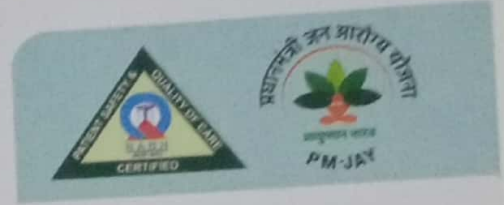




ALEXIS

HOSPITAL BHOPAL



DEPARTMENT OF PATHOLOGY

Date :

Patient Name : MR. ALBIN LONA
 Registration No. : AH-A-001695
 Lab No : 4
 Age & Sex : 31 Years / Male
 Referring Doctor :
 Report Status : Final

Registration Date/Time : 23/07/2022 09:51AM
 Accession Date/Time : 23/07/2022 09:58AM
 Report Date/Time : 24/07/2022 12:13PM
 Print Date/Time : 24/07/2022 12:14 pm

TEST(S)	RESULT(S)	UNITS	BIOLOGICAL REFERENCE RANGE
LIPID PROFILE			
SAMPLE TYPE : SERUM METHOD :			
S.Cholesterol	: 215.5	mg/dL	No risk < 200 mg/dl Moderate risk 200 - 239 mg/dl High risk >240 mg/dl Upto 150
S.Triglycerides	: 123.1	mg/dl	
HDL Cholesterol	: 48.1	mg/dl	Major risk < 40 mg/dl Negative risk > 60 mg/dl
LDL Cholesterol	: 142.78	mg/dl	Optimum < 100 mg/dl Near/above optimum 100 - 129 mg/dl Boderline high 130 - 159 mg/dl High 160 - 189 mg/dl Very high > 190 mg/dl Upto 30
VLDL Cholesterol	: 24.62	mg/dl	
S.Cholesterol/HDL Ratio	: 4.4802		4.4-11



***** End Of Report *****

Dr. Rajesh Kumar Chaurasia
 MBBS, M.D. (Pathology)
 Reg. No. MP-2808

Technician

Subject of following conditions :- The science of pathological diagnosis is based on interpretation of immunobiochemical ELISA, RIA and other reactions which are influenced by several factors. Assays are performed in accordance with standard procedures. The reported results are dependent on individual assay methods, equipment, specificity, sensitivity, drug interaction and the quality of the specimen (s) samples receiver. The most sophisticated computerized blood analyzer techniques are subject to unaccountable variations. Results of test vary from laboratory to laboratory and also in some parameters from time to time for the same patients. So the final diagnosis, conclusion should not be draw from these test only hence kindly correlate clinically. Adviced follow up. The laboratory investigations should always be interpreted in the light of clinical features and other related investigations. Subject to corrections of typing / printing / humanly mistakes. Shuold the result indicate an unexpected abnormality, reconfirmation shall be sought. This report is not vailed for



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TEST(S)

BLOOD GROUP & RH FACTOR

SAMPLE TYPE : EDTA WB
 METHOD : Slide and Tube m

Blood Group : * O *
 Rh Factor : POSITIVE

FBS

SAMPLE TYPE : WHOLE BLOOD FLUORIDE F
 METHOD : Hexokinase Enzym

Result : 122.6 mg/dl 70-100

LFT

SAMPLE TYPE : SERUM
 METHOD :

Bilirubin- Total : 0.35 mg/dL 0.15-1.1
 Bilirubin- Direct : 0.18 mg/dL 0-0.2
 Billirubin- Indirect : 0.17
 SGOT : 19.6 U/L 05-50
 SGPT : 16.4 U/L 07-40
 Alkaline Phosphatase : 100.9 U/L 5-128

RFT

SAMPLE TYPE : SERUM
 METHOD :

BLOOD UREA : 28.6 mg/dL 20-45
 SERUM CREATININE : 1.1 mg/dL 0.67-1.20

URIC ACID

SAMPLE TYPE : SERUM
 METHOD : Uricase POD End

Result : 6.3 mg/dL 3.4-7.0

***** End Of Report *****



Technician

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