

: 2202102816 Reg. No

Name : Diksha Methi Age/Sex : 28 Years / Female

Ref. By

Client : MEDIWHEEL WELLNESS Reg. Date

: 27-Feb-2022

Collected On : 27-Feb-2022 09:45

Approved On : 27-Feb-2022 11:19

Printed On : 08-Mar-2022 14:58

<u>Parameter</u>	<u>Result</u>	<u>Unit</u>	Reference Interval	
	KIDNEY FL	JNCTION TEST		
UREA (Urease & glutamate dehydrogenase)	18.4	mg/dL	10 - 50	
Creatinine (Jaffe method)	1.05	mg/dL	0.5 - 1.2	
Uric Acid (Enzymatic colorimetric)	3.3	mg/dL	2.5 - 7.0	
End Of Report				

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Parameter

Hemoglobin

RBC Count

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<u>Result</u>	<u>Unit</u>	Reference Interval	
	E BLOOD COUN CIMEN: EDTA BLOOD	•	
13.1	g/dL	12.0 - 15.0	

3.8 - 4.8

million/cmm

TO COUNT			0.0
Hematrocrit (PCV)	38.0	%	40 - 54
MCH	30.5	Pg	27 - 32
MCV	88.6	fL	83 - 101
MCHC	34.5	%	31.5 - 34.5
RDW	11.0	%	11.5 - 14.5
WBC Count	7300	/cmm	4000 - 11000
DIFFERENTIAL WBC COUNT (Flow of	cytometry)		
Neutrophils (%)	60	%	38 - 70
Lymphocytes (%)	30	%	20 - 40
Monocytes (%)	07	%	2 - 8
Eosinophils (%)	03	%	0 - 6
Basophils (%)	00	%	0 - 2
Neutrophils	4380	/cmm	
Lymphocytes	2190	/cmm	
Monocytes	511	/cmm	
Eosinophils	219	/cmm	
Basophils	0	/cmm	
Platelet Count (Flow cytometry)	59000	/cmm	150000 - 450000
MPV	11.7	fL	7.5 - 11.5
ERYTHROCYTE SEDIMENTATION R	<u>ATE</u>		
ESR (After 1 hour)	08	mm/hr	0 - 21

4.29

----- End Of Report -----

MD Pathologist

Modified Westergren Method

		TEST REPORT		
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Client	: MEDIWHEEL WELLNESS			
Paramet	<u>er</u>	Result		
	Specime	BLOOD GROUP & RH n: EDTA and Serum; Method: Haemagg	utination	
ABO		'O'		
Rh (D)		Positive		
		End Of Report		



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Age/Sex

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<u>Unit</u> Reference Interval Result

PLASMA GLUCOSE

Fasting Blood Sugar (FBS) 85.9 mg/dL 70 - 110

Hexokinase Method

70 - 140 Post Prandial Blood Sugar (PPBS) 105.5 mg/dL

Hexokinase Method

Criteria for the diagnosis of diabetes1. HbA1c >/= 6.5 *

2. Fasting plasma glucose >126 gm/dL. Fasting is defined as no caloric intake at least for 8 hrs.

3. Two hour plasma glucose >/= 200mg/dL during an oral glucose tolerence test by using a glucose load containing equivalent of 75 gm anhydrous glucose dissolved in water.

4. In a patient with classic symptoms of hyperglycemia or hyperglycemic crisis, a random plasma glucose >/= 200 mg/dL.

*In the absence of unequivocal hyperglycemia, criteria 1-3 should be confirmed by repeat testing.

American diabetes association. Standards of medical care in diabetes 2011. Diabetes care 2011;34;S11.

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<u>Parameter</u>	Result	<u>Unit</u>	Reference Interval		
LIPID PROFILE					
Cholesterol (Enzymatic colorimetric)	189.1	mg/dL	Desirable : < 200.0 Borderline High : 200-239 High : > 240.0		
Triglyceride (Enzymatic colorimetric)	67.3	mg/dL	Normal : < 150.0 Borderline : 150-199 High : 200-499 Very High : > 500.0		
VLDL	13.46	mg/dL	15 - 35		
Calculated					
LDL CHOLESTEROL	121.34	mg/dL	Optimal: < 100.0 Near / above optimal: 100-129 Borderline High: 130-159 High: 160-189 Very High: >190.0		
HDL Cholesterol	54.3	mg/dL	30 - 85		
Homogeneous enzymatic colorimetri	С				
Cholesterol /HDL Ratio Calculated	3.48		0 - 5.0		
LDL / HDL RATIO Calculated	2.23		0 - 3.5		



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Parameter Result <u>Unit</u> Reference Interval

NEW ATP III GUIDELINES (MAY 2001), MODIFICATION OF NCEP<?xml:namespace prefix = "o" ns = "urn:schemasmicrosoft-com:office:office" />

> LDL CHOLESTEROL **CHOLESTEROL HDL CHOLESTEROL TRIGLYCERIDES** Optimal<100 Desirable<200 Low<40 Normal<150 Near Optimal 100-129 Border Line 200-239

High >60 Border High 150-199 Borderline 130-159 High >240

> High 200-499 High 160-189

- LDL Cholesterol level is primary goal for treatment and varies with risk category and assesment
- For LDL Cholesterol level Please consider direct LDL value

Risk assessment from HDL and Triglyceride has been revised. Also LDL goals have changed.

- Detail test interpreation available from the lab
- All tests are done according to NCEP guidelines and with FDA approved kits.
- LDL Cholesterol level is primary goal for treatment and varies with risk category and assesment

For test performed on specimens received or collected from non-KSHIPRA locations, it is presumed that the specimen belongs to the patient named or identified as labeled on the container/test request and such verification has been carried out at the point generation of the said specimen by the sender.

KSHIPRA will be responsible Only for the analytical part of test carried out. All other responsibility will be of referring Laboratory.

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<u>Parameter</u>	Result	<u>Unit</u>	Reference Interval		
LIVER FUNCTION TEST					
Total Bilirubin	0.44	mg/dL	0.20 - 1.0		
Colorimetric diazo method					
Conjugated Bilirubin	0.28	mg/dL	0.0 - 0.3		
Sulph acid dpl/caff-benz					
Unconjugated Bilirubin	0.16	mg/dL	0.0 - 1.1		
Sulph acid dpl/caff-benz					
SGOT	33.3	U/L	0 - 31		
(Enzymatic)					
SGPT	33.8	U/L	0 - 31		
(Enzymatic)					
Alakaline Phosphatase	96.9	U/L	42 - 141		
(Colorimetric standardized method)					
Protien with ratio					
Total Protein	8.7	g/dL	6.5 - 8.7		
(Colorimetric standardized method)					
Albumin	5.1	mg/dL	3.5 - 4.94		
(Colorimetric standardized method)					
Globulin	3.60	g/dL	2.3 - 3.5		
Calculated					
A/G Ratio	1.42		0.8 - 2.0		
Calculated					

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MD Pathologist

TEST REPORT

Reg. No : 2202102816 Name : Diksha Methi

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<u>Parameter</u> <u>Result</u> <u>Unit</u> <u>Reference Interval</u>

HEMOGLOBIN A1 C ESTIMATION

Specimen: Blood EDTA

Hb A1C 5.4

Boronate Affinity with Fluorescent Quenching

% of Total Hb

Poor Control: > 7.0 % Good Control: 6.2-7.0 % Non-diabetic Level: 4.3-6.2 %

Mean Blood Glucose 114.94 mg/dL

Calculated

Degree of Glucose Control Normal Range:

Poor Control >7.0% *

Good Control 6.0 - 7.0 %**Non-diabetic level < 6.0 %

- * High risk of developing long term complication such as retinopathy, nephropathy, neuropathy, cardiopathy, etc.
- * Some danger of hypoglycemic reaction in Type I diabetics.
- * Some glucose intolerant individuals and "subclinical" diabetics may demonstrate HbA1c levels in this area.

EXPLANATION:-

*Total haemoglobin A1 c is continuously symthesised in the red blood cell throught its 120 days life span. The concentration of HBA1c in the cell reflects the average blood glucose concentration it encounters.

*The level of HBA1c increases proportionately in patients with uncontrolled diabetes. It reflects the average blood glucose oncentration over an extended time period and remains unaffected by short-term fluctuations in blood glucose levels.

*The measurement of HbA1c can serve as a convenient test for evaluating the adequacy of diabetic control and in preventing various diabetic complications. Because the average half life of a red blood cell is sixty days, HbA1c has been accepted as a measurnment which effects the mean daily blood glucose concentration, better than fasting blood glucose determination, and the degree of carbohydrate imbalance over the preceding two months.

*It may also provide a better index of control of the diabetic patient without resorting to glucose loading procedures.

HbA1c assay Interferences:

*Errneous values might be obtained from samples with abnormally elevated quantities of other Haemoglobins as a result of either their simultaneous elution with HbA1c(HbF) or differences in their glycation from that of HbA(HbS)

----- End Of Report -----

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Test done from collected sample

Approved by: DR PS RAO MD Pathologist



Reg. No : 2202102816
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: MEDIWHEEL WELLNESS

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: 27-Feb-2022

<u>Parameter</u>	Result	<u>Unit</u>	Reference Interval		
THYROID FUNCTION TEST					
T3 (Triiodothyronine)	1.07	ng/mL	0.87 - 1.78		
Chemiluminescence					
T4 (Thyroxine)	10.40	μg/dL	5.89 - 14.9		
Chemiluminescence					
TSH (ultra sensitive)	2.980	μIU/ml	0.34 - 5.6		
Chemiluminescence					

SUMMARY The hypophyseal release of TSH (thyrotropic hormone) is the central regulating mechanism for the biological action of thyroid hormones. TSH is a very sensitive and specific parameter for assessing thyroid function and is particularly suitable for early detection or exclusion of disorders in the central regulating circuit between the hypothalamus, pituitary and thyroid. LIMITATION Presence of autoantibodies may cause unexpected high value of TSH

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URINE ROUTINE EXAMINATION

PHYSICAL EXAMINATION

Quantity 20 cc

Pale Yellow Colour

Clear **Appearance**

CHEMICAL EXAMINATION (BY REFLECTANCE PHOTOMETRIC METHOD)

Result

5.0 - 8.0рΗ 7.0 1.020 1.002 - 1.03 Sp. Gravity

Nil Protein Glucose Nil **Ketone Bodies** Nil Urine Bile salt and Bile Pigment Nil Urine Bilirubin Nil Nitrite Nil Leucocytes Nil Blood Nil

MICROSCOPIC EXAMINATION (MANUAL BY MCIROSCOPY)

Nil

Leucocytes (Pus Cells) Nil Erythrocytes (Red Cells) Nil **Epithelial Cells** 1-2/hpf **Amorphous Material** Nil Casts Nil Nil Crystals **Bacteria** Nil

----- End Of Report -----

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DR PS RAO MD Pathologist