOD-POD
URINE BILIRUBIN	NEGATIVE		NEGATIVE	Diazonium Salt
URINE KETONES (RANDOM)	NEGATIVE		NEGATIVE	Sodium nitro prusside
UROBILINOGEN	NORMAL		NORMAL (0.1-1.8mg/dl)	Diazonium salt
NITRITE	NEGATIVE		NEGATIVE	Sulfanilic acid
LEUCOCYTE ESTERASE	NEGATIVE		NEGATIVE	Diazonium salt
CENTRIFUGED SEDIMENT WET MOUNT AND MICROSCOPY				
PUS CELLS	1 - 2	/hpf	0-5	Microscopy
EPITHELIAL CELLS	1 - 2	/hpf	< 10	Microscopy
RBC	0	/hpf	0-2	Microscopy
CASTS	NEGATIVE	/lpf	0-2 Hyaline Cast	Microscopy
CRYSTALS	NEGATIVE	/hpf	Occasional-Few	Microscopy

Comment:

All urine samples are checked for adequacy and suitability before examination. All abnormal chemical examination are rechecked and verified by manual methods.

Microscopy findings are reported as an average of 10 high power fields.

*** End Of Report ***

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TERMS AND CONDITIONS GOVERNING THIS REPORT

The reported results are for information and interpretation of the referring doctor or such other medical professionals, who understand reporting units, reference ranges and limitations of technologies.


Laboratories not be responsible for any interpretation whatsoever.

It is presumed that the tests performed are, on the specimen / sample being to the patient named or identified and the verifications of the particulars have been cleared out by the patient or his / her representative at the point of generation of said specimen.

The reported results are restricted to the given specimen only. Results may vary from lab to lab and from time to time for the same parameter for the same patient.

Assays are performed in accordance with standard procedures, The reported results are dependent on individual assay methods / equipment used and quality of specimen received.

This report is not valid for medico legal purposes.


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