PKR JAIN HEALTHCARE INSTITUTE NASIRPUR, Hissar Road, AMBALA CITY- (Haryana)

A PIONEER DIAGNOSTIC CENTRE

🔽 0171-2532620, 8222896961 🛛 🖾 pkrjainhealthcare@gmail.com

| RI | ATIENT ID EG. NO./LAB NO. EGISTRATION DATE | : 1693137 : 122412070009 |
|--------------------|--|---|
| RI | | : 122412070009 |
| | ECISTRATION DATE | |
| | EUISTIKATION DATE | :07/Dec/2024 11:04 AM |
| CO | OLLECTION DATE | :07/Dec/2024 11:07AM |
| TUTE R I | EPORTING DATE | :07/Dec/2024 12:30PM |
| BALA CITY - HARY | ANA | |
| | | |
| Value | Unit | Biological Reference interval |
| L CHEMISTI | RY/BIOCHEMIST | RY |
| GLUCOSE F A | ASTING (F) | |
| 84.3 | mg/dL | NORMAL: < 100.0 PREDIABETIC: 100.0 - 125.0 DIABETIC: > 0R = 126.0 |
| | ON GUIDELINES: | DN GUIDELINES: |

A fasting plasma glucose level below 100 mg/dl is considered normal.
A fasting plasma glucose level between 100 - 125 mg/dl is considered as glucose intolerant or prediabetic. A fasting and post-prandial blood test (after consumption of 75 gms of glucose) is recommended for all such patients.
A fasting plasma glucose level of above 125 mg/dl is highly suggestive of diabetic state. A repeat post-prandial is strongly recommended for all such patients. A fasting plasma glucose level in excess of 125 mg/dl on both occasions is confirmatory for diabetic state.





DR.VINAY CHOPRA CONSULTANT PATHOLOGIST MBBS, MD (PATHOLOGY & MICROBIOLOGY)

DR.YUGAM CHOPRA CONSULTANT PATHOLOGIST MBBS, MD (PATHOLOGY)



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| NAME | : Mr. OM PARKASH | | | | |
|---|-----------------------------------|--|--------------------------|--|--|
| AGE/ GENDER | : 65 YRS/MALE | | PATIENT ID | : 1693137 | |
| COLLECTED BY | : | | REG. NO./LAB NO. | : 122412070009 | |
| REFERRED BY | : | | REGISTRATION DATE | :07/Dec/2024 11:04 AM | |
| BARCODE NO. | : 12506044 | | COLLECTION DATE | :07/Dec/2024 11:07AM | |
| CLIENT CODE. | : P.K.R JAIN HEALTHCARE INS | STITUTE | REPORTING DATE | :07/Dec/2024 12:30PM | |
| CLIENT ADDRESS | : NASIRPUR, HISSAR ROAD, A | : NASIRPUR, HISSAR ROAD, AMBALA CITY - HARYANA | | | |
| Test Name | | Value | Unit | Biological Reference interval | |
| | | LIPID PR | OFILE : BASIC | | |
| CHOLESTEROL TO | TAL: SERUM | 117.51 | mg/dL | OPTIMAL: < 200.0 | |
| by CHOLESTEROL O | | | 0 | BORDERLINE HIGH: 200.0 - 239.0 HIGH CHOLESTEROL: > OR = 240.0 | |
| TRIGLYCERIDES: S by GLYCEROL PHOSE | ERUM PHATE OXIDASE (ENZYMATIC) | 121.51 | mg/dL | OPTIMAL: < 150.0 BORDERLINE HIGH: 150.0 - 199.0 HIGH: 200.0 - 499.0 VERY HIGH: > OR = 500.0 | |
| HDL CHOLESTERO by SELECTIVE INHIBIT | L (DIRECT): SERUM 710N | 36.44 | mg/dL | LOW HDL: < 30.0 BORDERLINE HIGH HDL: 30.0 60.0 HIGH HDL: > OR = 60.0 | |
| LDL CHOLESTERO | | 56.77 | mg/dL | OPTIMAL: < 100.0 ABOVE OPTIMAL: 100.0 - 129. BORDERLINE HIGH: 130.0 - 159.0 HIGH: 160.0 - 189.0 VERY HIGH: > OR = 190.0 | |
| NON HDL CHOLES' by calculated, spe | | 81.07 | mg/dL | OPTIMAL: < 130.0 ABOVE OPTIMAL: 130.0 - 159 BORDERLINE HIGH: 160.0 - 189.0 HIGH: 190.0 - 219.0 VERY HIGH: > OR = 220.0 | |
| VLDL CHOLESTER | | 24.3 | mg/dL | 0.00 - 45.00 | |
| TOTAL LIPIDS: SEF by CALCULATED, SPE | RUM | 356.53 | mg/dL | 350.00 - 700.00 | |
| CHOLESTEROL/HI by CALCULATED, SPE | | 3.22 | RATIO | LOW RISK: 3.30 - 4.40 AVERAGE RISK: 4.50 - 7.0 MODERATE RISK: 7.10 - 11.0 HIGH RISK: > 11.0 | |



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| | | | |

| Test Name | Value | Unit | Biological Reference interval |
|--|-------|-------|---|
| LDL/HDL RATIO: SERUM by CALCULATED, SPECTROPHOTOMETRY | 1.56 | RATIO | LOW RISK: 0.50 - 3.0 MODERATE RISK: 3.10 - 6.0 HIGH RISK: > 6.0 |
| TRIGLYCERIDES/HDL RATIO: SERUM by CALCULATED, SPECTROPHOTOMETRY | 3.33 | RATIO | 3.00 - 5.00 |

INTERPRETATION:

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.

2. As per NLA-2014 guidelines, all adults above the age of 20 years should be screened for lipid status. Selective screening of children above the age of 2 years with a family history of premature cardiovascular disease or those with at least one parent with high total cholesterol is recommended.

3. Low HDL levels are associated with increased risk for Atherosclerotic Cardiovascular disease (ASCVD) due to insufficient HDL being available

to participate in reverse cholesterol transport, the process by which cholesterol is eliminated from peripheral tissues. 4. 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 HDL.

5. Additional testing for Apolipoprotein B, hsCRP,Lp(a) & LP-PLA2 should be considered among patients with moderate risk for ASCVD for risk refinement





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| Test Name | | Value | Unit | Biological Reference interval | |
| | | URIC | ACID | | |
| URIC ACID: SERUM | ſ | 4.97 | mg/dL | 3.60 - 7.70 | |
| by URICASE - OXIDAS | | 4.37 | iiig/ uL | 3.00 - 1.10 | |
| 5.Psoriasis. 6.Sickle cell anaemia (B).DUE TO DECREASI 1.Alcohol ingestion. 2.Thiazide diuretics. 3.Lactic acidosis. 4.Aspirin ingestion (I | ED EXCREATION (BY KIDNEYS) ess than 2 grams per day). isis or starvation. | | | | |
| 6.Renal failure due to | o any cause ere. | | | | |
| 6.Renal failure due to DECREASED:- (A).DUE TO DIETARY I | DEFICIENCY | | | | |
| 6.Renal failure due to DECREASED:- (A).DUE TO DIETARY I 1.Dietary deficiency 2.Fanconi syndrome | DEFICIENCY of Zinc, Iron and molybdenum. & Wilsons disease. | | | | |
| 6.Renal failure due to DECREASED:- (A).DUE TO DIETARY I 1.Dietary deficiency 2.Fanconi syndrome 3.Multiple sclerosis 4.Syndrome of inapp (B).DUE TO INCREASE | DEFICIENCY of Zinc, Iron and molybdenum. & Wilsons disease. ropriate antidiuretic hormone (SIA D EXCREATION | | | | |
| 6.Renal failure due to DECREASED:- (A).DUE TO DIETARY I 1.Dietary deficiency 2.Fanconi syndrome 3.Multiple sclerosis 4.Syndrome of inapp (B).DUE TO INCREASE | DEFICIENCY of Zinc, Iron and molybdenum. & Wilsons disease. ropriate antidiuretic hormone (SIA D EXCREATION | | | ls and ACTH, anti-coagulants and estrogens e | |





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