









Name : Ms. USHA JAIN

Lab No.

A/c Status

Mo. Gonz Gain

Ref By: SELF

Age: 72 Years Gender: Female

Collected Received

: 9/4/2021 6:50:00AM

Reported

: 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

Report Status : Final

Test Name Results Units Bio. Ref. Interval

SWASTHFIT TAX SAVER ADVANCE PACKAGE

156976105

HEMOGRAM			
Hemoglobin (Photometry)	11.00	g/dL	12.00 - 15.00
Packed Cell Volume (PCV) (Calculated)	34.30	%	36.00 - 46.00
RBC Count (Electrical Impedence)	3.88	mill/mm3	3.80 - 4.80
MCV (Electrical Impedence)	88.40	fL	83.00 - 101.00
MCH (Calculated)	28.40	pg	27.00 - 32.00
MCHC (Calculated)	32.20	g/dL	31.50 - 34.50
Red Cell Distribution Width (RDW) (Electrical Impedence)	15.30	%	11.60 - 14.00
Total Leukocyte Count (TLC) (Electrical Impedence)	9.80	thou/mm3	4.00 - 10.00
Differential Leucocyte Count (DLC) (VCS Technology)			
Segmented Neutrophils	57.70	%	40.00 - 80.00
Lymphocytes	27.60	%	20.00 - 40.00
Monocytes	10.10	%	2.00 - 10.00
Eosinophils	3.50	%	1.00 - 6.00
Basophils	1.10	%	<2.00
Absolute Leucocyte Count (Calculated)			
Neutrophils	5.65	thou/mm3	2.00 - 7.00
Lymphocytes	2.70	thou/mm3	1.00 - 3.00
Monocytes	0.99	thou/mm3	0.20 - 1.00
Eosinophils	0.34	thou/mm3	0.02 - 0.50
Basophils	0.11	thou/mm3	0.02 - 0.10



Page 1 of 15





Female

Gender:







Name : Ms. USHA JAIN

Collected

Lab No.

156976105 Age: 72 Years

Received Reported : 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

A/c Status : P

Ref By: SELF

Report Status : Final

Test Name	Results	Units	Bio. Ref. Interval
Platelet Count (Electrical impedence)	225.0	thou/mm3	150.00 - 410.00
Mean Platelet Volume (Electrical Impedence)	7.8	fL	6.5 - 12.0
ESR (Capillary photometry)	37	mm/hr	0.00 - 30.00

Note

- As per the recommendation of International council for Standardization in Hematology, the differential leucocyte counts are additionally being reported as absolute numbers of each cell in per unit volume of blood
- 2. Test conducted on EDTA whole blood









Collected





Name : Ms. USHA JAIN

156976105 Age: 7

Age: 72 Years Gender:

Received Female

: 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM

Reported : 9/4/2021 4:15:30PM

A/c Status : P Ref By : SELF Report Status : Final

Test Name	Results	Units	Bio. Ref. Interval
AMYLASE, SERUM	45.00	U/L	28.00 - 100.00
(IFCC)			

Comments

Lab No.

Amylase is produced in the Pancreas and most of the elevation in serum is due to increased rate of Amylase entry into the blood stream / decreased rate of clearance or both. Serum Amylase rises within 6 to 48 hours of onset of Acute pancreatitis in 80% of patients, but is not proportional to the severity of the disease. Activity usually returns to normal in 3-5 days in patients with milder edematous form of the disease. Values persisting longer than this period suggest continuing necrosis of pancreas or Pseudocyst formation. Approximately 20% of patients with Pancreatitis have normal or near normal activity. Hyperlipemic patients with Pancreatitis also show spuriously normal Amylase levels due to suppression of Amylase activity by triglyceride. Low Amylase levels are seen in Chronic Pancreatitis, Congestive Heart failure, 2nd & 3rd trimesters of pregnancy, Gastrointestinal cancer & bone fractures.

GLUCOSE, FASTING (F), PLASMA (Hexokinase)	100.00	mg/dL	70 - 100
CARDIO C-REACTIVE PROTEIN (hsCRP), SERUM (Immunoturbidimetry)	>10.00	mg/L	<1.00
APOLIPOPROTEINS A1 & B, SERUM (Immunoturbidimetry)			
Apolipoprotein (Apo A1)	120	mg/dL	105.00 - 205.00
Apolipoprotein (Apo B)	66	mg/dL	55.00 - 130.00
Apo B / Apo A1 Ratio	0.55		0.35 - 0.98

Comments

Apolipoprotein B is a more powerful independent predictor of Coronary Heart Disease (CAD) than LDL Cholesterol. It is useful in assessing the risk of CAD and to classify Hyperlipidemias. Apolipoprotein studies help in monitoring coronary bypass surgery patients with regard to risk and severity of re-stenosis. They are also useful in assessing risk of re-infarction in patients of Myocardial infarction.

Apolipoprotein A1 is one of the apoproteins of high density lipoproteins (HDL) which is inversely related to the risk of CAD. Individuals with Tangier disease have < 1% of normal Apo A1. Levels <90mg/dL indicate



Page 3 of 15











Name : Ms. USHA JAIN

USHA JAIN Collected

Collected : 9/4/2021 6:50:00AM Received : 9/4/2021 6:52:14AM

Lab No. : 156976105

Age: 72 Years Gender: Female

Received : 9/4/2021 6:52:14AN Reported : 9/4/2021 4:15:30PM

A/c Status : P

Ref By: SELF

Report Status : Final

Test Name Results Units Bio. Ref. Interval

increased risk of Atherosclerotic disease.

As per recommendations of National Cholesterol Education Program (NCEP) the clinical significance of results is as follows:

Apolipoprotein B

ļ	RESULT IN mg/dL	REMARKS
ļ	<23	Abetalipoproteinemia/Hypobetalipoproteinemia
	23-45	 Hypobetalipoproteinemia
ļ	46-135	Normal
İ	>135	 Hyperapobetalipoproteinemia/Increased CAD risk

Apo B to A1 Ratio

RATIO	REMARKS
0.35-0.98	Desirable
>0.98	Increased CAD risk

IRON STUDIES, SERUM (Spectrophotometry)			
Iron	63.00	ug/dL	50.00 - 170.00
Total Iron Binding Capacity (TIBC)	262.00	μg/dL	250.00 - 425.00
Transferrin Saturation	24.05	%	15.00 - 50.00

Comments

Iron is an essential trace mineral element which forms an important component of hemoglobin, metallocompounds and Vitamin A. Deficiency of iron, leads to microcytic hypochromic anemia. The toxic effects of iron are deposition of iron in various organs of the body and hemochromatosis.

Total Iron Binding capacity (TIBC) is a direct measure of the protein Transferrin which transports iron from the gut to storage sites in the bone marrow. In iron deficiency anemia, serum iron is reduced and TIBC increases.







Female



Collected





: 9/4/2021 6:50:00AM

Name : Ms. USHA JAIN

156976105

Is. USHA JAIN

Age: 72 Years

Received : 9/4/2021 6:52:14AM Reported : 9/4/2021 4:15:30PM

A/c Status : P Ref By : SELF Report Status : Final

Gender:

Test Name Results Units Bio. Ref. Interval

Transferrin Saturation occurs in Idiopathic hemochromatosis and Transfusional hemosiderosis where no unsaturated iron binding capacity is available for iron mobilization. Similar condition is seen in congenital deficiency of Transferrin.

 VITAMIN B12; CYANOCOBALAMIN, SERUM
 >2000
 pg/mL
 211.00 - 911.00

 (CLIA)

Notes

Lab No.

- 1. Interpretation of the result should be considered in relation to clinical circumstances.
- It is recommended to consider supplementary testing with plasma Methylmalonic acid (MMA) or
 plasma homocysteine levels to determine biochemical cobalamin deficiency in presence of clinical
 suspicion of deficiency but indeterminate levels. Homocysteine levels are more sensitive but MMA is
 more specific
- 3. False increase in Vitamin B12 levels may be observed in patients with intrinsic factor blocking antibodies, MMA measurement should be considered in such patients
- 4. The concentration of Vitamin B12 obtained with different assay methods cannot be used interchangeably due to differences in assay methods and reagent specificity

VITAMIN D, 25 - HYDROXY, SERUM	131.10	nmol/L	75.00 - 250.00
(CLIA)			

Interpretation

LEVEL	REFERENCE RANGE IN nmol/L	COMMENTS
Deficient	< 50 	High risk for developing bone disease
Insufficient 	 50-74 	Vitamin D concentration Which normalizes Parathyroid hormone concentration
Sufficient	75-250 	Optimal concentration for maximal health benefit
Potential intoxication	>250 	 High risk for toxic













Name : Ms. USHA JAIN

156976105 Age: 72 Years

Gender: Female

Collected Received Reported : 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

A/c Status : P Ref By : SELF Report Status : Final

Test Name Results Units Bio. Ref. Interval

Note

Lab No.

• The assay measures both D2 (Ergocalciferol) and D3 (Cholecalciferol) metabolites of vitamin D.

25 (OH)D is influenced by sunlight, latitude, skin pigmentation, sunscreen use and hepatic function.

• Optimal calcium absorption requires vitamin D 25 (OH) levels exceeding 75 nmol/L.

It shows seasonal variation, with values being 40-50% lower in winter than in summer.

• Levels vary with age and are increased in pregnancy.

• A new test Vitamin D, Ultrasensitive by LC-MS/MS is also available

Comments

Vitamin D promotes absorption of calcium and phosphorus and mineralization of bones and teeth. Deficiency in children causes Rickets and in adults leads to Osteomalacia. It can also lead to Hypocalcemia and Tetany. Vitamin D status is best determined by measurement of 25 hydroxy vitamin D, as it is the major circulating form and has longer half life (2-3 weeks) than 1,25 Dihydroxy vitamin D (5-8 hrs).

Decreased Levels

- · Inadequate exposure to sunlight
- Dietary deficiency
- Vitamin D malabsorption
- · Severe Hepatocellular disease
- Drugs like Anticonvulsants
- Nephrotic syndrome

Increased levels

Vitamin D intoxication













Rio Pof Intorval

Name : Ms. USHA JAIN

Lab No. : 156976105

A/c Status

Toet Namo

: 156976105 Age: 72 Years

Ref By: SELF

Gender: Female

Doculte

Collected Received Reported

Linite

: 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM

: 9/4/2021 4:15:30PM

Report Status : Final

Test Name	Results	Units	Bio. Ref. Interval
LIVER & KIDNEY PANEL, SERUM			
Bilirubin Total (DPD)	0.35	mg/dL	0.20 - 1.10
Bilirubin Direct (DPD)	0.13	mg/dL	<0.3
Bilirubin Indirect (Calculated)	0.22	mg/dL	<1.10
AST (SGOT) (IFCC without P5P)	17	U/L	9.00 - 36.00
ALT (SGPT) (IFCC without P5P)	12	U/L	10.00 - 49.00
GGTP (IFCC)	26	U/L	0 - 38
Alkaline Phosphatase (ALP) (IFCC-AMP)	61	U/L	30.00 - 120.00
Total Protein (Biuret)	5.90	g/dL	5.70 - 8.20
Albumin (BCG)	4.10	g/dL	3.20 - 4.60
A : G Ratio (Calculated)	2.28		0.90 - 2.00
Urease UV)	42.00	mg/dL	17.00 - 49.00
Creatinine (Modified Jaffe,Kinetic)	1.46	mg/dL	0.55 - 1.02













Name : Ms. USHA JAIN

; P

Lab No.

A/c Status

: 156976105

Age: 72 Years

Gender: Female

Received male Reported

Collected

: 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

Ref By: SELF Report Status : Final

Test Name Uric Acid (Uricase)	Results 4.50	Units mg/dL	Bio. Ref. Interval 2.60 - 6.00
Calcium, Total (Arsenazo III)	8.80	mg/dL	8.80 - 10.20
Phosphorus (Molybdate UV)	3.47	mg/dL	2.80 - 4.00
Sodium (Indirect ISE)	145.00	mEq/L	136.00 - 145.00
Potassium (Indirect ISE)	4.80	mEq/L	3.50 - 5.10
Chloride (Indirect ISE)	109.00	mEq/L	98.00 - 107.00

ADVICE: CKD RISK MAP

KDIGO guideline, 2012 recommends Chronic Kidney disease (CKD) should be classified based on cause, GFR category and albuminuria (ACR) category. GFR & ACR category combined together reflect risk of progression and helps clinician to identify individuals who are progressing at more rapid rate than anticipated







Female







Name : Ms. USHA JAIN

Lab No.

USHA JAIN Collected

Gender:

Age: 72 Years

Collected : 9/4/2021 6:50:00AM
Received : 9/4/2021 6:52:14AM
Reported : 9/4/2021 4:15:30PM

A/c Status : P Ref By : SELF

156976105

Report Status : Final

Light Yellow <=1.005 8.5 Negative Present 3+(1.0 g/dL)		Pale yellow 1.001 - 1.030 5.0 - 8.0 Negative
<=1.005 8.5 Negative		1.001 - 1.030 5.0 - 8.0 Negative
<=1.005 8.5 Negative		1.001 - 1.030 5.0 - 8.0 Negative
8.5 Negative		5.0 - 8.0 Negative
Negative		Negative
Present 3+(1.0 g/dL)		
		Negative
Negative		Negative
Negative		0.0 - 2.0 RBC/hpf
Negative		0-5 WBC / hpf
0-1 Epi Cells/hpf		0.0 - 5.0 Epi cells/hpf
None seen		None seen/Lpf
None seen		None seen
None seen		None seen
	Negative Negative Negative Negative Negative Negative Negative O-1 Epi Cells/hpf None seen None seen	Negative Negative Negative Negative Negative Negative Negative Negative Negative Negative Negative None seen

Result Rechecked, Please Correlate Clinically.



Page 9 of 15







Reported





: 9/4/2021 6:50:00AM

: 9/4/2021 6:52:14AM

: 9/4/2021 4:15:30PM

Name Ms. USHA JAIN

156976105

Lab No.

Collected Received Gender:

Age: 72 Years

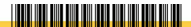
A/c Status Ref By: SELF **Report Status** : Final

Test Name Results **Units** Bio. Ref. Interval

IMPORTANT INSTRUCTIONS

Female

*Test results released pertain to the specimen submitted.*All test results are dependent on the quality of the sample received by the Laboratory *Laboratory investigations are only a tool to facilitate in arriving at a diagnosis and should be clinically correlated by the Referring Physician .*Sample repeats are accepted on request of Referring Physician within 7 days post reporting.*Report delivery may be delayed due to unforeseen circumstances. Inconvenience is regretted.*Certain tests may require further testing at additional cost for derivation of exact value. Kindly submit request within 72 hours post reporting.*Test results may show interlaboratory variations.*The Courts/Forum at Delhi shall have exclusive jurisdiction in all disputes/claims concerning the test(s) & or results of test(s).*Test results are not valid for medico legal purposes. *Contact customer care Tel No. +91-11-39885050 for all queries related to test results. (#) Sample drawn from outside source.













Name : Ms. USHA JAIN

156976105

Age: 72 Years

SELF

Ref By:

Gender: Female

Collected Received Reported : 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

Report Status : Final

Test Name	Results	Units	Bio. Ref. Interval
HbA1c (GLYCOSYLATED HEMOGLOBIN), BLOOD (HPLC, NGSP certified)			
HbA1c Estimated average glucose (eAG)	6.3 134	% mg/dL	4.00 - 5.60

Interpretation

Lab No.

A/c Status

HbA1c result is suggestive of at risk for Diabetes (Prediabetes)/ well controlled Diabetes in a known Diabetic

Note: Presence of Hemoglobin variants and/or conditions that affect red cell turnover must be considered, particularly when the HbA1C result does not correlate with the patient's blood glucose levels.

FACTORS THAT INTERFERE WITH HbA1C MEASUREMENT	FACTORS THAT AFFECT INTERPRETATION OF HBA1C RESULTS
Hemoglobin variants,elevated fetal hemoglobin (HbF) and chemically modified derivatives of hemoglobin (e.g. carbamylated Hb in patients with renal failure) can affect the accuracy of HbA1c measurements	Any condition that shortens erythrocyte survival or decreases mean erythrocyte age (e.g.,recovery from acute blood loss, hemolytic anemia, HbSS, HbCC, and HbSC) will falsely lower HbA1c test results regardless of the assay method used.Iron deficiency anemia is associated with higher HbA1c













Name : Ms. USHA JAIN

: 156976105

Age: 72 Years

Ref By: SELF

Gender: Female

Collected Received Reported : 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

Report Status : Final

Test Name	Results	Units	Bio. Ref. Interval
THYROID PROFILE,TOTAL, SERUM (CLIA)			
T3, Total	0.81	ng/mL	0.60 - 1.81
T4, Total	8.20	μg/dL	5.01 - 12.45
TSH	3.64	μIU/mL	0.35 - 5.50

Note

Lab No.

A/c Status

- 1. TSH levels are subject to circadian variation, reaching peak levels between 2 4.a.m. and at a minimum between 6-10 pm . The variation is of the order of 50% . hence time of the day has influence on the measured serum TSH concentrations.
- 2. Alteration in concentration of Thyroid hormone binding protein can profoundly affect Total T3 and/or Total T4 levels especially in pregnancy and in patients on steroid therapy.
- 3. Unbound fraction (Free,T4 /Free,T3) of thyroid hormone is biologically active form and correlate more closely with clinical status of the patient than total T4/T3 concentration
- 4. Values <0.03 uIU/mL need to be clinically correlated due to presence of a rare TSH variant in some individuals













Name : Ms. USHA JAIN

Lab No. : 156976105

A/c Status

976105 Age: 72 Years

Ref By: SELF

Gender: Female

Collected Received : 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM

Reported : 9/4/2021 4:15:30PM

Report Status : Final

Test Name	Results	Units	Bio. Ref. Interval
LIPID SCREEN, SERUM (Spectrophotometry)			
Cholesterol, Total	144.00	mg/dL	<200.00
Triglycerides	197.00	mg/dL	<150.00
HDL Cholesterol	41.30	mg/dL	>50.00
LDL Cholesterol, Calculated	63.30	mg/dL	<100.00
VLDL Cholesterol,Calculated	39.40	mg/dL	<30.00
Non-HDL Cholesterol *	103	mg/dL	<130

Interpretation

REMARKS 	TOTAL CHOLESTEROL in mg/dL	TRIGLYCERIDE in mg/dL	LDL CHOLESTEROL in mg/dL	NON HDL CHOLESTEROL in mg/dL
Optimal	<200	<150	<100	<130
Above Optimal			100-129	130 - 159
Borderline High	200-239	150-199	130-159	160 - 189
High	>=240	200-499	160-189	190 - 219
Very High	-	>=500	>=190	>=220

Note

- 1. Measurements in the same patient can show physiological & analytical variations. Three serial samples 1 week apart are recommended for Total Cholesterol, Triglycerides, HDL& LDL Cholesterol.
- NLA-2014 recommends a complete lipoprotein profile as the initial test for evaluating cholesterol.



Page 13 of 15

^{*} Not in NABL scope











Name : Ms. USHA JAIN

Collected

Female

: 9/4/2021 6:50:00AM

Lab No. ; 1

156976105 Age: 72 Years

Received Reported : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

A/c Status : P

Ref By: SELF

Report Status

: Final

Test Name Results Units Bio. Ref. Interval

Friedewald equation to calculate LDL cholesterol is most accurate when Triglyceride level is < 400 mg/dL. Measurement of Direct LDL cholesterol is recommended when Triglyceride level is > 400 mg/dL

Gender:

- NLA-2014 identifies Non HDL Cholesterol(an indicator of all atherogeniclipoproteins such as LDL, VLDL, IDL, Lpa, Chylomicron remnants)along with LDL-cholesterol as co- primary target for cholesterol lowering therapy. Note that major risk factors can modify treatment goals for LDL &Non HDI
- 5. Apolipoprotein B is an optional, secondary lipid target for treatment once LDL & Non HDL goals have been achieved
- 6. Additional testing for Apolipoprotein B, hsCRP,Lp(a) & LP-PLA2 should be considered among patients with moderate risk for ASCVD for risk refinement

Treatment Goals as per Lipid Association of India 2016

RISK	TREATMENT GOAL		CONSIDER THERAPY	
CATEGORY	LDL CHOLESTEROL (LDL-C)(mg/dL)	NON HDL CHLOESTEROL (NON HDL-C) (mg/dL)	LDL CHOLESTEROL (LDL-C)(mg/dL)	NON HDL CHLOESTEROL (NON HDL-C) (mg/dL)
Very High	<50		>=50	>=80
High	<70	<100	>=70	>=100
Moderate	<100	<130	>=100	>=130
Low	<100	<130	>=130*	>=160*

^{*}In low risk patient, consider therapy after an initial non-pharmacological intervention for at least 3 months

* Not in NABL scope



Page 14 of 15











Name : Ms. USHA JAIN

Lab No. : 156976105 A

Age: 72 Years

Female

Collected Received Reported : 9/4/2021 6:50:00AM : 9/4/2021 6:52:14AM : 9/4/2021 4:15:30PM

A/c Status : P Ref By : SELF Report Status : Final

Gender:

Test Name Results Units Bio. Ref. Interval

Dr.Kamal Modi

Dr.Kamal Modi MD, Biochemistry Consultant Biochemist NRL - Dr Lal PathLabs Ltd Dr Gurleen Oberoi

DM(Hematopathology), MD, DNB, MNAMS
Consultant & Technical Lead
-Hematopathology
NRL - Dr Lal Pathl abs Ltd

Brangeher

Dr Himangshu Mazumdar MD, Biochemistry Senior Consultant - Clinical Chemistry & Biochemical Genetics NRL - Dr Lal PathLabs Ltd

Kitu

Dr Ritu Nayar MD, Microbiology Deputy HOD - Microbiology & Serology NRL - Dr Lal PathLabs Ltd Dr Nimmi Kansal

MD, Biochemistry
National Head - Clinical Chemistry &
Biochemical Genetics

NRL - Dr Lal PathLabs Ltd

Dr Anil Arora
MD, Pathology
HOD Hematology &
Immunohematology
NRL - Dr Lal PathLabs Ltd

Dr Sunanda MD, Pathology Consultant

NRL - Dr Lal PathLabs Ltd

IMPORTANT INSTRUCTIONS

---End of report -

*Test results released pertain to the specimen submitted.*All test results are dependent on the quality of the sample received by the Laboratory.

*Laboratory investigations are only a tool to facilitate in arriving at a diagnosis and should be clinically correlated by the Referring Physician.*Sample repeats are accepted on request of Referring Physician within 7 days post reporting.*Report delivery may be delayed due to unforeseen circumstances. Inconvenience is regretted.*Certain tests may require further testing at additional cost for derivation of exact value. Kindly submit request within 72 hours post reporting.*Test results may show interlaboratory variations.*The Courts/Forum at Delhi shall have exclusive jurisdiction in all disputes/claims concerning the test(s) & or results of test(s).*Test results are not valid for medico legal purposes. *Contact customer care Tel No. +91-11-39885050 for all queries related to test results.

(#) Sample drawn from outside source.

* Not in NABL scope



Page 15 of 15