



CLIENT CODE: C000138354 **CLIENT'S NAME AND ADDRESS:**

ACROFEMI HEALTHCARE LTD (MEDIWHEEL) F-703, LADO SARAI, MEHRAULI SOUTH WEST DELHI

NEW DELHI 110030 DELHI INDIA 8800465156

SRL Ltd

Shop CG 017, PALM SPRINGS PLAZA GURUGRAM, 122001

HARYANA, INDIA Tel: 9111591115

PATIENT NAME: NISHAT SAINI PATIENT ID: NISHM250885282

ACCESSION NO : **0282WD00001** AGE: 37 Years SEX: Male ABHA NO:

RECEIVED: 01/04/2023 08:23 REPORTED: 03/04/2023 11:44 DRAWN:

REFERRING DOCTOR: SELF CLIENT PATIENT ID:

Test Report Status Results **Biological Reference Interval Units** <u>Final</u>

MEDI WHEEL FULL BODY HEALTH CHECK UP BELOW 40 MALE

BLOOD COUNTS,EDTA WHOLE BLOOD				
HEMOGLOBIN (HB)	12.5	Low	13.0 - 17.0	g/dL
METHOD: SPECTROPHOTOMETRY				
RED BLOOD CELL (RBC) COUNT	6.16	High	4.5 - 5.5	mil/μL
METHOD: IMPEDANCE				
WHITE BLOOD CELL (WBC) COUNT	7.51		4.0 - 10.0	thou/µL
METHOD: IMPEDANCE				
PLATELET COUNT	150		150 - 410	thou/µL
METHOD: IMPEDANCE				
RBC AND PLATELET INDICES				
HEMATOCRIT (PCV)	37.7	Low	40 - 50	%
METHOD: CALCULATED				
MEAN CORPUSCULAR VOLUME (MCV)	61.1	Low	83 - 101	fL
METHOD : DERIVED FROM IMPEDANCE MEASURE				
MEAN CORPUSCULAR HEMOGLOBIN (MCH)	20.3	Low	27.0 - 32.0	pg
METHOD : CALCULATED PARAMETER				
MEAN CORPUSCULAR HEMOGLOBIN CONCENTRATION (MCHC) METHOD: CALCULATED PARAMETER	33.3		31.5 - 34.5	g/dL
RED CELL DISTRIBUTION WIDTH (RDW)	18.5	High	11.6 - 14.0	%
METHOD: DERIVED FROM IMPEDANCE MEASURE				
MENTZER INDEX	9.9			
MEAN PLATELET VOLUME (MPV)	8.9		6.8 - 10.9	fL
METHOD: DERIVED FROM IMPEDANCE MEASURE				
WBC DIFFERENTIAL COUNT				
NEUTROPHILS	49		40 - 80	%
METHOD: DHSS FLOWCYTOMETRY				
LYMPHOCYTES	41	High	20 - 40	%
METHOD: DHSS FLOWCYTOMETRY				
MONOCYTES	8		2 - 10	%
METHOD: DHSS FLOWCYTOMETRY				
EOSINOPHILS	2		1 - 6	%
METHOD: DHSS FLOWCYTOMETRY				
BASOPHILS	0		0 - 2	%
METHOD: IMPEDANCE				
ABSOLUTE NEUTROPHIL COUNT	3.69		2.0 - 7.0	thou/µL









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METHOD: DHSS FLOWCYTOME	ETDY CALCULATED				
ABSOLUTE LYMPHOCYTE	•	3.07	High	1 - 3	thou/µL
METHOD : DHSS FLOWCYTOME		3.07	iligii	1 - 3	tilou/μL
ABSOLUTE MONOCYTE O	•	0.58		0.20 - 1.00	thou/µL
METHOD : DHSS FLOWCYTOME		0.36		0.20 - 1.00	tilou/μL
ABSOLUTE EOSINOPHIL	•	0.12		0.02 - 0.50	thou/µL
		0.12		0.02 - 0.30	tilou/μL
METHOD: DHSS FLOWCYTOME ABSOLUTE BASOPHIL CO	·	0.02		0.02 - 0.10	thou/µL
		0.02		0.02 - 0.10	tilou/μL
METHOD: DHSS FLOWCYTOME	•	1.2			
NEUTROPHIL LYMPHOCY	IE RAIIO (NLR)	1.2			
METHOD : CALCULATED	ENTATION DATE (EC	D) WHOLE			
ERYTHROCYTE SEDIM BLOOD	ENTATION KATE (ES	K),WHOLE			
E.S.R		3		0 - 14	mm at 1 hr
METHOD : AUTOMATED (PHOTO	OMETRICAL CAPILLARY STOPPE	D FLOW KINETIC ANALYSIS)			
GLUCOSE FASTING,FL	UORIDE PLASMA				
FBS (FASTING BLOOD S	UGAR)	95		Normal 75 - 99 Pre-diabetics: 100 - 125 Diabetic: > or = 126	mg/dL
METHOD: SPECTROPHOTOMET	TRY HEXOKINASE				
GLYCOSYLATED HEMO BLOOD	GLOBIN(HBA1C), ED	TA WHOLE			
HBA1C		4.9		Non-diabetic: < 5.7 Pre-diabetics: 5.7 - 6.4 Diabetics: > or = 6.5 ADA Target: 7.0 Action suggested: > 8.0	%
METHOD : CAPILLARY ELECTRO		00.0			
ESTIMATED AVERAGE G	LUCOSE(EAG)	93.9		< 116	mg/dL



METHOD: CALCULATED PARAMETER







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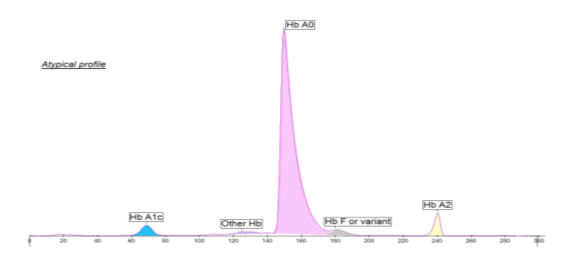
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ID: 28214764368 Sample num. 10 Date 02/04/2023

Birth: Depart:



A1c Haemoglobin Electrophoresis

Fractions	%	mmol/mol	Cal. %	
Hb A1c (*)	-	30	4.9	
Other Hb	1.7			
Hb A0	88.1			
Hb F or variant	2.1			
Hb A2 (!)	4.4			

HbA1c % cal :(*) = 4.9 %

HbA1c mmol/mol :(*) = 30 mmol/mol

(*) Atypical profile

GLUCOSE, POST-PRANDIAL, PLASMA

PPBS(POST PRANDIAL BLOOD SUGAR) SAMPLE NOT RECEIVED 70 - 139 mg/dL





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Test Report Status	<u>Final</u>	Results		Biological Reference Interv	al Units
METHOD : SPECTROPHOTOME	ETRY, HEXOKINASE				
LIPID PROFILE, SERU	JM				
CHOLESTEROL, TOTAL	DIMETERS ASSAY	133		Desirable : < 200 Borderline : 200 - 239 High : > / = 240	mg/dL
METHOD : ENZYMATIC COLOR	RIMETRIC ASSAY	120		Name	
TRIGLYCERIDES		130		Normal: < 150 Borderline high: 150 - 199 High: 200 - 499 Very High: >/= 500	mg/dL
METHOD : ENZYMATIC COLOR	RIMETRIC ASSAY				
HDL CHOLESTEROL		33	Low	At Risk: < 40 Desirable: > or = 60	mg/dL
METHOD: HOMOGENEOUS E	NZYMATIC COLORIMETRIC ASSAY				
CHOLESTEROL LDL		81		Adult levels: Optimal < 100 Near optimal/above optimal: 129 Borderline high: 130-159 High: 160-189 Very high: = 190	mg/dL 100-
METHOD: HOMOGENEOUS E	NZYMATIC COLORIMETRIC ASSAY				
NON HDL CHOLESTERO	L	100		Desirable: < 130 Above Desirable: 130 -159 Borderline High: 160 - 189 High: 190 - 219 Very high: > / = 220	mg/dL
METHOD : CALCULATED PARA	AMETER				
VERY LOW DENSITY LIF	POPROTEIN	26.0		< OR = 30.0	mg/dL
METHOD : CALCULATED PARA	AMETER				
CHOL/HDL RATIO		4.0		Low Risk: 3.3 - 4.4 Average Risk: 4.5 - 7.0 Moderate Risk: 7.1 - 11.0 High Risk: > 11.0	
METHOD : CALCULATED PARA	AMETER				
LDL/HDL RATIO		2.5		0.5 - 3.0 Desirable/Low Risk 3.1 - 6.0 Borderline/Moderate >6.0 High Risk	Risk
METHOD : CALCULATED PARA	AMETER			-	









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Test Report Status Results Biological Reference Interval Units <u>Final</u>

Interpretation(s)

- 1) Cholesterol levels help assess the patient risk status and to follow the progress of patient under treatment to lower serum cholesterol concentrations.
- 2) Serum Triglyceride (TG) are a type of fat and a major source of energy for the body. Both quantity and composition of the diet impact on plasma triglyceride concentrations. Elevations in TG levels are the result of overproduction and impaired clearance. High TG are associated with increased risk for CAD (Coronary artery disease) in patients with other risk factors, such as low HDL-C, some patient groups with elevated apolipoprotein B concentrations, and patients with forms of LDL that may be particularly atherogenic.
- 3)HDL-C plays a crucial role in the initial step of reverse cholesterol transport, this considered to be the primary atheroprotective function of HDL
- 4) LDL -C plays a key role in causing and influencing the progression of atherosclerosis and, in particular, coronary sclerosis. The majority of cholesterol stored in atherosclerotic plaques originates from LDL, thus LDL-C value is the most powerful clinical predictor.
- 5)Non HDL cholesterol: Non-HDL-C measures the cholesterol content of all atherogenic lipoproteins, including LDL hence it is a better marker of risk in both primary and secondary prevention studies. Non-HDL-C also covers, to some extent, the excess ASCVD risk imparted by the sdLDL, which is significantly more atherogenic than the normal large buoyant particles, an elevated non-HDL-C indirectly suggests greater proportion of the small, dense variety of LDL particles

Serum lipid profile is measured for cardiovascular risk prediction. Lipid Association of India recommends LDL-C as primary target and Non HDL-C as co-primary treatment target.

Risk Stratification for ASCVD (Atherosclerotic cardiovascular disease) by Lipid Association of India

Risk Category		
Extreme risk group	A.CAD with > 1 feature of high risk group	
	B. CAD with > 1 feature of Very high risk g	group or recurrent ACS (within 1 year) despite LDL-C
	< or = 50 mg/dl or polyvascular disease	
Very High Risk	1. Established ASCVD 2. Diabetes with 2 1	najor risk factors or evidence of end organ damage 3.
	Familial Homozygous Hypercholesterolemi	a
High Risk	1. Three major ASCVD risk factors. 2. Dia	betes with 1 major risk factor or no evidence of end
		DL >190 mg/dl 5. Extreme of a single risk factor. 6.
	Coronary Artery Calcium - CAC >300 AU.	7. Lipoprotein a >/= 50mg/dl 8. Non stenotic carotid
	plaque	
Moderate Risk	2 major ASCVD risk factors	
Low Risk	0-1 major ASCVD risk factors	
Major ASCVD (Ath	erosclerotic cardiovascular disease) Risk Fa	ctors
1. Age $>$ or $=$ 45 year	s in males and $>$ or $= 55$ years in females	3. Current Cigarette smoking or tobacco use
2. Family history of p	remature ASCVD	4. High blood pressure
5. Low HDL		

Newer treatment goals and statin initiation thresholds based on the risk categories proposed by LAI in 2020.

Risk Group	Treatment Goals		Consider Drug Thera	py
	LDL-C (mg/dl)	Non-HDL (mg/dl)	LDL-C (mg/dl)	Non-HDL (mg/dl)
Extreme Risk Group	<50 (Optional goal	< 80 (Optional goal	>OR = 50	>OR = 80
Category A	< OR = 30)	<OR = 60)		









Units

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Extreme Risk Group	<or 30<="" =="" td=""><td><or 60<="" =="" td=""><td>> 30</td><td>>60</td></or></td></or>	<or 60<="" =="" td=""><td>> 30</td><td>>60</td></or>	> 30	>60
Category B				
Very High Risk	<50	<80	>OR= 50	>OR= 80
High Rick	<70	<100	>OR= 70	>OR= 100

Results

Entreme rusk Group	1010	-010	. 50	. 00
Category B				
Very High Risk	<50	<80	>OR= 50	>OR= 80
High Risk	<70	<100	>OR= 70	>OR= 100
Moderate Risk	<100	<130	>OR= 100	>OR= 130
Low Risk	<100	<130	>OR= 130*	>OR= 160
*After an adequate non n	harmacological interventi	ion for at least 3 months		

<u>Final</u>

References: Management of Dyslipidaemia for the Prevention of Stroke: Clinical Practice Recommendations from the Lipid Association of India. Current Vascular Pharmacology, 2022, 20, 134-155.

LIVER FUNCTION PROFILE, SERUM

BILIRUBIN, TOTAL	1.5	High	Upto 1.2	mg/dL
METHOD: COLORIMETRIC DIAZO METHOD				
BILIRUBIN, DIRECT	0.6	High	< 0.30	mg/dL
METHOD: COLORIMETRIC DIAZO METHOD				
BILIRUBIN, INDIRECT	0.90		0.1 - 1.0	mg/dL
METHOD: CALCULATED PARAMETER				
TOTAL PROTEIN	7.4		6.0 - 8.0	g/dL
METHOD: SPECTROPHOTOMETRY, BIURET				
ALBUMIN	4.6		3.97 - 4.94	g/dL
METHOD: SPECTROPHOTOMETRY, BROMOCRESOL GREEN(BCG) -	DYE BINDING			
GLOBULIN	2.8		2.0 - 3.5	g/dL
METHOD: CALCULATED PARAMETER				
ALBUMIN/GLOBULIN RATIO	1.6		1.0 - 2.1	RATIO
METHOD: CALCULATED PARAMETER				
ASPARTATE AMINOTRANSFERASE (AST/SGOT)	27		< OR = 50	U/L
METHOD: SPECTROPHOTOMETRY, WITH PYRIDOXAL PHOSPHATE	ACTIVATION-IFCC			
ALANINE AMINOTRANSFERASE (ALT/SGPT)	51	High	< OR = 50	U/L
METHOD: SPECTROPHOTOMETRY, WITH PYRIDOXAL PHOSPHATE	ACTIVATION-IFCC			
ALKALINE PHOSPHATASE	69		40 - 129	U/L
METHOD: SPECTROPHOTOMETRY, PNPP, AMP BUFFER - IFCC				
GAMMA GLUTAMYL TRANSFERASE (GGT)	14		0 - 60	U/L
METHOD: ENZYMATIC COLORIMETRIC ASSAY STANDARDIZED AC	GAINST IFCC / SZASZ			
LACTATE DEHYDROGENASE	155		125 - 220	U/L
METHOD: SPECTROPHOTOMETRY, LACTATE TO PYRUVATE - UV-IF-	CC			
BLOOD UREA NITROGEN (BUN), SERUM				

6 - 20 **BLOOD UREA NITROGEN** 11.0 mg/dL

 ${\tt METHOD: SPECTROPHOTOMETRY, KINETIC TEST WITH UREASE AND GLUTAMATE DEHYDROGENASE}$

CREATININE, SERUM



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^{*}After an adequate non-pharmacological intervention for at least 3 months.





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CREATININE		0.70		0.7 - 1.2	mg/dL
METHOD : SPECTROPHOTON	METRIC, JAFFE'S KINETICS				3,
BUN/CREAT RATIO					
BUN/CREAT RATIO		15.71	High	8.0 - 15.0	
METHOD : CALCULATED PA	RAMETER				
URIC ACID, SERUM					
URIC ACID		5.8		3.4 - 7.0	mg/dL
METHOD : SPECTROPHOTON	METRY, URICASE				-
TOTAL PROTEIN, SE	RUM				
TOTAL PROTEIN		7.4		6.0 - 8.0	g/dL
METHOD : SPECTROPHOTON	METRY, BIURET				
ALBUMIN, SERUM					
ALBUMIN		4.6		3.97 - 4.94	g/dL
METHOD : SPECTROPHOTON	METRY, BROMOCRESOL GRE	EN(BCG) - DYE BINDING			
GLOBULIN					
GLOBULIN		2.8		2.0 - 3.5	g/dL
METHOD : CALCULATED PA	RAMETER				
ELECTROLYTES (NA	/K/CL), SERUM				
SODIUM, SERUM		137		136 - 145	mmol/L
METHOD : ISE INDIRECT					
POTASSIUM, SERUM		4.0		3.5 - 5.1	mmol/L
METHOD : ISE INDIRECT					
CHLORIDE, SERUM		102		98 - 107	mmol/L
METHOD : ISE INDIRECT					









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Interpretation(s)

Sodium	Potassium	Chloride
Decreased in: CCF, cirrhosis,	Decreased in: Low potassium	Decreased in: Vomiting, diarrhea,
vomiting, diarrhea, excessive	intake,prolonged vomiting or diarrhea,	renal failure combined with salt
sweating, salt-losing	RTA types I and II,	deprivation, over-treatment with
nephropathy, adrenal insufficiency,	hyperaldosteronism, Cushing's	diuretics, chronic respiratory acidosis,
nephrotic syndrome, water	syndrome,osmotic diuresis (e.g.,	diabetic ketoacidosis, excessive
intoxication, SIADH. Drugs:	hyperglycemia),alkalosis, familial	sweating, SIADH, salt-losing
thiazides, diuretics, ACE inhibitors,	periodic paralysis,trauma	nephropathy, porphyria, expansion of
chlorpropamide,carbamazepine,anti	(transient).Drugs: Adrenergic agents,	extracellular fluid volume,
depressants (SSRI), antipsychotics.	diuretics.	adrenalinsufficiency,
		hyperaldosteronism, metabolic
		alkalosis. Drugs: chronic
		laxative,corticosteroids, diuretics.
Increased in: Dehydration	Increased in: Massive hemolysis,	Increased in: Renal failure, nephrotic
(excessivesweating, severe	severe tissue damage, rhabdomyolysis,	syndrome, RTA,dehydration,
vomiting or diarrhea),diabetes	acidosis, dehydration,renal failure,	overtreatment with
mellitus, diabetesinsipidus,	Addison's disease, RTA type IV,	saline,hyperparathyroidism, diabetes
hyperaldosteronism, inadequate	hyperkalemic familial periodic	insipidus, metabolic acidosis from
water intake. Drugs: steroids,	paralysis. Drugs: potassium salts,	diarrhea (Loss of HCO3-), respiratory
licorice,oral contraceptives.	potassium- sparing diuretics,NSAIDs,	alkalosis, hyperadrenocorticism.
	beta-blockers, ACE inhibitors, high-	Drugs: acetazolamide,androgens,
	dose trimethoprim-sulfamethoxazole.	hydrochlorothiazide, salicylates.
Interferences: Severe lipemia or	Interferences: Hemolysis of sample,	Interferences:Test is helpful in
hyperproteinemi, if sodium analysis	delayed separation of serum,	assessing normal and increased anion
involves a dilution step can cause	prolonged fist clenching during blood	gap metabolic acidosis and in
spurious results. The serum sodium	drawing, and prolonged tourniquet	distinguishing hypercalcemia due to
falls about 1.6 mEq/L for each 100	placement. Very high WBC/PLT counts	hyperparathyroidism (high serum
mg/dL increase in blood glucose.	may cause spurious. Plasma potassium	chloride) from that due to malignancy
	levels are normal.	(Normal serum chloride)

PHYSICAL EXAMINATION, URINE

COLOR PALE YELLOW APPEARANCE CLEAR

Comments

NOTE: MICROSCOPIC EXAMINATION OF URINE IS PERFORMED ON CENTRIFUGED URINARY SEDIMENT. IN NORMAL URINE SAMPLES CAST AND CRYSTALS ARE NOT DETECTED.

CHEMICAL EXAMINATION, URINE

PH 7.0 4.7 - 7.5 SPECIFIC GRAVITY 1.010 1.003 - 1.035 **PROTEIN** NOT DETECTED NOT DETECTED **GLUCOSE** NOT DETECTED NOT DETECTED **KETONES** NOT DETECTED NOT DETECTED NOT DETECTED NOT DETECTED **BLOOD** NOT DETECTED NOT DETECTED **BILIRUBIN UROBILINOGEN NORMAL NORMAL**











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NITRITE	NOT DETECTED	NOT DETECTED	
LEUKOCYTE ESTERASE	NOT DETECTED	NOT DETECTED	
MICROSCOPIC EXAMINATION, URINE			
RED BLOOD CELLS	NOT DETECTED	NOT DETECTED	/HPF
PUS CELL (WBC'S)	0-1	0-5	/HPF
EPITHELIAL CELLS	0-1	0-5	/HPF
CASTS	NOT DETECTED		
CRYSTALS	NOT DETECTED		
BACTERIA	NOT DETECTED	NOT DETECTED	
METHOD: DIP STICK/MICRO SCOPY/REFLECTANCE SPECTROPHOT	OMETRY		
YEAST	NOT DETECTED	NOT DETECTED	









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Interpretation(s)

The following table describes the probable conditions, in which the analytes are present in urine

Presence of	Conditions			
Proteins	Inflammation or immune illnesses			
Pus (White Blood Cells)	Urinary tract infection, urinary tract or kidney stone, tumors or any kind			
CI	of kidney impairment			
Glucose	Diabetes or kidney disease			
Ketones	Diabetic ketoacidosis (DKA), starvation or thirst			
Urobilinogen	Liver disease such as hepatitis or cirrhosis			
Blood	Renal or genital disorders/trauma			
Bilirubin	Liver disease			
Erythrocytes	Urological diseases (e.g. kidney and bladder cancer, urolithiasis), urinary tract infection and glomerular diseases			
Leukocytes	Urinary tract infection, glomerulonephritis, interstitial nephritis either acute or chronic, polycystic kidney disease, urolithiasis, contamination by genital secretions			
Epithelial cells	Urolithiasis, bladder carcinoma or hydronephrosis, ureteric stents or			
	bladder catheters for prolonged periods of time			
Granular Casts	Low intratubular pH, high urine osmolality and sodium concentration, interaction with Bence-Jones protein			
Hyaline casts	Physical stress, fever, dehydration, acute congestive heart failure, renal diseases			
Calcium oxalate	Metabolic stone disease, primary or secondary hyperoxaluria, intravenous infusion of large doses of vitamin C, the use of vasodilator naftidrofuryl oxalate or the gastrointestinal lipase inhibitor orlistat, ingestion of ethylene glycol or of star fruit (Averrhoa carambola) or its juice			
Uric acid	arthritis			
Bacteria	Urinary infectionwhen present in significant numbers & with pus cells.			
Trichomonas vaginalis	Vaginitis, cervicitis or salpingitis			

THYROID PANEL, SERUM

ТЗ	91.1	80 - 200	ng/dL
METHOD: ELECTROCHEMILUMINESCENCE IMMUNO ASSAY			
T4	6.40	5.1 - 14.1	μg/dL
METHOD: ELECTROCHEMILUMINESCENCE IMMUNO ASSAY			
TSH (ULTRASENSITIVE)	1.950	0.27 - 4.2	uIU/mL

METHOD: ELECTROCHEMILUMINESCENCE IMMUNO ASSAY









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F-703, LADO SARAI, MEHRAULI SOUTH WEST DELHI **NEW DELHI 110030**

DELHI INDIA 8800465156

Shop CG 017, PALM SPRINGS PLAZA

GURUGRAM, 122001 HARYANA, INDIA Tel: 9111591115

PATIENT NAME: NISHAT SAINI PATIENT ID: NISHM250885282

0282WD00001 AGE: 37 Years ACCESSION NO: SEX: Male ABHA NO:

RECEIVED: 01/04/2023 08:23 03/04/2023 11:44 DRAWN: REPORTED:

REFERRING DOCTOR: SELF CLIENT PATIENT ID:

Test Report Status Results **Biological Reference Interval** Units <u>Final</u>

Interpretation(s)

Triiodothyronine T3, Thyroxine T4, and Thyroid Stimulating Hormone TSH are thyroid hormones which affect almost every physiological process in the body, including growth, development, metabolism, body temperature, and heart rate.

Production of T3 and its prohormone thyroxine (T4) is activated by thyroid-stimulating hormone (TSH), which is released from the pituitary gland. Elevated concentrations of T3, and T4 in the blood inhibit the production of TSH.

Excessive secretion of thyroxine in the body is hyperthyroidism, and deficient secretion is called hypothyroidism.

In primary hypothyroidism, TSH levels are significantly elevated, while in secondary and tertiary hyporthyroidism, TSH levels are low. Below mentioned are the guidelines for Pregnancy related reference ranges for Total T4, TSH & Total T3. Measurement of the serum TT3 level is a more sensitive test for the diagnosis of hyperthyroidism, and measurement of TT4 is more useful in the diagnosis of hypothyroidism, Most of the thyroid hormone in blood is bound to transport proteins. Only a very small fraction of the circulating hormone is free and biologically active. It is advisable to detect Free T3, FreeT4 along with TSH, instead of testing for albumin bound Total T3, Total T4.

Sr. No.	TSH	Total T4	FT4	Total T3	Possible Conditions
1	High	Low	Low	Low	(1) Primary Hypothyroidism (2) Chronic autoimmune Thyroiditis (3)
					Post Thyroidectomy (4) Post Radio-Iodine treatment
2	High	Normal	Normal	Normal	(1)Subclinical Hypothyroidism (2) Patient with insufficient thyroid
					hormone replacement therapy (3) In cases of Autoimmune/Hashimoto
					thyroiditis (4). Isolated increase in TSH levels can be due to Subclinical
					inflammation, drugs like amphetamines, Iodine containing drug and
					dopamine antagonist e.g. domperidone and other physiological reasons.
3	Normal/Low	Low	Low	Low	(1) Secondary and Tertiary Hypothyroidism
4	Low	High	High	High	(1) Primary Hyperthyroidism (Graves Disease) (2) Multinodular Goitre
					(3)Toxic Nodular Goitre (4) Thyroiditis (5) Over treatment of thyroid
					hormone (6) Drug effect e.g. Glucocorticoids, dopamine, T4
					replacement therapy (7) First trimester of Pregnancy
5	Low	Normal	Normal	Normal	(1) Subclinical Hyperthyroidism
6	High	High	High	High	(1) TSH secreting pituitary adenoma (2) TRH secreting tumor
7	Low	Low	Low	Low	(1) Central Hypothyroidism (2) Euthyroid sick syndrome (3) Recent
					treatment for Hyperthyroidism
8	Normal/Low	Normal	Normal	High	(1) T3 thyrotoxicosis (2) Non-Thyroidal illness
9	Low	High	High	Normal	(1) T4 Ingestion (2) Thyroiditis (3) Interfering Anti TPO antibodies

REF: 1. TIETZ Fundamentals of Clinical chemistry 2. Guidlines of the American Thyroid association during pregnancy and Postpartum, 2011. NOTE: It is advisable to detect Free T3, FreeT4 along with TSH, instead of testing for albumin bound Total T3, Total T4.TSH is not affected by variation in thyroid - binding protein. TSH has a diurnal rhythm, with peaks at 2:00 - 4:00 a.m. And troughs at 5:00 - 6:00 p.m. With ultradian variations.

MICROSCOPIC EXAMINATION, STOOL

TEST CANCELLED AS SPECIMEN NOT RECEIVED REMARK

METHOD: MICROSCOPIC EXAMINATION



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Interpretation(s)

Stool routine analysis is only a screening test for disorders of gastrointentestinal tract like infection, malabsorption, etc. The following table describes the probable conditions, in which the analytes are present in stool.

PRESENCE OF	CONDITION			
Pus cells	Pus in the stool is an indication of infection			
Red Blood cells	Parasitic or bacterial infection or an inflammatory bowel condition such as			
	ulcerative colitis			
Parasites	Infection of the digestive system. Stool examination for ova and parasite detects presence of parasitic infestation of gastrointestinal tract. Various forms of parasite that can be detected include cyst, trophozoite and larvae. One negative result does not rule out the possibility of parasitic infestation. Intermittent shedding of parasites warrants examinations of multiple specimens tested on consecutive days. Stool specimens for parasitic examination should be collected before initiation of antidiarrheal therapy or antiparasitic therapy. This test does not detect presence of opportunistic parasites like Cyclospora, Cryptosporidia and Isospora species. Examination of Ova and Parasite has been carried out by direct and concentration techniques.			
Mucus	Mucus is a protective layer that lubricates, protects& reduces damage due to bacteria or viruses.			
Charcot-Leyden crystal	Parasitic diseases.			
Ova & cyst	Ova & cyst indicate parasitic infestation of intestine.			
Frank blood	Bleeding in the rectum or colon.			
Occult blood	Occult blood indicates upper GI bleeding.			
Macrophages	Macrophages in stool are an indication of infection as they are protective cells.			
Epithelial cells	Epithelial cells that normally line the body surface and internal organs show up in stool when there is inflammation or infection.			
Fat	Increased fat in stool maybe seen in conditions like diarrhoea or malabsorption.			
рН	Normal stool pH is slightly acidic to neutral. Breast-fed babies generally have an acidic stool.			

ADDITIONAL STOOL TESTS:

- 1. Stool Culture:- This test is done to find cause of GI infection, make decision about best treatment for GI infection & to find out if treatment for GI infection worked.
- 2. Fecal Calprotectin: It is a marker of intestinal inflammation. This test is done to differentiate Inflammatory Bowel Disease (IBD) from Irritable Bowel Syndrome (IBS).
- Fecal Occult Blood Test(FOBT): This test is done to screen for colon cancer & to evaluate possible cause of unexplained anaemia. 3.
- Clostridium Difficile Toxin Assay: This test is strongly recommended in healthcare associated bloody or waterydiarrhoea, due to 4. overuse of broad spectrum antibiotics which alter the normal GI flora.
- 5. Biofire (Film Array) GI PANEL: In patients of Diarrhoea, Dysentry, Rice watery Stool, FDA approved, Biofire Film Array Test,(Real Time Multiplex PCR) is strongly recommended as it identifies organisms, bacteria, fungi, virus ,parasite and other opportunistic pathogens, Vibrio cholera infections only in 3 hours. Sensitivity 96% & Specificity 99%.
- Rota Virus Immunoassay: This test is recommended in severe gastroenteritis in infants & children associated with watery diarrhoea, vomitting& abdominal cramps. Adults are also affected. It is highly contagious in nature.









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Test Report Status Final Results Biological Reference Interval Units

ABO GROUP & RH TYPE, EDTA WHOLE BLOOD

ABO GROUP O

 ${\tt METHOD: HEMAGGLUTINATION \ REACTION \ ON \ SOLID \ PHASE}$

RH TYPE RH+

METHOD: HEMAGGLUTINATION REACTION ON SOLID PHASE

XRAY-CHEST

»» BOTH THE LUNG FIELDS ARE CLEAR

»» BOTH THE COSTOPHRENIC AND CARDIOPHRENIC ANGLES ARE CLEAR

»» BOTH THE HILA ARE NORMAL

»» CARDIAC AND AORTIC SHADOWS APPEAR NORMAL»» BOTH THE DOMES OF THE DIAPHRAGM ARE NORMAL

»» VISUALIZED BONY THORAX IS NORMAL

IMPRESSION NO ABNORMALITY DETECTED

TMT OR ECHO

TMT OR ECHO STRESS TEST IS NEGATIVE FOR RMI

ECG

ECG WITHIN NORMAL LIMITS

MEDICAL HISTORY

RELEVANT PRESENT HISTORY

RELEVANT PAST HISTORY

RELEVANT PERSONAL HISTORY

RELEVANT FAMILY HISTORY

OCCUPATIONAL HISTORY SERVICE
HISTORY OF MEDICATIONS SUPPLEMENTS

ANTHROPOMETRIC DATA & BMI

HEIGHT IN METERS 1.66 mts WEIGHT IN KGS. 77 Kgs

BMI & Weight Status as follows: kg/sqmts

Below 18.5: Underweight 18.5 - 24.9: Normal 25.0 - 29.9: Overweight 30.0 and Above: Obese

GENERAL EXAMINATION

MENTAL / EMOTIONAL STATE NORMAL PHYSICAL ATTITUDE NORMAL









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GENERAL APPEARANCE / NUTRITIONAL STATUS OVERWEIGHT
BUILT / SKELETAL FRAMEWORK AVERAGE
FACIAL APPEARANCE NORMAL
SKIN NORMAL
UPPER LIMB NORMAL
LOWER LIMB NORMAL
NECK NORMAL

NECK LYMPHATICS / SALIVARY GLANDS NOT ENLARGED OR TENDER

THYROID GLAND NOT ENLARGED

CAROTID PULSATION NORMAL TEMPERATURE NORMAL

PULSE 72/ MINUTE, REGULAR, ALL PERIPHERAL PULSES FELT.

RESPIRATORY RATE NORMAL

CARDIOVASCULAR SYSTEM

BP 120/84 MMHG mm/Hg

(SUPINE) NORMAL NORMAL NORMAL

ABSENT

RESPIRATORY SYSTEM

SIZE AND SHAPE OF CHEST NORMAL

MOVEMENTS OF CHEST SYMMETRICAL

BREATH SOUNDS INTENSITY NORMAL

BREATH SOUNDS QUALITY VESICULAR (NORMAL)

ADDED SOUNDS ABSENT

PER ABDOMEN

PERICARDIUM

HEART SOUNDS

APEX BEAT

MURMURS

APPEARANCE NORMAL
VENOUS PROMINENCE ABSENT
LIVER NOT PALPABLE
SPLEEN NOT PALPABLE

CENTRAL NERVOUS SYSTEM

HIGHER FUNCTIONS NORMAL
CRANIAL NERVES NORMAL
CEREBELLAR FUNCTIONS NORMAL









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Test Report Status Results **Biological Reference Interval** Units <u>Final</u> SENSORY SYSTEM NORMAL MOTOR SYSTEM NORMAL REFLEXES NORMAL **MUSCULOSKELETAL SYSTEM** SPINE NORMAL JOINTS **NORMAL BASIC EYE EXAMINATION** DISTANT VISION RIGHT EYE WITHOUT GLASSES 6/6 DISTANT VISION LEFT EYE WITHOUT GLASSES 6/6 NEAR VISION RIGHT EYE WITHOUT GLASSES N/6 NEAR VISION LEFT EYE WITHOUT GLASSES N/6 COLOUR VISION 17/17

SUMMARY

REMARKS / RECOMMENDATIONS

ADVISED

LIFESTYLE CHANGES

IRON RICH HIGH FIBRE DIET

LFT AFER ONE WEEK

FOLLOW UP WITH GE PHYSICIAN FOR FURTHER MANAGEMENT.

BLOOD COUNTS, EDTA WHOLE BLOOD-The cell morphology is well preserved for 24hrs. However after 24-48 hrs a progressive increase in MCV and HCT is observed leading to a decrease in MCHC. A direct smear is recommended for an accurate differential count and for examination of RBC morphology.

RBC AND PLATELET INDICES-Mentzer index (MCV/RBC) is an automated cell-counter based calculated screen tool to differentiate cases of Iron deficiency anaemia(>13) from Beta thalassaemia trait

(<13) in patients with microcytic anaemia. This needs to be interpreted in line with clinical correlation and suspicion. Estimation of HbA2 remains the gold standard for diagnosing a case of beta thalassaemia trait.

WBC DIFFERENTIAL COUNT-The optimal threshold of 3.3 for NLR showed a prognostic possibility of clinical symptoms to change from mild to severe in COVID positive

patients. When age = 49.5 years old and NLR = 3.3, 46.1% COVID-19 patients with mild disease might become severe. By contrast, when age < 49.5 years old and NLR < 3.3, COVID-19 patients tend to show mild disease.

(Reference to - The diagnostic and predictive role of NLR, d-NLR and PLR in COVID-19 patients A.-P. Yang, et al. International Immunopharmacology 84 (2020) 106504 This ratio element is a calculated parameter and out of NABL scope.

ERYTHROCYTE SEDIMENTATION RATE (ESR), WHOLE BLOOD-TEST DESCRIPTION:

Erythrocyte sedimentation rate (ESR) is a test that indirectly measures the degree of inflammation present in the body. The test actually measures the rate of fall (sedimentation) of erythrocytes in a sample of blood that has been placed into a tall, thin, vertical tube. Results are reported as the millimetres of clear fluid (plasma) that are present at the top portion of the tube after one hour. Nowadays fully automated instruments are available to measure ESR.

ESR is not diagnostic it is a non-specific test that may be elevated in a number of different conditions. It provides general information about the presence of an inflammatory condition.CRP is superior to ESR because it is more sensitive and reflects a more rapid change **TEST INTERPRETATION**

Increase in: Infections, Vasculities, Inflammatory arthritis, Renal disease, Anemia, Malignancies and plasma cell dyscrasias, Acute allergy Tissue injury, Pregnancy, Estrogen medication, Aging.

Finding a very accelerated ESR(>100 mm/hour) in patients with ill-defined symptoms directs the physician to search for a systemic disease (Paraproteinemias,

Disseminated malignancies, connective tissue disease, severe infections such as bacterial endocarditis). In pregnancy BRI in first trimester is 0-48 mm/hr(62 if anemic) and in second trimester (0-70 mm /hr(95 if anemic). ESR returns to normal 4th week post partum. Decreased in: Polycythermia vera, Sickle cell anemia



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LIMITATIONS

False elevated ESR: Increased fibrinogen, Drugs(Vitamin A, Dextran etc), Hypercholesterolemia

False Decreased: Poikilocytosis, (SickleCells, spherocytes), Microcytosis, Low fibrinogen, Very high WBC counts, Drugs (Quinine,

salicylates)

REFERENCE:

1. Nathan and Oski's Haematology of Infancy and Childhood, 5th edition 2. Paediatric reference intervals. AACC Press, 7th edition. Edited by S. Soldin 3. The reference for the adult reference range is "Practical Haematology by Dacie and Lewis,10th edition. GLUCOSE FASTING, FLUORIDE PLASMA-TEST DESCRIPTION

Normally, the glucose concentration in extracellular fluid is closely regulated so that a source of energy is readily available to tissues and sothat no glucose is excreted in the

Increased in:Diabetes mellitus, Cushing's syndrome (10 – 15%), chronic pancreatitis (30%). Drugs:corticosteroids,phenytoin, estrogen, thiazides.

Decreased in:Pancreatic islet cell disease with increased insulin,insulinoma,adrenocortical insufficiency,hypopituitarism,diffuse liver disease, malignancy(adrenocortical,stomach,fibrosarcoma),infant of a diabetic mother,enzyme deficiency diseases(e.g.galactosemia),Drugs-insulin,ethanol,propranolol

sulfonylureas,tolbutamide,and other oral hypoglycemic agents.

NOTE: While random serum glucose levels correlate with home glucose monitoring results (weekly mean capillary glucose values),there is wide fluctuation within individuals. Thus, glycosylated hemoglobin(HbA1c) levels are favored to monitor glycemic control.

High fasting glucose level in comparison to post prandial glucose level may be seen due to effect of Oral Hypoglycaemics & Insulin treatment, Renal Glyosuria, Glycaemic

index & response to food consumed, Alimentary Hypoglycemia, Increased insulin response & sensitivity etc. GLYCOSYLATED HEMOGLOBIN(HBA1C), EDTA WHOLE BLOOD-**Used For**:

- Evaluating the long-term control of blood glucose concentrations in diabetic patients.
 Diagnosing diabetes.

The ADA recommends measurement of HbA1c (typically 3-4 times per year for type 1 and poorly controlled type 2 diabetic patients, and 2 times per year for well-controlled type 2 diabetic patients) to determine whether a patients metabolic control has remained continuously within the target range.

- eAG (Estimated average glucose) converts percentage HbA1c to md/dl, to compare blood glucose levels.
 eAG gives an evaluation of blood glucose levels for the last couple of months.
 eAG is calculated as eAG (mg/dl) = 28.7 * HbA1c 46.7

HbA1c Estimation can get affected due to :

- 1. Shortened Erythrocyte survival : Any condition that shortens erythrocyte survival or decreases mean erythrocyte age (e.g. recovery from acute blood loss,hemolytic anemia) will falsely lower HbA1c test results. Fructosamine is recommended in these patients which indicates diabetes control over 15 days.
- 2.Vitamin C & E are reported to falsely lower test results. (possibly by inhibiting glycation of hemoglobin.

 3. Iron deficiency anemia is reported to increase test results. Hypertriglyceridemia, uremia, hyperbilirubinemia, chronic alcoholism, chronic ingestion of salicylates & opiates addiction are reported to interfere with some assay methods, falsely increasing results.
- 4. Interference of hemoglobinopathies in HbA1c estimation is seen in

- a) Homozygous hemoglobinopathy. Fructosamine is recommended for testing of HbA1c.
 b) Heterozygous state detected (D10 is corrected for HbS & HbC trait.)
 c) HbF > 25% on alternate paltform (Boronate affinity chromatography) is recommended for testing of HbA1c.Abnormal Hemoglobin electrophoresis (HPLC method) is

GLUCOSE, POST-PRANDIAL, PLASMA-High fasting glucose level in comparison to post prandial glucose level may be seen due to effect of Oral Hypoglycaemics & Insulin treatment, Renal Glyosuria, Glycaemic index & response to food consumed, Alimentary Hypoglycemia, Increased insulin response & sensitivity etc.Additional test HbA1c LIVER FUNCTION PROFILE, SERUM-

Bilirubin is a yellowish pigment found in bile and is a breakdown product of normal heme catabolism. Bilirubin is excreted in bile and urine, and elevated levels may give yellow discoloration in jaundice. **Elevated levels** results from increased bilirubin production (eg, hemolysis and ineffective erythropoiesis), decreased bilirubin excretion (eg, obstruction and hepatitis), and abnormal bilirubin metabolism (eg, hereditary and neonatal jaundice). Conjugated (direct) bilirubin is elevated more than unconjugated (indirect) bilirubin in Viral hepatitis, Drug reactions, Alcoholic liver disease Conjugated (direct) bilirubin is also elevated more than unconjugated (indirect) bilirubin when there is some kind of blockage of the bile ducts like in Gallstones getting into the bile ducts, tumors &Scarring of the bile ducts. Increased unconjugated (indirect) bilirubin may be a result of Hemolytic or pernicious anemia, Transfusion reaction & a common metabolic condition termed Gilbert syndrome, due to low levels of the enzyme that attaches sugar molecules to bilirubin. **AST** is an enzyme found in various parts of the body. AST is found in the liver, heart, skeletal muscle, kidneys, brain, and red blood cells, and it is commonly measured

clinically as a marker for liver health. AST levels increase during chronic viral hepatitis, blockage of the bile duct, cirrhosis of the liver,liver cancer,kidney failure,hemolytic anemia,pancreatitis,hemochromatosis. AST levels may also increase after a heart attack or strenuous activity.ALT test measures the amount of this enzyme in the blood.ALT is found mainly in the liver, but also in smaller amounts in the kidneys,heart,muscles, and pancreas.It is commonly measured as a part of a diagnostic evaluation of hepatocellular injury, to determine liver health AST levels increase during acute hepatitis, sometimes due to a viral infection, ischemia to the liver, chronic hepatitis, obstruction of bile ducts, cirrhosis.

ALP is a protein found in almost all body tissues. Tissues with higher amounts of ALP include the liver, bile ducts and bone. Elevated ALP levels are seen in Biliary obstruction, Osteoblastic bone tumors, osteomalacia, hepatitis, Hyperparathyroidism, Leukemia, Lymphoma, Pagets disease, Rickets, Sarcoidosis etc. Lower-than-normal ALP levels seen in Hypophosphatasia, Malnutrition, Protein deficiency, Wilsons disease.

GGT is an enzyme found in cell membranes of many tissues mainly in the liver, kidney and pancreas. It is also found in other tissues including intestine, spleen, heart, brain and seminal vesicles. The highest concentration is in the kidney, but the liver is considered the source of normal enzyme activity. Serum GGT has been widely used as an index of liver dysfunction. Elevated serum GGT activity can be found in diseases of the liver, biliary system and pancreas. Conditions that increase serum GGT are obstructive liver disease, high alcohol consumption and use of enzyme-inducing drugs etc.





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Total Protein also known as total protein, is a biochemical test for measuring the total amount of protein in serum. Protein in the plasma is made up of albumin and globulin. Higher-than-normal levels may be due to: Chronic inflammation or infection, including HIV and hepatitis B or C, Multiple myeloma, Waldenstroms disease. Lower-than-normal levels may be due to: Agammaglobulinemia, Bleeding (hemorrhage), Burns, Glomerulonephritis, Liver disease, Malabsorption, Malnutrition, Nephrotic syndrome, Protein-losing enteropathy etc. **Albumin** is the most abundant protein in human blood plasma. It is produced in the liver. Albumin constitutes about half of the blood serum protein. Low blood albumin levels

(hypoalbuminemia) can be caused by:Liver disease like cirrhosis of the liver, nephrotic syndrome,protein-losing enteropathy,Burns,hemodilution,increased vascular permeability or decreased lymphatic clearance,malnutrition and wasting etc

BLOOD UREA NITROGEN (BUN), SERUM-**Causes of Increased** levels include Pre renal (High protein diet, Increased protein catabolism, GI haemorrhage, Cortisol, Dehydration, CHF Renal), Renal Failure, Post Renal (Malignancy, Nephrolithiasis, Prostatism) **Causes of decreased** level include Liver disease, SIADH.

OREATININE, SERUM-**Higher than normal level may be due to:**• Blockage in the urinary tract, Kidney problems, such as kidney damage or failure, infection, or reduced blood flow, Loss of body fluid (dehydration), Muscle problems, such as breakdown of muscle fibers, Problems during pregnancy, such as seizures (eclampsia)), or high blood pressure caused by pregnancy (preeclampsia) Lower than normal level may be due to:

Lower than normal level may be due to:

Myasthenia Gravis, Muscuophy

URIC ACID, SERUM-Causes of Increased levels:-Dietary(High Protein Intake,Prolonged Fasting,Rapid weight loss),Gout,Lesch nyhan syndrome,Type 2 DM,Metabolic syndrome Causes of decreased levels-Low Zinc intake,OCP,Multiple Sclerosis

TOTAL PROTEIN, SERUM-is a biochemical test for measuring the total amount of protein in serum.Protein in the plasma is made up of albumin and globulin.

Higher-than-normal levels may be due to: Chronic inflammation or infection, including HIV and hepatitis B or C, Multiple myeloma, Waldenstroms disease.

Lower-than-normal levels may be due to: Agammaglobulinemia, Bleeding (hemorrhage), Burns, Glomerulonephritis, Liver disease, Malabsorption, Malnutrition, Nephrotic

syndrome, Protein-losing enteropathy etc.
ALBUMIN, SERUMHuman serum albumin is the most abundant protein in human blood plasma. It is produced in the liver. Albumin constitutes about half of the blood serum protein. **Low** blood albumin levels (hypoalbuminemia) can be caused by: Liver disease like cirrhosis of the liver, nephrotic syndrome, protein-losing enteropathy, Burns, hemodilution, increased vascular permeability or decreased lymphatic clearance, malnutrition and wasting etc.

ABO GROUP & RH TYPE, EDTA WHOLE BLOOD-Blood group is identified by antigens and antibodies present in the blood. Antigens are protein molecules found on the surface of red blood cells. Antibodies are found in plasma. To determine blood group, red cells are mixed with different antibody solutions to give A,B,O or AB.

Disclaimer: "Please note, as the results of previous ABO and Rh group (Blood Group) for pregnant women are not available, please check with the patient records for availability of the same.

The test is performed by both forward as well as reverse grouping methods.

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GURUGRAM, 122001 HARYANA, INDIA Tel: 9111591115

PATIENT NAME: NISHAT SAINI PATIENT ID: NISHM250885282

0282WD00001 AGE: 37 Years ACCESSION NO: SEX: Male ABHA NO:

DRAWN: RECEIVED: 01/04/2023 08:23 REPORTED: 03/04/2023 11:44

REFERRING DOCTOR: SFLF CLIENT PATIENT ID:

Test Report Status Results Units <u>Final</u>

MEDI WHEEL FULL BODY HEALTH CHECK UP BELOW 40 MALE

ULTRASOUND ABDOMEN ULTRASOUND ABDOMEN U.S.G Scan S/o Grade I fatty liver. No other significant abnormality detected.

Please correlate clinically.

End Of Report Please visit www.srlworld.com for related Test Information for this accession

Dr. Deblina Naithani **Consultant Physician**

CONDITIONS OF LABORATORY TESTING & REPORTING

- 1. It is presumed that the test sample belongs to the patient named or identified in the test requisition form.
- 2. All tests are performed and reported as per the
- turnaround time stated in the SRL Directory of Services. 3. Result delays could occur due to unforeseen
- circumstances such as non-availability of kits / equipment breakdown / natural calamities / technical downtime or any other unforeseen event.
- 4. A requested test might not be performed if:
 - i. Specimen received is insufficient or inappropriate
 - ii. Specimen quality is unsatisfactory
 - iii. Incorrect specimen type
 - iv. Discrepancy between identification on specimen container label and test requisition form

- 5. SRL confirms that all tests have been performed or assayed with highest quality standards, clinical safety & technical integrity.
- 6. Laboratory results should not be interpreted in isolation; it must be correlated with clinical information and be interpreted by registered medical practitioners only to determine final diagnosis.
- Test results may vary based on time of collection, physiological condition of the patient, current medication or nutritional and dietary changes. Please consult your doctor or call us for any clarification.
- 8. Test results cannot be used for Medico legal purposes.
- 9. In case of gueries please call customer care (91115 91115) within 48 hours of the report.

SRL Limited

Fortis Hospital, Sector 62, Phase VIII, Mohali 160062



