



Tests you can trust

Pco g : XXXXXXXXXXXXXXXXXXXX

F cvg : XXXXXXXXXX

VguV'Cungf : J aanch Diabetes Comprehensive

T gr qtv"Uvcwu : Complete Report



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to have **100%** of its Labs with  
**NABL Accreditation<sup>#</sup>**

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Patient Name : XXXXXXXXXXXXX  
 Referred By : XXXXXXXXXXXXX  
 Address : XXXXXXXXXXXXX

Tests Done : JAANCH DIABETES COMPREHENSIVE



## Report Availability Summary

**Note:** Please refer to the table below for status of your tests.

✓ 19 Ready
🟡 0 Ready with Cancellation
🔄 0 Processing
✗ 0 Cancelled in Lab

### TEST DETAILS

### REPORT STATUS

|                                              |         |
|----------------------------------------------|---------|
| <b>JAANCH DIABETES COMPREHENSIVE</b>         | Ready ✓ |
| CHLORIDE                                     | Ready ✓ |
| C-PEPTIDE                                    | Ready ✓ |
| FRUCTOSAMINE                                 | Ready ✓ |
| HIGH SENSITIVITY C-REACTIVE PROTEIN (HS-CRP) | Ready ✓ |
| LIPASE                                       | Ready ✓ |
| Lipoprotein (a) [Lp(a)]                      | Ready ✓ |
| HOMA INSULIN RESISTANCE INDEX                | Ready ✓ |
| SODIUM                                       | Ready ✓ |
| URINARY MICROALBUMIN                         | Ready ✓ |
| VITAMIN B-12                                 | Ready ✓ |
| HBA PROFILE                                  | Ready ✓ |
| HEMOGRAM - 6 PART (DIFF)                     | Ready ✓ |
| LIVER FUNCTION TESTS                         | Ready ✓ |
| ROUTINE URINE ANALYSIS                       | Ready ✓ |
| KIDPRO                                       | Ready ✓ |
| LIPID PROFILE                                | Ready ✓ |
| APOLIPROTEIN RATIO                           | Ready ✓ |
| AMYLASE                                      | Ready ✓ |
| BLOOD KETONE (D3HB)                          | Ready ✓ |

Patient Name : XXXXXXXXXXXXX  
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Tests Done : JAANCH DIABETES COMPREHENSIVE



## Tests Outside Reference Range

**Note:** Please refer to the table below for tests outside reference range.

| Test Name                                | Observed Value    | Units                 | Bio. Ref. Interval. |
|------------------------------------------|-------------------|-----------------------|---------------------|
| <b>COMPLETE HEMOGRAM</b>                 |                   |                       |                     |
| HEMATOCRIT(PCV)                          | <b>33.6</b>       | %                     | 40.0-50.0           |
| HEMOGLOBIN                               | <b>10.4</b>       | g/dL                  | 13.0-17.0           |
| LYMPHOCYTE                               | <b>17.4</b>       | %                     | 20-40               |
| MEAN CORP.HEMO.CONC(MCHC)                | <b>31</b>         | g/dL                  | 31.5-34.5           |
| MEAN CORPUSCULAR HEMOGLOBIN(MCH)         | <b>26.8</b>       | pq                    | 27.0-32.0           |
| RED CELL DISTRIBUTION WIDTH (RDW-CV)     | <b>15.4</b>       | %                     | 11.6-14             |
| RED CELL DISTRIBUTION WIDTH - SD(RDW-SD) | <b>48.9</b>       | fL                    | 39-46               |
| TOTAL RBC                                | <b>3.88</b>       | X 10 <sup>6</sup> /μL | 4.5-5.5             |
| <b>DIABETES</b>                          |                   |                       |                     |
| FRUCTOSAMINE                             | <b>322</b>        | μmol/L                | <=286               |
| HOMA INSULIN RESISTANCE INDEX            | <b>2.77</b>       | Index                 | <2.0                |
| <b>LIPID</b>                             |                   |                       |                     |
| LDL / HDL RATIO                          | <b>0.9</b>        | Ratio                 | 1.5-3.5             |
| TC/ HDL CHOLESTEROL RATIO                | <b>2.2</b>        | Ratio                 | 3 - 5               |
| <b>RENAL</b>                             |                   |                       |                     |
| CREATININE - SERUM                       | <b>0.7</b>        | mg/dL                 | 0.72-1.18           |
| <b>URINOGRAM</b>                         |                   |                       |                     |
| SPECIFIC GRAVITY                         | <b>&lt; 1.003</b> | -                     | 1.003-1.030         |
| <b>VITAMIN</b>                           |                   |                       |                     |
| VITAMIN B-12                             | <b>981</b>        | pg/mL                 | 211-911             |



Patient Name : XXXXXXXXXXXXX  
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Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                           | TECHNOLOGY | VALUE  | UNITS |
|-------------------------------------|------------|--------|-------|
| C-PEPTIDE<br>Bio. Ref. Interval. :- | E.C.L.I.A  | < 2.33 | ng/mL |

1.10 – 4.40 ng/ml

**Clinical Significance**

C-peptide, a polypeptide consisting of 31 amino acids (MW~3000), is stored in the secretory granules of the beta cells and released into circulation in equimolar amounts with insulin. The determination of C-peptide provides an assessment of endogenous insulin secretory reserves in patients with diabetes mellitus and is considered a more reliable indicator of insulin secretion than insulin itself. The primary indication for measuring C-peptide is for the evaluation of fasting hypoglycemia. It is also used to monitor patient's response to pancreatic surgery. C-peptide levels increase in insulinomas and beta-cell tumors.

Specifications: Precision: Intra assay (%CV): 2.9%, Inter assay (%CV): 3.6%; Sensitivity: 0.02 ng/ml

**Kit Validation reference:**

Clerk PM, Assays for insulin, proinsulin (s) and c-peptide. Ann clin biochem 1999;36(5):541-564

**Please correlate with clinical conditions.**

**Method:-** FULLY AUTOMATED ELECTROCHEMILUMINESCENCE IMMUNOASSAY

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



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Patient Name : XXXXXXXXXXXXX  
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Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          | TECHNOLOGY         | VALUE | UNITS |
|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------|-------|-------|
| APOLIPOPROTEIN - A1 (APO-A1)<br><b>Bio. Ref. Interval. :</b><br>Male : 86 - 152<br>Female : 94 - 162<br><b>Method :</b> FULLY AUTOMATED RATE IMMUNOTURBIDIMETRY - BECKMAN COULTER                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | IMMUNOTURBIDIMETRY | < 119 | mg/dL |
| APOLIPOPROTEIN - B (APO-B)<br><b>Bio. Ref. Interval. :</b><br>Male : 56 - 145<br>Female : 53 - 138<br><b>Method :</b> FULLY AUTOMATED RATE IMMUNOTURBIDIMETRY - BECKMAN COULTER                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | IMMUNOTURBIDIMETRY | < 101 | mg/dL |
| APO B / APO A1 RATIO (APO B/A1)<br><b>Bio. Ref. Interval. :</b><br>Male : 0.40 - 1.26<br>Female : 0.38 - 1.14<br><br>Clinical Significance :<br><ul style="list-style-type: none"><li>• Apolipoprotein B is a more potent and independent predictor of Coronary artery disease (CAD) than LDL Cholesterol.</li><li>• Apolipoprotein A1 is one of the apoproteins of HDL and is inversely related to risk of CAD.</li><li>• The Apolipoprotein studies help in monitoring risk of restenosis in patients with myocardial infarction, Coronary bypass surgery etc.</li><li>• An increased ratio of Apo B to A1 beyond the defined normal range is indicative of CAD risk.</li><li>• All results have to be interpreted in Conjunction with clinical history and other findings.</li></ul> <b>Method :</b> Derived from serum Apo A1 and Apo B values | CALCULATED         | 0.8   | Ratio |

**Please correlate with clinical conditions.**

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Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                              | TECHNOLOGY | VALUE | UNITS |
|----------------------------------------|------------|-------|-------|
| VITAMIN B-12<br>Bio. Ref. Interval. :- | C.L.I.A    | 981   | pg/mL |

Normal : 211 - 911 pg/ml

Clinical significance :

Vitamin B12 or cyanocobalamin, is a complex corrinoid compound found exclusively from animal dietary sources, such as meat, eggs and milk. It is critical in normal DNA synthesis, which in turn affects erythrocyte maturation and in the formation of myelin sheath. Vitamin-B12 is used to find out neurological abnormalities and impaired DNA synthesis associated with macrocytic anemias. For diagnostic purpose, results should always be assessed in conjunction with the patients medical history, clinical examination and other findings.

Specifications: Intra assay (%CV):5.0%, Inter assay (%CV):9.2 %;Sensitivity:45 pg/ml

Kit Validation reference:

Chen IW, Sperling MI, Heminger LA. Vitamin B12. In: Pesce AJ, Kaplan LA, eds. Methods in Clinical Chemistry. St. Louis: CV Mosby; 1987:569-73.

**Please correlate with clinical conditions.**

**Method:-** COMPETITIVE CHEMI LUMINESCENT IMMUNO ASSAY

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



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Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                   | TECHNOLOGY | VALUE   | UNITS |
|---------------------------------------------|------------|---------|-------|
| INSULIN - FASTING<br>Bio. Ref. Interval. :- | E.C.L.I.A  | < 12.45 | µU/mL |

Normal: 2.6 - 24.9 µU/mL

#### Clinical Significance

Type I (Insulin dependent: "Juvenile") diabetes is due to a destruction of the beta cells, with a consequence of absolute lack of insulin. In type II (Non insulin-dependent: "Maturity onset") diabetes, insulin resistance may play an important role; However after several years of evolution, beta-cells failure may occur, leading to a relative insulinopenia requiring, in some cases, insulin administration. Insulin resistance is associated with high circulation levels of the hormone. For diagnostic purpose, results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings.

#### Specifications:

Precision: Intra Assay (%CV): 4.3 %, Inter Assay (%CV): 5.3%; Sensitivity: 0.4 µU/mL.

#### Kit validation references:

Lang DA, Matthews DR, Peto J, et al. Cyclic oscillations of basal plasma glucose and insulin concentrations in human beings. N Engl J Med 1979;301:1023-1027.

#### Please correlate with clinical conditions.

Method:- FULLY AUTOMATED ELECTROCHEMILUMINESCENCE IMMUNOASSAY

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

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Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                            | TECHNOLOGY | VALUE  | UNITS |
|------------------------------------------------------|------------|--------|-------|
| BLOOD KETONE (D3HB)<br><b>Bio. Ref. Interval. :-</b> | PHOTOMETRY | < 1.51 | mg/dL |

0.21-2.81 mg/dL

**Clinical Significance:**

Three types of ketones can be produced in body D-3- Hydroxybutyrate, Acetoacetate and Acetone. D-3- Hydroxybutyrate accounts for approximately 75% of the ketone bodies. During periods of ketosis, D-3- Hydroxybutyrate increases more than the other two. It has been shown to be a better index of ketoacidosis. In diabetics, D-3- Hydroxybutyrate is needed for the assessment of the severity of diabetic coma and to calculate insulin requirements.

**Specification:**

Precision: Intra assay (%CV): 4.53, Inter assay (%CV): 2.9, Sensitivity: 10.41 mg/dL.

**Kit validation references:**

Mcmurray, C.H., Blanchflower, W.J., Rice, D.A., ClinChem., 1984;30:No.3.

**Please correlate with clinical conditions.**

**Method:-** ENZYMATIC (KINETIC)

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

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Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                              | TECHNOLOGY | VALUE | UNITS  |
|----------------------------------------|------------|-------|--------|
| FRUCTOSAMINE<br>Bio. Ref. Interval. :- | PHOTOMETRY | 322   | µmol/L |

Normal < 286 µmol/L

Clinical Significance:

Fructosamine assay is useful in monitoring the degree of glycemia over short-to-intermediate time frames (1-3 weeks) concentration greater than the established normal range is an indication of prolonged hyperglycemia of 1-3 weeks or longer. The higher fructosamine value, poorer is the degree of glycemia control.

Specifications:

Precision %CV : Intra assay %CV- 3.2% , Inter assay %CV-4.0%, Sensitivity:- 290 umol/L

Kit Validation Reference:

Howey JEA, Browning MCK, Fraser CG. Assay of serum fructosamine that minimizes standardization and matrix problems: Use to assess components of biological va-riation. Clin Chem 1987; 33: 269- 272.

**Please correlate with clinical conditions.**

**Method:-** NITROBLUE TETRAZOLIUM ASSAY (NBT)

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

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Patient Name : XXXXXXXXXXXXX  
Referred By : XXXXXXXXXXXXX  
Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                                | TECHNOLOGY         | VALUE | UNITS |
|----------------------------------------------------------|--------------------|-------|-------|
| Lipoprotein (a) [Lp(a)]<br><b>Bio. Ref. Interval. :-</b> | IMMUNOTURBIDIMETRY | 10    | mg/dL |

Adults : < 30.0 mg/dl

**Clinical Significance:**

Determination of LPA may be useful to guide management of individuals with a family history of CHD or with existing disease. The levels of LPA in the blood depends on genetic factors; The range of variation in a population is relatively large and hence for diagnostic purpose, results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings.

**Specifications:**

Precision %CV :- Intra assay %CV- 4.55% , Inter assay %CV-0.86 %

**Kit Validation Reference:**

Tietz NW, Clinical Guide to Laboratory Tests Philadelphia WB. Saunders 1995 : 442-444

**Please correlate with clinical conditions.**

**Method:-** LATEX ENHANCED IMMUNOTURBIDIMETRY

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

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Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                                                     | TECHNOLOGY         | VALUE | UNITS |
|-------------------------------------------------------------------------------|--------------------|-------|-------|
| HIGH SENSITIVITY C-REACTIVE PROTEIN (HS-CRP)<br><b>Bio. Ref. Interval. :-</b> | IMMUNOTURBIDIMETRY | 1.1   | mg/L  |

< 1.00 - Low Risk  
1.00 - 3.00 - Average Risk  
>3.00 - 10.00 - High Risk  
> 10.00 - Possibly due to Non-Cardiac Inflammation

Disclaimer: Persistent unexplained elevation of HSCRP >10 should be evaluated for non-cardiovascular etiologies such as infection, active arthritis or concurrent illness.

Clinical significance:

High sensitivity C- reactive Protein ( HSCRP) can be used as an independent risk marker for the identification of Individuals at risk for future cardiovascular Disease. A coronary artery disease risk assessment should be based on the average of two hs-CRP tests, ideally taken two weeks apart.

Kit Validation Reference:

- 1.Clinical management of laboratory data in medical practice 2003-3004, 207(2003).
- 2.Tietz : Textbook of Clinical Chemistry and Molecular diagnostics :Second edition :Chapter 47:Page no.1507- 1508.

**Please correlate with clinical conditions.**

**Method:-** FULLY AUTOMATED LATEX AGGLUTINATION – BECKMAN COULTER

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

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Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                | TECHNOLOGY | VALUE | UNITS |
|------------------------------------------|------------|-------|-------|
| AMYLASE<br><b>Bio. Ref. Interval. :-</b> | PHOTOMETRY | 69.34 | U/L   |

Adults : 28-100 U/L

**Interpretation:**

Lipemic Sera (Hypertriglyceridemia) may contain inhibitors, Which falsely depress results. About 20% of patients with Acute Pancreatitis have abnormal lipids. Normal serum amylase may occur in Pancreatitis, Especially relapsing and chronic pancreatitis. Moderate increases may be reported in normal pregnancy.

**Clinical Significance:**

Causes of high Serum Amylase include Acute Pancreatitis, Pancreatic Pseudocyst, Pancreatic Ascites, Pancreatic Abscess, Neoplasm in or adjacent to Pancreas, Trauma to Pancreas, and common Duct Stones. Nonpancreatic Causes include inflammatory salivary lesions (Eg, Mumps), Perforated Peptic Ulcer, Intestinal Obstruction, Biliary Tract Disease, Peritonitis, Acute Appendicitis, Diabetic Ketoacidosis, and Extrapaneatic Carcinomas. Amylase levels more than 25-fold the upper limit of normal are often found when metastatic tumors produce Ectopic Amylase.

**Specifications:**

Precision: Intra assay (%CV): 2.82, Inter assay (%CV): 2.49, Sensitivity: 10.9 U/L.

**Kit Validation References:**

Rauscher, E., et coll., Fresenius Z. Analyt. Chem. 324 (1986) 304-305.

**Please correlate with clinical conditions.**

**Method:-** ENZYMATIC COLORIMETRIC TEST

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



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Patient Name : XXXXXXXXXXXXX  
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Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                               | TECHNOLOGY | VALUE | UNITS |
|-----------------------------------------|------------|-------|-------|
| LIPASE<br><b>Bio. Ref. Interval. :-</b> | PHOTOMETRY | 40.09 | U/L   |

Adults : 5.0 - 60.0 U/L

Interpretation:

For diagnostic purposes, the results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings like serum amylase. Serum Lipase is usually normal in patients with elevated serum amylase, having peptic ulcer, salivary adenitis, inflammatory bowel disease, intestinal obstruction, and macroamylasemia. Lipemic sera may interfere with results.

Clinical Significance:

High serum Lipase is a specific marker for pancreatitis; after acute pancreatitis the Lipase activity increases within 4-8 hours, reaches a peak after 24 hours and decreases after 8 to 14 days. However, there is no correlation between the Lipase activity determined in serum and the extent of damage to the pancreas.

Specifications:

Precision: Intra assay (%CV): 3.35, Inter assay (%CV): 2.46, Sensitivity: 3.5 U/L.

Kit Validation References:

Tietz Nw Et Al. Lipase In Serum - The Elusive Enzyme: An Overview. Clin Chem 1993; 39:746-756.

**Please correlate with clinical conditions.**

**Method:-** ENZYMATIC COLORIMETRIC ASSAY

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



Patient Name : XXXXXXXXXXXXX  
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Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT): XXXXXXXXXXXXX  
Report Released on (RRT): XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                        | TECHNOLOGY        | VALUE      | UNITS        | Bio. Ref. Interval. |
|----------------------------------|-------------------|------------|--------------|---------------------|
| TOTAL CHOLESTEROL                | PHOTOMETRY        | 123        | mg/dL        | < 200               |
| HDL CHOLESTEROL - DIRECT         | PHOTOMETRY        | 56         | mg/dL        | 40-60               |
| LDL CHOLESTEROL - DIRECT         | PHOTOMETRY        | 53.5       | mg/dL        | < 100               |
| TRIGLYCERIDES                    | PHOTOMETRY        | 120        | mg/dL        | < 150               |
| <b>TC/ HDL CHOLESTEROL RATIO</b> | <b>CALCULATED</b> | <b>2.2</b> | <b>Ratio</b> | <b>3 - 5</b>        |
| TRIG / HDL RATIO                 | CALCULATED        | 2.12       | Ratio        | < 3.12              |
| <b>LDL / HDL RATIO</b>           | <b>CALCULATED</b> | <b>0.9</b> | <b>Ratio</b> | <b>1.5-3.5</b>      |
| HDL / LDL RATIO                  | CALCULATED        | 1.05       | Ratio        | > 0.40              |
| NON-HDL CHOLESTEROL              | CALCULATED        | 66.2       | mg/dL        | < 160               |
| VLDL CHOLESTEROL                 | CALCULATED        | 23.94      | mg/dL        | 5 - 40              |

**Please correlate with clinical conditions.**

#### Method :

CHOL - Cholesterol Oxidase, Esterase, Peroxidase  
HCHO - Direct Enzymatic Colorimetric  
LDL - Direct Measure  
TRIG - Enzymatic, End Point  
TC/H - Derived from serum Cholesterol and Hdl values  
TRI/H - Derived from TRIG and HDL Values  
LDL/ - Derived from serum HDL and LDL Values  
HD/LD - Derived from HDL and LDL values.  
NHDH - Derived from serum Cholesterol and HDL values  
VLDL - Derived from serum Triglyceride values

#### \*REFERENCE RANGES AS PER NCEP ATP III GUIDELINES:

| TOTAL CHOLESTEROL | (mg/dl) | HDL  | (mg/dl) | LDL             | (mg/dl) | TRIGLYCERIDES   | (mg/dl) |
|-------------------|---------|------|---------|-----------------|---------|-----------------|---------|
| DESIRABLE         | <200    | LOW  | <40     | OPTIMAL         | <100    | NORMAL          | <150    |
| BORDERLINE HIGH   | 200-239 | HIGH | >60     | NEAR OPTIMAL    | 100-129 | BORDERLINE HIGH | 150-199 |
| HIGH              | >240    |      |         | BORDERLINE HIGH | 130-159 | HIGH            | 200-499 |
|                   |         |      |         | HIGH            | 160-189 | VERY HIGH       | >500    |
|                   |         |      |         | VERY HIGH       | >190    |                 |         |

**Alert !!! 10-12 hours fasting is mandatory for lipid parameters. If not, values might fluctuate.**

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



Patient Name : XXXXXXXXXXXXX  
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Address : XXXXXXXXXXXXX

Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT): XXXXXXXXXXXXX  
Report Released on (RRT): XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                          | TECHNOLOGY | VALUE | UNITS | Bio. Ref. Interval. |
|------------------------------------|------------|-------|-------|---------------------|
| ALKALINE PHOSPHATASE               | PHOTOMETRY | 105.1 | U/L   | 45-129              |
| BILIRUBIN - TOTAL                  | PHOTOMETRY | 0.45  | mg/dL | 0.3-1.2             |
| BILIRUBIN -DIRECT                  | PHOTOMETRY | 0.1   | mg/dL | 0 - 0.20            |
| BILIRUBIN (INDIRECT)               | CALCULATED | 0.35  | mg/dL | 0-0.9               |
| GAMMA GLUTAMYL TRANSFERASE (GGT)   | PHOTOMETRY | 13.4  | U/L   | < 55                |
| ASPARTATE AMINOTRANSFERASE (SGOT ) | PHOTOMETRY | 24.3  | U/L   | < 35                |
| ALANINE TRANSAMINASE (SGPT)        | PHOTOMETRY | 32.9  | U/L   | < 45                |
| SGOT / SGPT RATIO                  | CALCULATED | 0.74  | Ratio | < 2                 |
| PROTEIN - TOTAL                    | PHOTOMETRY | 6.98  | gm/dL | 5.7-8.2             |
| ALBUMIN - SERUM                    | PHOTOMETRY | 4.14  | gm/dL | 3.2-4.8             |
| SERUM GLOBULIN                     | CALCULATED | 2.84  | gm/dL | 2.5-3.4             |
| SERUM ALB/GLOBULIN RATIO           | CALCULATED | 1.46  | Ratio | 0.9 - 2             |

**Please correlate with clinical conditions.**

**Method :**

ALKP - Modified IFCC method  
BILT - Diazonium salt DPD method  
BILD - Diazonium salt DPD method  
BILI - Derived from serum Total and Direct Bilirubin values  
GGT - Modified IFCC method  
SGOT - IFCC\* Without Pyridoxal Phosphate Activation  
SGPT - IFCC\* Without Pyridoxal Phosphate Activation  
OT/PT - Derived from SGOT and SGPT values.  
PROT - Biuret Method  
SALB - Albumin Bcg<sup>1</sup>method (Colorimetric Assay Endpoint)  
SEGB - DERIVED FROM SERUM ALBUMIN AND PROTEIN VALUES  
A/GR - Derived from serum Albumin and Protein values

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



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**First National Diagnostic Chain to have 100% of its Labs with NABL Accreditation<sup>#</sup>**

Patient Name : XXXXXXXXXXXXX  
Referred By : XXXXXXXXXXXXX  
Address : XXXXXXXXXXXXX

Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT): XXXXXXXXXXXXX  
Report Released on (RRT): XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME | TECHNOLOGY       | VALUE | UNITS  |
|-----------|------------------|-------|--------|
| CHLORIDE  | I.S.E - INDIRECT | 101   | mmol/L |

**Bio. Ref. Interval. :**  
ADULTS: 98-107 MMOL/L

Clinical Significance :

An increased level of blood chloride (called hyperchloremia) usually indicates dehydration, but can also occur with other problems that cause high blood sodium, such as Cushing syndrome or kidney disease. Hyperchloremia also occurs when too much base is lost from the body (producing metabolic acidosis) or when a person hyperventilates (causing respiratory alkalosis). A decreased level of blood chloride (called hypochloremia) occurs with any disorder that causes low blood sodium. Hypochloremia also occurs with congestive heart failure, prolonged vomiting or gastric suction, Addison disease, emphysema or other chronic lung diseases (causing respiratory acidosis), and with loss of acid from the body (called metabolic alkalosis).

**Method :** ION SELECTIVE ELECTRODE - INDIRECT

|        |                  |     |        |
|--------|------------------|-----|--------|
| SODIUM | I.S.E - INDIRECT | 137 | mmol/L |
|--------|------------------|-----|--------|

**Bio. Ref. Interval. :**  
Adults: 136-145 mmol/l

**Method :** ION SELECTIVE ELECTRODE - INDIRECT

**Please correlate with clinical conditions.**

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



Patient Name : XXXXXXXXXXXXX  
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Address : XXXXXXXXXXXXX

Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                  | TECHNOLOGY        | VALUE      | UNITS        | Bio. Ref. Interval. |
|----------------------------|-------------------|------------|--------------|---------------------|
| BLOOD UREA NITROGEN (BUN)  | PHOTOMETRY        | 12.6       | mg/dL        | 7.94 - 20.07        |
| <b>CREATININE - SERUM</b>  | <b>PHOTOMETRY</b> | <b>0.7</b> | <b>mg/dL</b> | <b>0.72-1.18</b>    |
| BUN / Sr.CREATININE RATIO  | CALCULATED        | 18         | Ratio        | 9:1-23:1            |
| UREA (CALCULATED)          | CALCULATED        | 26.96      | mg/dL        | Adult : 17-43       |
| UREA / SR.CREATININE RATIO | CALCULATED        | 38.52      | Ratio        | < 52                |
| CALCIUM                    | PHOTOMETRY        | 9.62       | mg/dL        | 8.8-10.6            |
| URIC ACID                  | PHOTOMETRY        | 6.8        | mg/dL        | 4.2 - 7.3           |

**Please correlate with clinical conditions.**

**Method :**

BUN - Kinetic UV Assay.  
SCRE - Creatinine Enzymatic Method  
B/CR - Derived from serum Bun and Creatinine values  
UREAC - Derived from BUN Value.  
UR/CR - Derived from UREA and Sr.Creatinine values.  
CALC - Arsenazo III Method, End Point.  
URIC - Uricase / Peroxidase Method

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



Patient Name : XXXXXXXXXXXXX  
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Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                                               | TECHNOLOGY | VALUE | UNITS          |
|-------------------------------------------------------------------------|------------|-------|----------------|
| EST. GLOMERULAR FILTRATION RATE (eGFR)<br><b>Bio. Ref. Interval. :-</b> | CALCULATED | 134   | mL/min/1.73 m2 |

> = 90 : Normal  
60 - 89 : Mild Decrease  
45 - 59 : Mild to Moderate Decrease  
30 - 44 : Moderate to Severe Decrease  
15 - 29 : Severe Decrease  
<15 : Kidney Failure

#### Clinical Significance

The normal serum creatinine reference interval does not necessarily reflect a normal GFR for a patient. Because mild and moderate kidney injury is poorly inferred from serum creatinine alone. Thus, it is recommended for clinical laboratories to routinely estimate glomerular filtration rate (eGFR), a "gold standard" measurement for assessment of renal function, and report the value when serum creatinine is measured for patients 18 and older, when appropriate and feasible. It cannot be measured easily in clinical practice, instead, GFR is estimated from equations using serum creatinine, age, race and sex. This provides easy to interpret information for the doctor and patient on the degree of renal impairment since it approximately equates to the percentage of kidney function remaining. Application of CKD-EPI equation together with the other diagnostic tools in renal medicine will further improve the detection and management of patients with CKD.

#### Reference

Levey AS, Stevens LA, Schmid CH, Zhang YL, Castro AF, 3rd, Feldman HI, et al. A new equation to estimate glomerular filtration rate. Ann Intern Med. 2009;150(9):604-12.

#### Please correlate with clinical conditions.

**Method:-** 2021 CKD EPI Creatinine Equation

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



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Address : XXXXXXXXXXXXX

Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                                             | TECHNOLOGY        | VALUE       | UNITS        |
|-----------------------------------------------------------------------|-------------------|-------------|--------------|
| <b>HOMA INSULIN RESISTANCE INDEX</b><br><b>Bio. Ref. Interval. :-</b> | <b>CALCULATED</b> | <b>2.77</b> | <b>Index</b> |

Normal : < 2.0

**Clinical Significance:**

Homeostatic Model Assessment for Insulin Resistance (HOMA-IR )-is an index used to determine insulin resistance .Obesity, lack of Physical activity are few of the factors which contribute to insulin resistance which may lead to increase the risk of Cardiovascular disease.It is recommended to see HOMA-IR index as a cause for concern, not as an exact diagnosis.

**Kit Validation Reference:**

Matthews, D. R.etc."Homeostasis model assessment: insulin resistance and beta-cell function from fasting plasma glucose and insulin concentrations in man" Diabetologia (July 1985).

**Please correlate with clinical conditions.**

**Method:-** Derived from Insulin and Blood Sugar values

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005003360/IT011

Doctor 1 Sign

Doctor 2 Sign



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Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                                                                                                                               | TECHNOLOGY | VALUE   | UNITS               |
|---------------------------------------------------------------------------------------------------------------------------------------------------------|------------|---------|---------------------|
| <b>DIABETES SCREEN (URINE)</b>                                                                                                                          |            |         |                     |
| URINARY MICROALBUMIN                                                                                                                                    | PHOTOMETRY | < 12.46 | µg/mL               |
| <b>Bio. Ref. Interval. :</b><br>Adults: Less than 25 µg/ml<br><b>Method :</b> Fully Automated Immuno Turbidometry                                       |            |         |                     |
| CREATININE - URINE                                                                                                                                      | PHOTOMETRY | < 149   | mg/dL               |
| <b>Bio. Ref. Interval. :</b><br>Male: 39 - 259 mg/dl<br>Female: 28 - 217 mg/dl<br><b>Method :</b> Creatinine Jaffe Method, Rate-Blanked and Compensated |            |         |                     |
| URI. ALBUMIN/CREATININE RATIO (UA/C)                                                                                                                    | CALCULATED | 8.4     | µg/mg of Creatinine |
| <b>Bio. Ref. Interval. :</b><br>Adults : Less than 30 µg/mg of Creatinine<br><b>Method :</b> Derived from Albumin and Creatinine values                 |            |         |                     |
| <b>Please correlate with clinical conditions.</b>                                                                                                       |            |         |                     |

Tests Done : JAANCH DIABETES COMPREHENSIVE

Report Remarks : Labcode:2005124173/IT011

Doctor 1 Sign

Doctor 2 Sign



Patient Name : XXXXXXXXXXXXX  
Referred By : XXXXXXXXXXXXX  
Address : XXXXXXXXXXXXX

Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT): XXXXXXXXXXXXX  
Report Released on (RRT): XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                             | METHODOLOGY          | VALUE             | UNITS     | Bio. Ref. Interval. |
|---------------------------------------|----------------------|-------------------|-----------|---------------------|
| <b>Complete Urinogram</b>             |                      |                   |           |                     |
| <b><u>Physical Examination</u></b>    |                      |                   |           |                     |
| VOLUME                                | Visual Determination | 1                 | mL        | -                   |
| COLOUR                                | Visual Determination | PALE YELLOW       | -         | Pale Yellow         |
| APPEARANCE                            | Visual Determination | CLEAR             | -         | Clear               |
| <b>SPECIFIC GRAVITY</b>               | <b>pKa change</b>    | <b>&lt; 1.003</b> | -         | <b>1.003-1.030</b>  |
| PH                                    | pH indicator         | 5                 | -         | 5-8                 |
| <b><u>Chemical Examination</u></b>    |                      |                   |           |                     |
| URINARY PROTEIN                       | PEI                  | ABSENT            | mg/dL     | Absent              |
| URINARY GLUCOSE                       | GOD-POD              | ABSENT            | mg/dL     | Absent              |
| URINE KETONE                          | Nitroprusside        | ABSENT            | mg/dL     | Absent              |
| URINARY BILIRUBIN                     | Diazo coupling       | ABSENT            | mg/dL     | Absent              |
| UROBILINOGEN                          | Diazo coupling       | Normal            | mg/dL     | <=0.2               |
| URINE BLOOD                           | Peroxidase reaction  | ABSENT            | -         | Absent              |
| NITRITE                               | Diazo coupling       | ABSENT            | -         | Absent              |
| LEUCOCYTE ESTERASE                    | Esterase reaction    | ABSENT            | -         | Absent              |
| <b><u>Microscopic Examination</u></b> |                      |                   |           |                     |
| URINARY LEUCOCYTES (PUS CELLS)        | Microscopy           | 2.5               | cells/HPF | 0-5                 |

(Reference : \*PEI - Protein error of indicator, \*GOD-POD - Glucose oxidase-peroxidase)

Tests Done : ROUTINE URINE ANALYSIS, URINARY MICROALBUMIN

Report Remarks : Labcode:2005124173/IT011

Doctor 1 Sign

Doctor 2 Sign



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Patient Name : XXXXXXXXXXXXX  
Referred By : XXXXXXXXXXXXX  
Address : XXXXXXXXXXXXX

Sample Collected on (SCT): XXXXXXXXXXXXX  
Sample Received on (SRT): XXXXXXXXXXXXX  
Report Released on (RRT): XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME | TECHNOLOGY | VALUE | UNITS |
|-----------|------------|-------|-------|
| HbA1c     | H.P.L.C    | 5.1   | %     |

**Bio. Ref. Interval. :**

**As per ADA Guidelines**

Below 5.7% : Normal  
5.7% - 6.4% : Prediabetic  
>=6.5% : Diabetic

**Guidance For Known Diabetics**

Below 6.5% : Good Control  
6.5% - 7% : Fair Control  
7.0% - 8% : Unsatisfactory Control  
>8% : Poor Control

**Method :** Fully Automated H.P.L.C method

AVERAGE BLOOD GLUCOSE (ABG) CALCULATED 100 mg/dL

**Bio. Ref. Interval. :**

90 - 120 mg/dl : Good Control  
121 - 150 mg/dl : Fair Control  
151 - 180 mg/dl : Unsatisfactory Control  
> 180 mg/dl : Poor Control

**Method :** Derived from HbA1c values

**Please correlate with clinical conditions.**

Tests Done : HBA PROFILE,HEMOGRAM

Report Remarks : Labcode:2005124175/IT011

Doctor 1 Sign

Doctor 2 Sign



Patient Name : XXXXXXXXXXXXX  
Referred By : XXXXXXXXXXXXX  
Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                                        | METHODOLOGY                  | VALUE       | UNITS                        | Bio. Ref. Interval. |
|--------------------------------------------------|------------------------------|-------------|------------------------------|---------------------|
| <b>HEMOGLOBIN</b>                                | <b>SLS-Hemoglobin Method</b> | <b>10.4</b> | <b>g/dL</b>                  | <b>13.0-17.0</b>    |
| <b>Hematocrit (PCV)</b>                          | <b>CPH Detection</b>         | <b>33.6</b> | <b>%</b>                     | <b>40.0-50.0</b>    |
| <b>Total RBC</b>                                 | <b>HF &amp; EI</b>           | <b>3.88</b> | <b>X 10<sup>6</sup>/μL</b>   | <b>4.5-5.5</b>      |
| Mean Corpuscular Volume (MCV)                    | Calculated                   | 86.6        | fL                           | 83.0-101.0          |
| <b>Mean Corpuscular Hemoglobin (MCH)</b>         | <b>Calculated</b>            | <b>26.8</b> | <b>pq</b>                    | <b>27.0-32.0</b>    |
| <b>Mean Corp.Hemo. Conc (MCHC)</b>               | <b>Calculated</b>            | <b>31</b>   | <b>g/dL</b>                  | <b>31.5-34.5</b>    |
| <b>Red Cell Distribution Width - SD (RDW-SD)</b> | <b>Calculated</b>            | <b>48.9</b> | <b>fL</b>                    | <b>39-46</b>        |
| <b>Red Cell Distribution Width (RDW - CV)</b>    | <b>Calculated</b>            | <b>15.4</b> | <b>%</b>                     | <b>11.6-14</b>      |
| RED CELL DISTRIBUTION WIDTH INDEX (RDWI)         | Calculated                   | 343.7       | -                            | *Refer Note below   |
| MENTZER INDEX                                    | Calculated                   | 22.3        | -                            | *Refer Note below   |
| <b>TOTAL LEUCOCYTE COUNT (WBC)</b>               | <b>HF &amp; FC</b>           | <b>7.77</b> | <b>X 10<sup>3</sup> / μL</b> | <b>4.0 - 10.0</b>   |
| <b>DIFFERENTIAL LEUCOCYTE COUNT</b>              |                              |             |                              |                     |
| Neutrophils Percentage                           | Flow Cytometry               | 75.2        | %                            | 40-80               |
| <b>Lymphocytes Percentage</b>                    | <b>Flow Cytometry</b>        | <b>17.4</b> | <b>%</b>                     | <b>20-40</b>        |
| Monocytes Percentage                             | Flow Cytometry               | 2.7         | %                            | 2-10                |
| Eosinophils Percentage                           | Flow Cytometry               | 3.9         | %                            | 1-6                 |
| Basophils Percentage                             | Flow Cytometry               | 0.5         | %                            | 0-2                 |
| Immature Granulocyte Percentage (IG%)            | Flow Cytometry               | 0.3         | %                            | 0-0.5               |
| Nucleated Red Blood Cells %                      | Flow Cytometry               | 0.1         | %                            | 0.0-5.0             |
| <b>ABSOLUTE LEUCOCYTE COUNT</b>                  |                              |             |                              |                     |
| Neutrophils - Absolute Count                     | Calculated                   | 5.84        | X 10 <sup>3</sup> / μL       | 2.0-7.0             |
| Lymphocytes - Absolute Count                     | Calculated                   | 1.35        | X 10 <sup>3</sup> / μL       | 1.0-3.0             |
| Monocytes - Absolute Count                       | Calculated                   | 0.21        | X 10 <sup>3</sup> / μL       | 0.2 - 1.0           |
| Basophils - Absolute Count                       | Calculated                   | 0.04        | X 10 <sup>3</sup> / μL       | 0.02 - 0.1          |
| Eosinophils - Absolute Count                     | Calculated                   | 0.3         | X 10 <sup>3</sup> / μL       | 0.02 - 0.5          |
| Immature Granulocytes (IG)                       | Calculated                   | 0.02        | X 10 <sup>3</sup> / μL       | 0-0.3               |
| Nucleated Red Blood Cells                        | Calculated                   | 0.01        | X 10 <sup>3</sup> / μL       | 0.0-0.5             |
| <b>PLATELET COUNT</b>                            | <b>HF &amp; EI</b>           | <b>260</b>  | <b>X 10<sup>3</sup> / μL</b> | <b>150-410</b>      |
| Mean Platelet Volume (MPV)                       | Calculated                   | 10.3        | fL                           | 6.5-12              |
| Platelet Distribution Width (PDW)                | Calculated                   | 11.2        | fL                           | 9.6-15.2            |
| Platelet to Large Cell Ratio (PLCR)              | Calculated                   | 26.6        | %                            | 19.7-42.4           |
| Plateletcrit (PCT)                               | Calculated                   | 0.27        | %                            | 0.19-0.39           |

\*Note - Mentzer index (MI), RDW-CV and RDWI are hematological indices to differentiate between Iron Deficiency Anemia (IDA) and Beta Thalassemia Trait (BTT). MI >13, RDWI >220 and RDW-CV >14 more likely to be IDA. MI <13, RDWI <220, and RDW-CV <14 more likely to be BTT. Suggested Clinical correlation. BTT to be confirmed with HB electrophoresis if clinically indicated.

Method : Fully automated bidirectional analyser

(Reference : \*FC- flowcytometry, \*HF- hydrodynamic focussing, \*EI- Electric Impedance, \*Hb- hemoglobin, \*CPH- Cumulative pulse height)

Tests Done : HBA PROFILE,HEMOGRAM

Doctor 1 Sign

Doctor 2 Sign

Report Remarks : Labcode:2005124175/IT011



Patient Name : XXXXXXXXXXXXX  
Referred By : XXXXXXXXXXXXX  
Address : XXXXXXXXXXXXX

Sample Collected on (SCT) : XXXXXXXXXXXXX  
Sample Received on (SRT) : XXXXXXXXXXXXX  
Report Released on (RRT) : XXXXXXXXXXXXX  
Sample Type | Barcode : XXXXXXXXXXXXX

| TEST NAME                    | TECHNOLOGY | VALUE | UNITS |
|------------------------------|------------|-------|-------|
| FASTING BLOOD SUGAR(GLUCOSE) | PHOTOMETRY | 90.1  | mg/dL |

**Bio. Ref. Interval. :-**

| As per ADA Guideline: Fasting Plasma Glucose (FPG) |                        |
|----------------------------------------------------|------------------------|
| Normal                                             | 70 to 100 mg/dl        |
| Prediabetes                                        | 100 mg/dl to 125 mg/dl |
| Diabetes                                           | 126 mg/dl or higher    |

**Note :**

The assay could be affected mildly and may result in anomalous values if serum samples have heterophilic antibodies, hemolyzed, icteric or lipemic. The concentration of Glucose in a given specimen may vary due to differences in assay methods, calibration and reagent specificity. For diagnostic purposes results should always be assessed in conjunction with patients medical history, clinical findings and other findings.

**Please correlate with clinical conditions.**

**Method:-** GOD-POD METHOD

~~ End of report ~~

Tests Done : BLOOD SUGAR (F)

Report Remarks : Labcode:2005124176/IT011

Doctor 1 Sign

Doctor 2 Sign

## CONDITIONS OF REPORTING

- ✓ The reported results are for information and interpretation of the referring doctor only.
- ✓ It is presumed that the tests performed on the specimen belong to the patient; named or identified.
- ✓ Results of tests may vary from laboratory to laboratory and also in some parameters from time to time for the same patient.
- ✓ Should the results indicate an unexpected abnormality, the same should be reconfirmed.
- ✓ Only such medical professionals who understand reporting units, reference ranges and limitations of technologies should interpret results.
- ✓ This report is not valid for medico-legal purpose.
- ✓ Neither Thyrocare, nor its employees/representatives assume: (a) any liability, responsibility for any loss or damage that may be incurred by any person as a result of presuming the meaning or contents of the report, (b) any claims of any nature whatsoever arising from or relating to the performance of the requested tests as well as any claim for indirect, incidental or consequential damages. The total liability, in any case, of Thyrocare shall not exceed the total amount of invoice for the services provided and paid for.
- ✓ Thyrocare Discovery video link :- <https://youtu.be/nbdYeRgYyQc>

## EXPLANATIONS

- ✓ Majority of the specimen processed in the laboratory are collected by Pathologists and Hospitals we call them as "Clients".
- ✓ **Name** - The name is as declared by the client and recored by the personnel who collected the specimen.
- ✓ **Ref.Dr** - The name of the doctor who has recommended testing as declared by the client.
- ✓ **Labcode** - This is the accession number in our laboratory and it helps us in archiving and retrieving the data.
- ✓ **Barcode** - This is the specimen identity number and it states that the results are for the specimen bearing the barcode (irrespective of the name).
- ✓ **SCP** - Specimen Collection Point - This is the location where the blood or specimen was collected as declared by the client.
- ✓ **SCT** - Specimen Collection Time - The time when specimen was collected as declared by the client.
- ✓ **SRT** - Specimen Receiving Time - This time when the specimen reached our laboratory.
- ✓ **RRT** - Report Releasing Time - The time when our pathologist has released the values for Reporting.
- ✓ **Reference Range** - Means the range of values in which 95% of the normal population would fall.

## SUGGESTIONS

- ✓ Values out of reference range requires reconfirmation before starting any medical treatment.
- ✓ Retesting is needed if you suspect any quality shortcomings.
- ✓ Testing or retesting should be done in accredited laboratories.
- ✓ For suggestions, complaints, clinical support or feedback, write to us at [customersupport@thyrocare.com](mailto:customersupport@thyrocare.com) or call us on **022-3090 0000**

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| Thyroid | Diabetes | STDs   | Skin Care                     |
|         |          |        | Hair Fall                     |

Tests you can trust

\* T&C Apply, #As on 5th December 2024 (Applicable for all company owned labs except Bhagalpur & Vijayawada),

\* As per survey on doctors' perception of laboratory diagnostics (IJARIIT, 2023), -Mumbai Reference Lab is CAP Accredited