Name	: Mrs. SANDHYA PODDAR			
PID No.	: MED120919744	Register On	: 23/02/2023 9:40 AM	$\mathbf{C}$
SID No.	: 522302802	<b>Collection On</b>	: 23/02/2023 11:04 AM	medall
Age / Sex	: 53 Year(s) / Female	Report On	: 23/02/2023 3:33 PM	DIAGNOSTICS
Туре	: OP	Printed On	: 01/03/2023 1:49 PM	
Ref. Dr	: MediWheel			

Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> Reference Interval
<b>HAEMATOLOGY</b>			
<b>Complete Blood Count With - ESR</b>			
Haemoglobin (EDTA Blood/Spectrophotometry)	12.8	g/dL	12.5 - 16.0
Packed Cell Volume(PCV)/Haematocrit (EDTA Blood)	39.1	%	37 - 47
RBC Count (EDTA Blood)	4.65	mill/cu.mm	4.2 - 5.4
Mean Corpuscular Volume(MCV) (EDTA Blood)	84.0	fL	78 - 100
Mean Corpuscular Haemoglobin(MCH) (EDTA Blood)	27.5	pg	27 - 32
Mean Corpuscular Haemoglobin concentration(MCHC) (EDTA Blood)	32.7	g/dL	32 - 36
RDW-CV	13.1	%	11.5 - 16.0
RDW-SD	38.51	fL	39 - 46
Total Leukocyte Count (TC) (EDTA Blood)	6900	cells/cu.mm	4000 - 11000
Neutrophils (Blood)	63.4	%	40 - 75
Lymphocytes (Blood)	27.7	%	20 - 45
Eosinophils (Blood)	2.2	%	01 - 06
Monocytes (Blood)	6.4	%	01 - 10



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> <u>Reference Interval</u>
Basophils (Blood)	0.3	%	00 - 02
INTERPRETATION: Tests done on Automated Five I	Part cell counter. All	abnormal results are	reviewed and confirmed microscopically.
Absolute Neutrophil count (EDTA Blood)	4.37	10^3 / µl	1.5 - 6.6
Absolute Lymphocyte Count (EDTA Blood)	1.91	10^3 / µl	1.5 - 3.5
Absolute Eosinophil Count (AEC) (EDTA Blood)	0.15	10^3 / µl	0.04 - 0.44
Absolute Monocyte Count (EDTA Blood)	0.44	10^3 / µl	< 1.0
Absolute Basophil count (EDTA Blood)	0.02	10^3 / µl	< 0.2
Platelet Count (EDTA Blood)	153	10^3 / µl	150 - 450
MPV (Blood)	11.9	fL	8.0 - 13.3
PCT (Automated Blood cell Counter)	0.18	%	0.18 - 0.28
ESR (Erythrocyte Sedimentation Rate) (Citrated Blood)	44	mm/hr	< 30

Dr.Arjun C.P MBBS,MD Pathology Reg No:KMC \$9655 APPROVED BY

The results pertain to sample tested.

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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> <u>Reference Interval</u>
<b>BIOCHEMISTRY</b>			
Liver Function Test			
Bilirubin(Total) (Serum/DCA with ATCS)	1.23	mg/dL	0.1 - 1.2
Bilirubin(Direct) (Serum/Diazotized Sulfanilic Acid)	0.43	mg/dL	0.0 - 0.3
Bilirubin(Indirect) (Serum/Derived)	0.80	mg/dL	0.1 - 1.0
SGOT/AST (Aspartate Aminotransferase) (Serum/ <i>Modified IFCC</i> )	18.83	U/L	5 - 40
SGPT/ALT (Alanine Aminotransferase) (Serum/ <i>Modified IFCC</i> )	17.30	U/L	5 - 41
GGT(Gamma Glutamyl Transpeptidase) (Serum/IFCC / Kinetic)	15.95	U/L	< 38
Alkaline Phosphatase (SAP) (Serum/ <i>Modified IFCC</i> )	105.5	U/L	53 - 141
Total Protein (Serum/Biuret)	7.54	gm/dl	6.0 - 8.0
Albumin (Serum/Bromocresol green)	4.72	gm/dl	3.5 - 5.2
Globulin (Serum/Derived)	2.82	gm/dL	2.3 - 3.6
A : G RATIO	1.67		1.1 - 2.2

(Serum/Derived)



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> Reference Interval
<u>Lipid Profile</u>			
Cholesterol Total (Serum/CHOD-PAP with ATCS)	133.13	mg/dL	Optimal: < 200 Borderline: 200 - 239 High Risk: >= 240
Triglycerides (Serum/ <i>GPO-PAP with ATCS</i> )	125.48	mg/dL	Optimal: < 150 Borderline: 150 - 199 High: 200 - 499 Very High: >= 500

**INTERPRETATION:** The reference ranges are based on fasting condition. Triglyceride levels change drastically in response to food, increasing as much as 5 to 10 times the fasting levels, just a few hours after eating. Fasting triglyceride levels show considerable diurnal variation too. There is evidence recommending triglycerides estimation in non-fasting condition for evaluating the risk of heart disease and screening for metabolic syndrome, as non-fasting sample is more representative of the `usual\_circulating level of triglycerides during most part of the day.

HDL Cholesterol (Serum/Immunoinhibition)	55.43	mg/dL	Optimal(Negative Risk Factor): >= 60 Borderline: 50 - 59 High Risk: < 50
LDL Cholesterol (Serum/ <i>Calculated</i> )	52.6	mg/dL	Optimal: < 100 Above Optimal: 100 - 129 Borderline: 130 - 159 High: 160 - 189 Very High: >=190
VLDL Cholesterol (Serum/Calculated)	25.1	mg/dL	< 30
Non HDL Cholesterol (Serum/ <i>Calculated</i> )	77.7	mg/dL	Optimal: < 130 Above Optimal: 130 - 159 Borderline High: 160 - 189 High: 190 - 219 Very High: >=220



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> <u>Reference Interval</u>
<b>INTERPRETATION:</b> 1.Non-HDL Cholesterol is nov 2.It is the sum of all potentially atherogenic proteins in co-primary target for cholesterol lowering therapy.	1		
Total Cholesterol/HDL Cholesterol Ratio (Serum/ <i>Calculated</i> )	2.4		Optimal: < 3.3 Low Risk: 3.4 - 4.4 Average Risk: 4.5 - 7.1 Moderate Risk: 7.2 - 11.0 High Risk: > 11.0
Triglyceride/HDL Cholesterol Ratio (TG/HDL) (Serum/Calculated)	2.3		Optimal: < 2.5 Mild to moderate risk: 2.5 - 5.0 High Risk: > 5.0
LDL/HDL Cholesterol Ratio (Serum/ <i>Calculated</i> )	0.9		Optimal: 0.5 - 3.0 Borderline: 3.1 - 6.0 High Risk: > 6.0



APPROVED BY

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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	Biological Reference Interval
<u>Glycosylated Haemoglobin (HbA1c)</u>			
HbA1C (Whole Blood/ <i>HPLC</i> )	6.3	%	Normal: 4.5 - 5.6 Prediabetes: 5.7 - 6.4 Diabetic: >= 6.5
INTEDDDETATION If Diskates Cood control (	1 70 % Esir control	.71 800 Dec	$\sim 10^{-10}$

**INTERPRETATION:** If Diabetes - Good control : 6.1 - 7.0 %, Fair control : 7.1 - 8.0 %, Poor control >= 8.1 %

(Whole Blood)

#### **INTERPRETATION:** Comments

HbA1c provides an index of Average Blood Glucose levels over the past 8 - 12 weeks and is a much better indicator of long term glycemic control as compared to blood and urinary glucose determinations.

Conditions that prolong RBC life span like Iron deficiency anemia, Vitamin B12 & Folate deficiency,

hypertriglyceridemia, hyperbilirubinemia, Drugs, Alcohol, Lead Poisoning, Asplenia can give falsely elevated HbA1C values.

Conditions that shorten RBC survival like acute or chronic blood loss, hemolytic anemia, Hemoglobinopathies, Splenomegaly, Vitamin E ingestion, Pregnancy, End stage Renal disease can cause falsely low HbA1c.



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> Reference Interval
<b>IMMUNOASSAY</b>			
<u>THYROID PROFILE / TFT</u>			
T3 (Triiodothyronine) - Total (Serum/ <i>ECLIA</i> )	1.30	ng/ml	0.4 - 1.81
<b>INTERPRETATION:</b> <b>Comment :</b> Total T3 variation can be seen in other condition like preg Metabolically active.	gnancy, drugs, nepł	rosis etc. In such cas	es, Free T3 is recommended as it is
T4 (Tyroxine) - Total (Serum/ <i>ECLIA</i> )	8.85	µg/dl	4.2 - 12.0
<b>INTERPRETATION:</b> <b>Comment :</b> Total T4 variation can be seen in other condition like pres Metabolically active.	gnancy, drugs, nepł	rrosis etc. In such cas	es, Free T4 is recommended as it is
TSH (Thyroid Stimulating Hormone) (Serum/ECLIA)	4.96	µIU/mL	0.35 - 5.50
INTERPRETATION: Reference range for cord blood - upto 20 1 st trimester: 0.1-2.5 2 nd trimester 0.2-3.0 3 rd trimester : 0.3-3.0 (Indian Thyroid Society Guidelines) Comment : 1.TSH reference range during pregnancy depends on Iodi	ine intake, TPO stat	rus, Serum HCG con	centration, race, Ethnicity and BMI.
2.TSH Levels are subject to circadian variation, reaching of the order of 50%,hence time of the day has influence of 3 Values& amplt0 03 uII /mL need to be clinically correl	peak levels betwee on the measured ser	n 2-4am and at a mir um TSH concentratio	nimum between 6-10PM. The variation can be ons.

3. Values&amplt,0.03 µIU/mL need to be clinically correlated due to presence of rare TSH variant in some individuals.



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> <u>Reference Interval</u>
<b>CLINICAL PATHOLOGY</b>			
<u>PHYSICAL EXAMINATION (URINE</u> <u>COMPLETE)</u>			
Colour (Urine)	Pale yellow		Yellow to Amber
Appearance (Urine)	Clear		Clear
Volume(CLU) (Urine)	10		
<u>CHEMICAL EXAMINATION (URINE</u> <u>COMPLETE)</u>			
pH (Urine)	5.0		4.5 - 8.0
Specific Gravity (Urine)	1.003		1.002 - 1.035
Ketone (Urine)	Negative		Negative
Urobilinogen (Urine)	Normal		Normal
Blood (Urine)	Negative		Negative
Nitrite (Urine)	Negative		Negative
Bilirubin (Urine)	Negative		Negative
Protein (Urine)	Negative		Negative



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	Biological Reference Interval
Glucose (Urine/GOD - POD)	Negative		Negative
Leukocytes(CP) (Urine)	Negative		Negative
<u>MICROSCOPIC EXAMINATION</u> (URINE COMPLETE)			
Pus Cells (Urine)	0-1	/hpf	NIL
Epithelial Cells (Urine)	0-1	/hpf	NIL
RBCs (Urine)	NIL	/HPF	NIL
Others (Urine)	NIL		

**INTERPRETATION:** Note: Done with Automated Urine Analyser & Automated urine sedimentation analyser. All abnormal reports are reviewed and confirmed microscopically.

mishall Dr Anusha.K.S Sr.Consultant Pathologist Reg No : 100674 APPROVED BY

The results pertain to sample tested.

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Investigation

<u>Observed</u> <u>Value</u>

<u>Unit</u>

Biological Reference Interval

# **IMMUNOHAEMATOLOGY**

BLOOD GROUPING AND Rh TYPING (EDTA Blood/Agglutination)

'B' 'Positive'

**INTERPRETATION:** Note: Slide method is screening method. Kindly confirm with Tube method for transfusion.



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Investigation	<u>Observed</u> <u>Value</u>	<u>Unit</u>	<u>Biological</u> Reference Interval
<b>BIOCHEMISTRY</b>			
BUN / Creatinine Ratio	10		6.0 - 22.0
Glucose Fasting (FBS) (Plasma - F/GOD-PAP)	113.31	mg/dL	Normal: < 100 Pre Diabetic: 100 - 125 Diabetic: >= 126

**INTERPRETATION:** Factors such as type, quantity and time of food intake, Physical activity, Psychological stress, and drugs can influence blood glucose level.

Glucose, Fasting (Urine)	Negative	Negative
(Urine - F/GOD - POD)		
Glucose Postprandial (PPBS)	121.85 mg/	dL 70 - 140
(Plasma - PP/GOD-PAP)		

#### **INTERPRETATION:**

Factors such as type, quantity and time of food intake, Physical activity, Psychological stress, and drugs can influence blood glucose level. Fasting blood glucose level may be higher than Postprandial glucose, because of physiological surge in Postprandial Insulin secretion, Insulin resistance, Exercise or Stress, Dawn Phenomenon, Somogyi Phenomenon, Anti- diabetic medication during treatment for Diabetes.

Urine Glucose(PP-2 hours) (Urine - PP)	Negative	Negative
Blood Urea Nitrogen (BUN) (Serum/Urease UV / derived)	7.8 mg/dL	7.0 - 21
Creatinine	0.78 mg/dL	0.6 - 1.1

## (Serum/Modified Jaffe)

**INTERPRETATION:** Elevated Creatinine values are encountered in increased muscle mass, severe dehydration, Pre-eclampsia, increased ingestion of cooked meat, consuming Protein/ Creatine supplements, Diabetic Ketoacidosis, prolonged fasting, renal dysfunction and drugs such as cefoxitin ,cefazolin, ACE inhibitors ,angiotensin II receptor antagonists,N-acetylcyteine , chemotherapeutic agent such as flucytosine etc.

Uric Acid	5.51	mg/dL	2.6 - 6.0
(Serum/Enzymatic)			



-- End of Report --



Name	MRS.SANDHYA PODDAR	ID	MED120919744
Age & Gender	53/FEMALE	Visit Date	23/02/2023
Ref Doctor Name	MediWheel		

X-ray mammogram (mediolateral oblique & craniocaudal views) followed by Sonomammography.

### **BILATERAL MAMMOGRAPHY**

Breast composition Type B (These are scattered areas of fibroglandular density).

No evidence of focal soft tissue lesion.

No evidence of cluster microcalcification.

Subcutaneous fat deposition is within normal limits.

#### **BILATERAL SONOMAMMOGRAPHY**

Both the breasts show normal echopattern.

No evidence of focal solid / cystic areas.

No evidence of ductal dilatation.

Breast parenchyma is seen in the bilateral axilla - axillary tail.

Bilateral axillary lymphnodes are noted with preserved fatty hilum.

#### **IMPRESSION:**

- No breast lesion.
- Bilateral benign axillary lymphnodes. •

#### **ASSESSMENT: BI-RADS CATEGORY -1**

#### **BI-RADS CLASSIFICATION**

#### CATEGORY RESULT

2

Benign finding. Routine mammogram in 1 year recommended.

#### REPORT DISCLAIMER 7.Results of the test are influenced by the various factors such as sensitivity, specificity of the

- 1. This is only a radiologincal imperssion. Like other investigations, radiological investication also have limitation. Therefore radiologincal reports should be interpreted in correlation with clinical and pathological findings
- 2. The results reported here in are subject to interpretation by qualified medical professionals only.
- 3.Customer identities are accepted provided by the customer or their representative. 4.information about the customer's condition at the time of sample collection such as fasting, food
- consumption, medication, etc are accepted as provided by the customer or representative and shall not be investigated for its truthfulness.

- procedures of the tests, quality of the samples and drug interactions etc., 8.If the test results are found not to be correlating clinically can contact the lab in charge for clarification or retesting where practicable within 24 hours from the time of issue of results. 9.Liability is limited to the extend of amount billed

10.Reports are subject to interpretation in their entirety.partial or selective interpretation may lead to false opinion.

11.Disputes, if any , with regard to the report findings are subject to the exclusive jurisdiction of the competent courts chennai only

<sup>5.</sup>If any specimen/sample is received from any others laboratory/hospital,its is presumed that the sample belongs to the patient identified or named.

<sup>6.</sup>Test results should be interpreted in context of clinical and other findings if any.In case of any clarification /doubt , the refrering doctor/patient can contact the respective section head of the laboratory.



Name	MRS.SANDHYA PODDAR	ID	MED120919744
Age & Gender	53/FEMALE	Visit Date	23/02/2023
Ref Doctor Name	MediWheel		

#### DR. HEMANANDINI V.N CONSULTANT RADIOLOGIST Hn/

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