



	MD (F	/inay Chopra Pathology & Microbiology) man & Consultant Pathologi		(Pathology)
NAME	: Mr. JAGTAR SING	H		
AGE/ GENDER	: 32 YRS/MALE		PATIENT ID	: 1654835
COLLECTED BY	:		REG. NO./LAB NO.	: 012410270045
REFERRED BY	:		REGISTRATION DATE	: 27/Oct/2024 01:48 PM
BARCODE NO.	:01519657		COLLECTION DATE	: 27/Oct/2024 01:57PM
CLIENT CODE.	: KOS DIAGNOSTIC	LAB	REPORTING DATE	: 27/Oct/2024 03:10PM
CLIENT ADDRESS	: 6349/1, NICHOLS	ON ROAD, AMBALA CANT	ſ	
T N		Value	Unit	Pialagical Defenses interval
Test Name		Value	Unit	Biological Reference interval
1 est Name			STRY/BIOCHEMIST	
I est Name		CLINICAL CHEMIS		
CHOLESTEROL TO by CHOLESTEROL OF		CLINICAL CHEMIS	STRY/BIOCHEMIST	

KOS Diagnostic Lab (A Unit of KOS Healthcare)

NATIONAL LIPID ASSOCIATION RECOMMENDATIONS (NLA-2014)	CHOLESTEROL IN ADULTS (mg/dL)	CHOLESTEROL IN ADULTS (mg/dL)
DESIRABLE	< 200.0	< 170.0
BORDERLINE HIGH	200.0 - 239.0	171.0 - 199.0
HIGH	>= 240.0	>= 200.0

NOTE:

 Molection
 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.
 As per National Lipid association - 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. high total cholesterol is recommended.





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

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







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Test Name		Value	Unit	Biological Reference interval
		U	RIC ACID	
URIC ACID: SERIIM			RIC ACID	3 60 - 7 70
by URICASE - OXIDAS <u>INTERPRETATION:-</u> 1.GOUT occurs when 2.Uric Acid is the end intestinal tract by mi INCREASED:- (A).DUE TO INCREASE 1.Idiopathic primary	E PEROXIDASE high levels of Uric A product of purine r crobial degradatior D PRODUCTION:- gout.	4.64 Acid in the blood cause crys netabolism . Uric acid is exc	mg/dL tals to form & accumulate ar	3.60 - 7.70 ound a joint. e kidneys and to a smaller degree in the
INTERPRETATION:- 1.GOUT occurs when 2.Uric Acid is the end intestinal tract by mi INCREASED:- (A).DUE TO INCREASE 1.Idiopathic primary 2.Excessive dietary po	E PEROXIDASE high levels of Uric A product of purine r crobial degradation D PRODUCTION:- gout. urines (organ meats of malignancies es & myeloid metaplas etc. D EXCREATION (BY H ess than 2 grams pensis or starvation. any cause etc. DEFICIENCY of Zinc, Iron and mo & Wilsons disease.	4.64 Acid in the blood cause crys netabolism . Uric acid is exc ulegumes, anchovies, etc). becially leukemais & lymph ia. IDNEYS)	mg/dL tals to form & accumulate ar reted to a large degree by the	ound a joint.





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 KOS Central Lab: 6349/1, Nicholson Road, Ambala Cantt -133 001, Haryana

 KOS Molecular Lab: IInd Floor, Parry Hotel, Staff Road, Opp. GPO, Ambala Cantt -133 001, Haryana

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	Dr. Vinay Chopra MD (Pathology & Micro Chairman & Consultant	biology)		(Pathology)
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Test Name		Value	Unit	Biological Reference interval
	IMMUNO	OPATHO	DLOGY/SEROLOGY	<i>t</i>
	RHEUMATOID FA	CTOR (R	A): QUANTITATIVE	- SERUM
RHEUMATOID (RA) SERUM by NEPHLOMETRY	FACTOR QUANTITATIVE:	5.3	IU/mL	NEGATIVE: < 18.0 BORDERLINE: 18.0 - 25.0 POSITIVE: > 25.0
RHEUMATOID ARTHIR 1. Rheumatoid Arthir membrane lining (syr 2. The disease spreda 3. The diagnosis of R. measurement of RA fa CAUTION (FALSE POS 1. RA factor is not spe 2. Non rheumatoid an RA patients have a no 3. Patients have a no 3. Patients with variou lupus erythematosus, 4. Anti-CCP have been specific (98%) than RA 5. Upto 30 % of patier	itis is a systemic autoimmune disease hovium) joints which ledas to progress as from small to large joints, with great A is primarily based on clinical, radiolo actor. TIVE):- cific for Rheumatoid arthiritis, as it is oft d rheumatoid arthritis (RA) populations nreactive titer and 8% of nonrheumatoid us nonrheumatoid diseases, characterized polymyositis, tuberculosis, syphilis, viral discovered in joints of patients with RA, factor. hts with Seronegative Rheumatoid arthir ive value of Anti-CCP antibodies for Rheu	that is mult ive joint des est damage gical & imm en present ii are not clead d patients ha d by chronic hepatitis, in but not in o ritis also sho	i-functional in origin and i struction and in most case in early phase. nunological features. The n n healthy individuals with o rly separate with regard to ave a positive titer). inflammation may have pos fectious mononucleosis, an ther form of joint disease. A w Anti-CCP antibodies. hiritis is far greater than Rh	ther autoimmune diseases and chronic infections the presence of rheumatoid factor (RF) (15% of sitive tests for RF. These diseases include systemic d influenza. nti-CCP2 is HIGHLY SENSITIVE (71%) & more
	L		port	
	there -		hopra	

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

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