

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT



	Dr. Vinay Chopr MD (Pathology & Micr Chairman & Consultar	robiology)	Dr. Yugam MD ( CEO & Consultant	(Pathology)
NAME	: Mr. RAJESH BABU			
AGE/ GENDER	: 62 YRS/MALE	]	PATIENT ID	: 1675875
COLLECTED BY	:	]	REG. NO./LAB NO.	: 012411190026
<b>REFERRED BY</b>	:	]	REGISTRATION DATE	: 19/Nov/2024 10:44 AM
BARCODE NO.	: 01521078		COLLECTION DATE	: 19/Nov/2024 10:47AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		REPORTING DATE	: 19/Nov/2024 11:07AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMB	ALA CANTT		
Test Name		Value	Unit	Biological Reference interval
	SWAST	THYA WE	LLNESS PANEL: D	
	COME	PLETE BLO	OD COUNT (CBC)	
<u>RED BLOOD CELL</u>	S (RBCS) COUNT AND INDICES			
HAEMOGLOBIN (H		13.5	gm/dL	12.0 - 17.0
by CALORIMETRIC RED BLOOD CELL		4.82	Millions/	cmm 3.50 - 5.00
	FOCUSING, ELECTRICAL IMPEDENCE			
PACKED CELL VOL	UME (PCV) AUTOMATED HEMATOLOGY ANALYZER	42.7	%	40.0 - 54.0
	AR VOLUME (MCV) AUTOMATED HEMATOLOGY ANALYZER	88.4	fL	80.0 - 100.0
MEAN CORPUSCUI	LAR HAEMOGLOBIN (MCH) AUTOMATED HEMATOLOGY ANALYZER	27.9	pg	27.0 - 34.0
MEAN CORPUSCUI	LAR HEMOGLOBIN CONC. (MCHC)	31.5 <sup>L</sup>	g/dL	32.0 - 36.0
	automated hematology analyzer BUTION WIDTH (RDW-CV)	14.5	%	11.00 - 16.00
	AUTOMATED HEMATOLOGY ANALYZER BUTION WIDTH (RDW-SD)	48.1	fL	35.0 - 56.0
	AUTOMATED HEMATOLOGY ANALYZER	40.1	IL	33.0 - 30.0
MENTZERS INDEX by CALCULATED		18.34	RATIO	BETA THALASSEMIA TRAIT: < 13.0
2, 0, 12002, 1122				IRON DEFICIENCY ANEMIA:
GREEN & KING IN	DEV	26.49	RATIO	>13.0 BETA THALASSEMIA TRAIT:<=
by CALCULATED	DEA	20.49	KATIO	65.0
				IRON DEFICIENCY ANEMIA: > 65.0
WHITE BLOOD CE	<u>ILLS (WBCS)</u>			03.0
TOTAL LEUCOCYT		4710	/cmm	4000 - 11000
	y by sf cube & microscopy BLOOD CELLS (nRBCS)	NIL		0.00 - 20.00
by AUTOMATED 6 PA	RT HEMATOLOGY ANALYZER		04	
	BLOOD CELLS (nRBCS) % AUTOMATED HEMATOLOGY ANALYZER	NIL	%	< 10 %
			0	





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Dr. Vinay Chopra Dr. Yugam Chopra MD (Pathology & Microbiology) MD (Pathology) Chairman & Consultant Pathologist **CEO & Consultant Pathologist** NAME : Mr. RAJESH BABU AGE/ GENDER : 62 YRS/MALE **PATIENT ID** :1675875 **COLLECTED BY** REG. NO./LAB NO. :012411190026 **REFERRED BY REGISTRATION DATE** : 19/Nov/2024 10:44 AM **BARCODE NO.** :01521078 **COLLECTION DATE** : 19/Nov/2024 10:47AM CLIENT CODE. : KOS DIAGNOSTIC LAB **REPORTING DATE** : 19/Nov/2024 11:07AM **CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT Test Name Value Unit **Biological Reference interval DIFFERENTIAL LEUCOCYTE COUNT (DLC)** NEUTROPHILS 44<sup>L</sup> % 50 - 70 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY LYMPHOCYTES 40 % 20 - 40 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY EOSINOPHILS 6 % 1 - 6 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY MONOCYTES 10 % 2 - 12by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY BASOPHILS 0 % 0 - 1 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY **ABSOLUTE LEUKOCYTES (WBC) COUNT** ABSOLUTE NEUTROPHIL COUNT 2072 2000 - 7500 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE LYMPHOCYTE COUNT 1884 800 - 4900 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE EOSINOPHIL COUNT 283 /cmm 40 - 440 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE MONOCYTE COUNT 471 /cmm 80 - 880 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE BASOPHIL COUNT 0 /cmm 0 - 110 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY PLATELETS AND OTHER PLATELET PREDICTIVE MARKERS. PLATELET COUNT (PLT) 150000 - 450000 138000<sup>L</sup> /cmm by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELETCRIT (PCT) 0.19 % 0.10 - 0.36 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE MEAN PLATELET VOLUME (MPV) 14<sup>H</sup> fL 6.50 - 12.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE 30000 - 90000 PLATELET LARGE CELL COUNT (P-LCC) 72000 /cmm by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET LARGE CELL RATIO (P-LCR) 52.4<sup>H</sup> % 11.0 - 45.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET DISTRIBUTION WIDTH (PDW) 15.0 - 17.0 % 17.2<sup>H</sup> by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE ADVICE **KINDLY CORRELATE CLINICALLY** 

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Test Name	Va	alue Unit	Biological Reference interval

NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD



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		gy & Microbiology) Consultant Pathologist	MD CEO & Consultant	(Pathology) : Pathologist
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ARCODE NO.	:01521078	COLL	ECTION DATE	: 19/Nov/2024 10:47AM
LIENT CODE.	: KOS DIAGNOSTIC LAB	REPO	RTING DATE	: 19/Nov/2024 11:08AM
LIENT ADDRESS	: 6349/1, NICHOLSON ROA	AD, AMBALA CANTT		
est Name		Value	Unit	Biological Reference interval
ITERPRETATION: ESR is a non-specif nmune disease, but	does not tell the health pract	esult often indicates the pro titioner exactly where the in	nflammation is in the	ion associated with infection, cancer and auto-
VIERPRETATION: . ESR is a non-specif nmune disease, but . An ESR can be affe s C-reactive protein . This test may also ystemic lupus erythe ONDITION WITH LOV . low ESR can be see bolycythaemia), sigr s sickle cells in sickl IOTE: . ESR and C - reactive . Generally, ESR doe . CRP is not affected . If the ESR is elevate	ic test because an elevated ro does not tell the health prac cted by other conditions besi be used to monitor disease a ematosus <b>W ESR</b> n with conditions that inhibit	TETRY esult often indicates the pro- titioner exactly where the in des inflammation. For this ctivity and response to the the normal sedimentation ll count (leucocytosis), and the ESR. where of inflammation. the SCRP, either at the start of s ESR, making it a better may vo types of proteins, globul	esence of inflammat filammation is in the eason, the ESR is ty rapy in both of the a of red blood cells, s some protein abno f inflammation or a: <b>rker of inflammatior</b> ins or fibrinogen.	ion associated with infection, cancer and auto e body or what is causing it. pically used in conjunction with other test such bove diseases as well as some others, such as uch as a high red blood cell count rmalities. Some changes in red cell shape (suc s it resolves. <b>n</b> .





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CLIENT ADDRESS	: 6349/1, NICHOLSON I	ROAD, AMBALA CANTT		
Test Name		Value	Unit	<b>Biological Reference interval</b>
	CI	LINICAL CHEMIS	<b>FRY/BIOCHEMIST</b>	'RY
		GLUCOSE	FASTING (F)	

IN ACCORDANCE WITH AMERICAN DIABETES ASSOCIATION 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.

test (after consumption of 75 gms of glucose) is recommended for all such patients. 3. 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.





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Test Name		Value	Unit	<b>Biological Reference interval</b>
		LIPID PROFI	LE : BASIC	
CHOLESTEROL TO	TAL: SERUM	227.32 <sup>H</sup>	mg/dL	OPTIMAL: < 200.0
by CHOLESTEROL OX		221.32 <sup></sup>	ing, all	BORDERLINE HIGH: 200.0 - 239.0 HIGH CHOLESTEROL: > OR = 240.0
TRIGLYCERIDES: S by GLYCEROL PHOSE	ERUM HATE OXIDASE (ENZYMATIC)	121.14	mg/dL	OPTIMAL: < 150.0 BORDERLINE HIGH: 150.0 - 199.0 HIGH: 200.0 - 499.0
HDL CHOLESTERO	L (DIRECT): SERUM	57.28	mg/dL	VERY HIGH: > OR = 500.0 LOW HDL: < 30.0 BORDERLINE HIGH HDL: 30.0 60.0
LDL CHOLESTEROI by CALCULATED, SPE		145.81 <sup>H</sup>	mg/dL	HIGH HDL: > OR = 60.0 OPTIMAL: < 100.0 ABOVE OPTIMAL: 100.0 - 129.0 BORDERLINE HIGH: 130.0 - 159.0 HIGH: 160.0 - 189.0 VEDV MCH = OD = 100.0
NON HDL CHOLEST by CALCULATED, SPE		170.04 <sup>H</sup>	mg/dL	VERY HIGH: > OR = 190.0 OPTIMAL: < 130.0 ABOVE OPTIMAL: 130.0 - 159.0 BORDERLINE HIGH: 160.0 - 189.0 HIGH: 190.0 - 219.0
VLDL CHOLESTER		24.23	mg/dL	VERY HIGH: > OR = 220.0 0.00 - 45.00
by CALCULATED, SPE TOTAL LIPIDS: SER by CALCULATED, SPE	CUM	575.78	mg/dL	350.00 - 700.00
by CALCULATED, SPE CHOLESTEROL/HE by CALCULATED, SPE	DL RATIO: SERUM	3.97	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: S by CALCULATED, SPE		2.55	RATIO	LOW RISK: 0.50 - 3.0 MODERATE RISK: 3.10 - 6.0 HIGH RISK: > 6.0
TRIGLYCERIDES/H by CALCULATED, SPE	IDL RATIO: SERUM	2.11 <sup>L</sup>	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.

 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.
 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
BILIRUBIN TOTAL	: SERUM	FUNCTION 0.97	I <b>TEST (COMPLETE)</b> mg/dL	INFANT: 0.20 - 8.00
	PECTROPHOTOMETRY	0.97	iiig/ uL	ADULT: 0.00 - 1.20
	C (CONJUGATED): SERUM	0.21	mg/dL	0.00 - 0.40
BILIRUBIN INDIRE	CT (UNCONJUGATED): SERUM	0.76	mg/dL	0.10 - 1.00
SGOT/AST: SERUM by IFCC, WITHOUT PY	[ /RIDOXAL PHOSPHATE	39.5	U/L	7.00 - 45.00
SGPT/ALT: SERUM by IFCC, WITHOUT PY	[ /RIDOXAL PHOSPHATE	29.4	U/L	0.00 - 49.00
AST/ALT RATIO: S by CALCULATED, SPE		1.34	RATIO	0.00 - 46.00
ALKALINE PHOSPI by PARA NITROPHEN PROPANOL	HATASE: SERUM YL PHOSPHATASE BY AMINO METHYL	105.21	U/L	40.0 - 130.0
GAMMA GLUTAMY by SZASZ, SPECTRON	L TRANSFERASE (GGT): SERUM PHTOMETRY	32.99	U/L	0.00 - 55.0
TOTAL PROTEINS: by BIURET, SPECTRO		7.14	gm/dL	6.20 - 8.00
ALBUMIN: SERUM by BROMOCRESOL G	REEN	4.24	gm/dL	3.50 - 5.50
GLOBULIN: SERUN by CALCULATED, SPE		2.9	gm/dL	2.30 - 3.50
A : G RATIO: SERUI		1.46	RATIO	1.00 - 2.00

by CALCULATED, SPECTROPHOTOMETRY

## INTERPRETATION

NOTE:- To be correlated in individuals having SGOT and SGPT values higher than Normal Referance Range.

USE:- Differential diagnosis of diseases of hepatobiliary system and pancreas.

# **INCREASED:**

DRUG HEPATOTOXICITY	> 2
ALCOHOLIC HEPATITIS	> 2 (Highly Suggestive)
CIRRHOSIS	1.4 - 2.0
INTRAHEPATIC CHOLESTATIS	> 1.5
HEPATOCELLULAR CARCINOMA & CHRONIC HEPATITIS	> 1.3 (Slightly Increased)





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# Test NameValueUnitBiological Reference interval

## DECREASED:

1. Acute Hepatitis due to virus, drugs, toxins (with AST increased 3 to 10 times upper limit of normal)

2. Extra Hepatic cholestatis: 0.8 (normal or slightly decreased).

PROGNOSTIC SIGNIFICANCE:	

NORMAL	< 0.65
GOOD PROGNOSTIC SIGN	0.3 - 0.6
POOR PROGNOSTIC SIGN	1.2 - 1.6



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50 9001 : 2008 CERTIFIED LAB		EXCELLENCE IN HEALTHCARE & DIAGNOSTICS			
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Test Name		Value	Unit	<b>Biological Reference interval</b>	
	KIDNE	EY FUNCTIO	N TEST (COMPLETE)		
UREA: SERUM		30.37	mg/dL	10.00 - 50.00	
	ATE DEHYDROGENASE (GLDH)		Ũ		
CREATININE: SERU by ENZYMATIC, SPEC		1.15	mg/dL	0.40 - 1.40	
	OGEN (BUN): SERUM	14.19	mg/dL	7.0 - 25.0	
BLOOD UREA NITE RATIO: SERUM by CALCULATED, SPE	COGEN (BUN)/CREATININE	12.34	RATIO	10.0 - 20.0	
UREA/CREATININ	E RATIO: SERUM	26.41	RATIO		
URIC ACID: SERUM		7.02	mg/dL	3.60 - 7.70	
by URICASE - OXIDAS CALCIUM: SERUM by ARSENAZO III, SPE		9.84	mg/dL	8.50 - 10.60	
PHOSPHOROUS: SE		2.71	mg/dL	2.30 - 4.70	
ELECTROLYTES					
SODIUM: SERUM by ISE (ION SELECTIV		146.3	mmol/L	135.0 - 150.0	
POTASSIUM: SERUI by ISE (ION SELECTIV	M	5.08 <sup>H</sup>	mmol/L	3.50 - 5.00	
CHLORIDE: SERUM by ISE (ION SELECTIV	1	109.73	mmol/L	90.0 - 110.0	
ESTIMATED GLOM	IERULAR FILTERATION RATE				
(eGFR): SERUM by CALCULATED INTERPRETATION:	ERULAR FILTERATION RATE een pre- and post renal azotemia.	72			

INCREASED RATIO (>20:1) WITH NORMAL CREATININE:

1. Prerenal azotemia (BUN rises without increase in creatinine) e.g. heart failure, salt depletion, dehydration, blood loss) due to decreased glomerular filtration rate.

2. Catabolic states with increased tissue breakdown.

3. GI haemorrhage.



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	٨	<b>Dr. Vinay Chopra</b> MD (Pathology & Microbiology) Chairman & Consultant Pathologist		Dr. Yugam Chopra MD (Pathology) t CEO & Consultant Pathologist				
NAME	: Mr. RAJESH H	ABU						
AGE/ GENDER	: 62 YRS/MALE		F	PATIENT ID	:1	675875		
COLLECTED BY			F	REG. NO./LAB NO.	. :	01241119002	26	
REFERRED BY				REGISTRATION D		19/Nov/2024 1		
BARCODE NO.	:01521078			COLLECTION DAT		19/Nov/2024 1		
CLIENT CODE.	: KOS DIAGNOS			REPORTING DATH	E : 1	19/Nov/2024 1	12:49PM	
CLIENT ADDRESS	: 6349/1, NICH	OLSON ROAD, AMB	ALA CANTT					
Test Name			Value	Uni	it	Biolog	ical Referen	nce interva
INCREASED RĂTIO (>2 1. Postrenal azotemia 2. Prerenal azotemia DECREASED RATIO (<	(BUN rises dispr superimposed or 0:1) WITH DECRE	red CREATININE LEVI oportionately more a renal disease.		e) (e.g. obstructive	e uropathy).			
NCREASED RATIO (>2 1. Postrenal azotemia 2. Prerenal azotemia	0:1) WITH ELEVA (BUN rises dispr superimposed or 0:1) WITH DECRE osis. Id starvation. 2. creased urea syn urea rather than monemias (urea of inappropiate ar 0:1) WITH INCREL py (accelerates c eleases muscle cl who develop ren : sis (acetoacetate creased BUN/cre apy (interferes w ULAR FILTERATION Norr Kid no	<b>FED CREATININE LEVI</b> oportionately more in renal disease. <b>ASED BUN :</b> thesis. creatinine diffuses of is virtually absent in htidiuretic harmone) <b>ASED CREATININE:</b> onversion of creating reatinine). al failure. causes false increas atinine ratio). ith creatinine measu	but of extrace blood). due to tubula e to creatinine e in creatinin rement).	Ilular fluid). ar secretion of urea	hodologies, ASSOCI No Presen	resulting in nor ATED FINDINGS proteinuria ce of Protein , or cast in urine		hen dehydra
NCREASED RATIO (>2 1. Postrenal azotemia 2. Prerenal azotemia DECREASED RATIO (< 1. Acute tubular necr 2. Low protein diet ar 3. Severe liver disease 4. Other causes of de 5. Repeated dialysis ( 6. Inherited hyperam 7. SIADH (syndrome of 8. Pregnancy. DECREASED RATIO (< 6. Phenacimide thera 2. Rhabdomyolysis (r 8. Muscular patients NAPPROPIATE RATIO 1. Diabetic ketoacido should produce an in 2. Cephalosporin ther <u>STIMATED GLOMERU</u> <u>G1</u> <u>G2</u> <u>G3a</u> <u>G3a</u> <u>G3b</u>	0:1) WITH ELEVA (BUN rises dispr superimposed or 0:1) WITH DECRE osis. Id starvation. 2. creased urea syn urea rather than monemias (urea if inappropiate ar 0:1) WITH INCRE py (accelerates c eleases muscle cl who develop ren : sis (acetoacetate creased BUN/cre apy (interferes w UAR FILTERATION Norr Kid no Mil	rED CREATININE LEVI oportionately more in renal disease. ASED BUN : thesis. creatinine diffuses of is virtually absent in ntidiuretic harmone) ASED CREATININE: onversion of creating reatinine). al failure. causes false increas atinine ratio). ith creatinine measu ith creatinine measu IRATE: DESCRIPTION nal kidney function ney damage with rmal or high GFR	but of extrace blood). due to tubula e to creatinine e in creatinin rement).	Ilular fluid). ar secretion of urea e). e with certain met <u>L/min/1.73m2 )</u> >90 >90 60 -89 30-59	hodologies, ASSOCI No Presen	ATED FINDINGS proteinuria ce of Protein ,		hen dehydra
NCREASED RATIO (>2 1. Postrenal azotemia 2. Prerenal azotemia DECREASED RATIO (< 1. Acute tubular necr 2. Low protein diet ar 3. Severe liver disease 4. Other causes of de 5. Repeated dialysis ( 6. Inherited hyperam 7. SIADH (syndrome of 8. Pregnancy. DECREASED RATIO (< 1. Phenacimide thera 2. Rhabdomyolysis (r 3. Muscular patients NAPPROPIATE RATIO 1. Diabetic ketoacido should produce an in 2. Cephalosporin ther ESTIMATED GLOMERU CKD STAGE G1 G2 G3a	0:1) WITH ELEVA (BUN rises dispr superimposed or 0:1) WITH DECRE osis. Id starvation. 2. creased urea syn urea rather than monemias (urea if inappropiate ar 0:1) WITH INCRE py (accelerates c eleases muscle cl who develop ren : sis (acetoacetate creased BUN/cre apy (interferes w UAR FILTERATION Norr Kid no Mill Mode Seve	rED CREATININE LEVI oportionately more in renal disease. ASED BUN : thesis. creatinine diffuses of is virtually absent in ntidiuretic harmone) ASED CREATININE: onversion of creatine reatinine). al failure. causes false increase atinine ratio). ith creatinine measu ith creatinine measu IRATE: DESCRIPTION nal kidney function ney damage with rmal or high GFR d decrease in GFR	but of extrace blood). due to tubula e to creatinine e in creatinin rement).	Ilular fluid). ar secretion of urea e). e with certain met <u>L/min/1.73m2 ) &gt;90 &gt;90 60 -89</u>	hodologies, ASSOCI No Presen	ATED FINDINGS proteinuria ce of Protein ,		hen dehydra





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

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









Test Name		Value Unit	Biological Reference interval
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMBA	ALA CANTT	
	: KOS DIAGNOSTIC LAB	REPORTING DATE	: 19/Nov/2024 12:49PM
CLIENT CODE.			
BARCODE NO.	: 01521078	COLLECTION DATE	: 19/Nov/2024 10:47AM
<b>REFERRED BY</b>	:	<b>REGISTRATION DATE</b>	: 19/Nov/2024 10:44 AM
COLLECTED BY	:	<b>REG. NO./LAB NO.</b>	:012411190026
AGE/ GENDER	: 62 YRS/MALE	PATIENT ID	: 1675875
NAME	: Mr. RAJESH BABU		
	Chairman & Consultan		
	Dr. Vinay Chopra MD (Pathology & Micro		m Chopra D (Pathology)

COMMENTS:

Estimated Glomerular filtration rate (eGFR) is the sum of filtration rates in all functioning nephrons and so an estimation of the GFR provides a measure of functioning nephrons of the kidney.
 eGFR calculated using the 2009 CKD-EPI creatinine equation and GFR category reported as per KDIGO guideline 2012
 In patients, with eGFR creatinine between 45-59 ml/min/1.73 m2 (G3) and without any marker of Kidney damage, It is recommended to measure of CFD with the commended to measure

3. In patients, with eGFR cleaning between 45-59 minimit 1.73 m2 (G3) and without any marker of Kidney damage, it is recommended to measure eGFR with Cystatin C for confirmation of CKD
4. eGFR category G1 OR G2 does not fulfill the criteria for CKD, in the absence of evidence of Kidney Damage
5. In a suspected case of Acute Kidney Injury (AKI), measurement of eGFR should be done after 48-96 hours of any Intervention or procedure
6. eGFR calculated by Serum Creatinine may be less accurate due to certain factors like Race, Muscle Mass, Diet, Certain Drugs. In such cases, eGFR should be calculated using Serum Cystatin C
7. A decrease in eGFR implies either progressive renal disease, or a reversible process causing decreased nephron function (eg, severe dehydration).

ADVICE:

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



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NME       : Mr. RAIRSH BABU         AGE/ GENDEE       : 02 XES/MALE       PATIENT ID       : 1075875         COLLECTED BY       ::       .:       REG. NO./LAB NO.       : 012411190028         EFFERRED BY       :       .:       REG. NO./LAB NO.       ::       19/Nov/2024 10:44 AM         BACODE NO.       ::       1521078       .:       OLLECTION DATE       ::       19/Nov/2024 10:47 AM         CUENT ODE       ::       :       SCOLLECTION DATE       ::       19/Nov/2024 10:47 AM         CUENT ODE       ::       :       SCOLLECTION DATE       ::       19/Nov/2024 10:47 AM         CUENT ODE       ::       :       SCOLLECTION DATE       ::       19/Nov/2024 10:47 AM         CUENT ODE       ::       ::       ::       NEPORTING DATE       ::       19/Nov/2024 10:47 AM         CUENT ODE       ::       ::       ::       ::       Nov:2024 10:47 AM       ::		Dr. Vinay Cho MD (Pathology & M Chairman & Consu	Microbiology)		(Pathology)
COLLECTED BY       ::       REG. NO./LAB NO.       ::       012411190026         REFERRED BY       ::       REGISTRATION DATE       ::       19/Nov/2024 10:44 AM         BARCODE NO.       ::       01521078       COLLECTION DATE       ::       19/Nov/2024 10:47AM         CLIENT CODE       ::       ::       SOS DIAGNOSTIC LAB       REPORTING DATE       ::       19/Nov/2024 11:57AM         CLIENT ADDRESS       :: <th>NAME</th> <th>: Mr. RAJESH BABU</th> <th></th> <th></th> <th></th>	NAME	: Mr. RAJESH BABU			
REFERRED BY       ::       REGISTRATION DATE       ::       19/Nov/2024 10:44 AM         BARCODE NO.       ::       01521078       COLLECTION DATE       ::       19/Nov/2024 10:47AM         CLIENT CODE.       ::       KOS DIAGNOSTIC LAB       REPORTING DATE       ::       19/Nov/2024 11:57AM         CLIENT ADDRESS       ::       :       Side Control (Control (Contro (Control (Control (Control (Control (Control (Control (Control (C	AGE/ GENDER	: 62 YRS/MALE		PATIENT ID	: 1675875
BARCODE NO.       : 01521078       COLLECTION DATE       : 19/Nov/2024 10:47AM         CLIENT CODE       : KOS DIAGNOSTIC LAB       REPORTING DATE       : 19/Nov/2024 11:57AM         CLIENT ADDRESS       : 6349/1, NICHOLSON ROAD, AMBALA CANTT       Image: College Col	<b>COLLECTED BY</b>	:		<b>REG. NO./LAB NO.</b>	: 012411190026
CLIENT CODE       KOS DIAGNOSTIC LAB       REPORTING DATE       : 19/Nov/2024 11:57AM         CLIENT ADDRESS       : 6349/1, NICHOLSON ROAD, AMBALA CANTT         Test Name       Value       Unit       Biological Reference interval         INTERVALUES         INTERVALUES         RHEUMATOID FACTOR (RA): QUANTITATIVE - SERUM         RHEUMATOID FACTOR (RA): QUANTITATIVE - SERUM         SERUM         BORDERLINE: 18.0 - 25.0         POSITIVE: > 25.0         MEEMATOID FACTOR (RA):         NEEMATOID FACTOR (RA):         NEEMATOID FACTOR (RA):         OPOSITIVE: > 25.0         POSITIVE: > 25.0 <td< th=""><th><b>REFERRED BY</b></th><th>:</th><th></th><th><b>REGISTRATION DATE</b></th><th>: 19/Nov/2024 10:44 AM</th></td<>	<b>REFERRED BY</b>	:		<b>REGISTRATION DATE</b>	: 19/Nov/2024 10:44 AM
CLIENT ADDRESS       : 6349/1, NICHOLSON ROAD, AMBALA CANTT         Test Name       Value       Unit       Biological Reference interval         International Control (RA)       Control (RA)       Control (RA)       Control (RA)       Control (RA)         REFUNATION FACTOR QUANTITATIVE:       2.74       IU/mL       NEGATIVE: < 18.0       Second (RA)         SERUM       BORDERLINE:       18.0 - 25.0       POSITIVE: > 25.0       POSITIVE: > 25.0         NTERPRETATION:       Relevance of RA is an antibodies that are directed against the Fc fragment of IgG altered in its tertiary structure.       0.0000 (RF) is diagnostically useful although it may not be etiologically related to RA.       0.0000 (RF) is diagnostically useful although it may not be etiologically related to RA.         1. The titer of RF correlately protein (CRP) are normal in about 60 % of patients with positive RA.       0.0000 (RF) is diagnostically useful although it may not disease activity, but those patients with high titers tend to have more severe disease course.         1. The titer of RF correlately protein (CRP) are normal in about 60 % of patients with positive RA.       0.0000 (RF) is diagnostically useful although it may not disease activity, but those patients with high titers tend to have more severe disease course.         1. The titer of RF correlates poorly with disease activity, but those patients with positive RA.       0.00000 (RF) is diagnostically useful and though it disease activity, but those and prognosis of rheumatoid arthritis.         1. The titer of RF correlates	BARCODE NO.	: 01521078		<b>COLLECTION DATE</b>	: 19/Nov/2024 10:47AM
Test Name         Value         Unit         Biological Reference interval           IMMUNOPATHOLOGY/SEROLOGY           BIEUMATOID FACTOR (RA): QUANTITATIVE - SERUM           RHEUMATOID FACTOR (RA): QUANTITATIVE - SERUM           SERUM         NEGATIVE: < 18.0           BORDERLINE: 18.0 - 25.0           POSITIVE: > 25.0           MEREMANDID FACTOR (RA):           I. Rheumatoid actors (RF) are antibodies that are directed against the Fc fragment of IgG altered in its tertiary structure.           2. Over 75% of patients with rheumatoid arthritis (RA) have an IgM antibody to IgG immunoglobulin. This aufoantibody (RF) is diagnostically useful although it may not be etiologically related to RA.           3. Inflammatory Markers such as ESR & C-Reactive protein (CRP) are normal in about 60 % of patients with positive RA.           A. The titre of RF correlates poorty with disease activity, but those patients with high titers tend to have more severe disease course.           5. The test is useful for diagnosis and prognosis of rheumatoid arthritis.           RHEUMATOID ARTHRITIS:           RHEUMATOID ARTHRITIS:           RHEUMATOID factor (RA):           1. Rheumatoid arthritis is a systemic autoimmune disease that is multi-functional in norigin and is characterized by chronic inflammation of the membrane lining (synovium) joints which leads to progr	CLIENT CODE.	: KOS DIAGNOSTIC LAB		REPORTING DATE	: 19/Nov/2024 11:57AM
IMMUNOPATHOLOGY/SEROLOGY         IMMUNOPATHOLOGY/SEROLOGY         RHEUMATOID FACTOR (RA): QUANTITATIVE - SERUM         RHEUMATOID FACTOR QUANTITATIVE: 2.74       IU/mL       NEGATIVE: < 18.0         SERUM       BORDERLINE: 18.0 - 25.0         by NEPHLOMETRY       DOSITIVE: > 25.0         MITERPRETATION:-         RHEUMATOID FACTOR (RA):         1. Over 75% of patients with rheumatoid arthritis (RA) have an IgM antibody to IgG immunoglobulin. This autoantibody (RF) is diagnostically useful although it may not be etiologically related to RA.         3. Inflammatory Markers such as ESR & C-Reactive protein (CRP) are normal in about 60 % of patients with positive RA.         4. The titer of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.         5. The test is useful for diagnosis and prognosis of rheumatoid arthritis.         RHEUMATOID ARTHIRITIS         1. Rheumatoid Arthritis is a systemic autoimmune disease that is multi-functional in origin and is characterized by chronic inflammation of the membrane lining (synovium) joints, with greatest damage in early phase.         2. Ne diagnosis of RA is primarily based on clinical, radiological & immunological features. The most frequent serological test is the measurement of R factor.         CAURING (RA) Expontedid arthritis, (RA) populations are not clearly	CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	MBALA CANTI	ſ	
IMMUNOPATHOLOGY/SEROLOGY         IMMUNOPATHOLOGY/SEROLOGY         RHEUMATOID FACTOR (RA): QUANTITATIVE - SERUM         RHEUMATOID FACTOR QUANTITATIVE: 2.74       IU/mL       NEGATIVE: < 18.0         SERUM       BORDERLINE: 18.0 - 25.0         by NEPHLOMETRY       DOSITIVE: > 25.0         INTERPRETATION:-         RHEUMATOID FACTOR (RA):         1. Over 75% of patients with rheumatoid arthritis (RA) have an IgM antibody to IgG immunoglobulin. This autoantibody (RF) is diagnostically useful although it may not be etiologically related to RA.         3. Inflammatory Markers such as ESR & C-Reactive protein (CRP) are normal in about 60 % of patients with positive RA.         4. The titer of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.         5. The test is useful for diagnosis and prognosis of rheumatoid arthritis.         RHEUMATOID ARTHRITIS         1. Rheumatoid Arthritis is a systemic autoimmune disease that is multi-functional in origin and is characterized by chronic inflammation of the membrane lining (synovium) joints which ledas to progressive joint destruction and in most cases to disability and reduction of quality life.         2. The disease spredas from small to large joints, with greatest damage in early phase.         3. The diagnosis of RA	Tost Namo		Valuo	Unit	Biological Poforonco interval
RHEUMATOID FACTOR (RA): QUANTITATIVE - SERUM         SERUM         by NEPHLOMETRY       2.74       IU/mL       BORDERLINE: 18.0 - 25.0         NITERPETATION:-         RHEUMATOID FACTOR (RA):         1. Rheumatoid factors (RF) are antibodies that are directed against the Fc fragment of IgG altered in its tertiary structure.         2. Over 75% of patients with rheumatoid arthritis (RA) have an IgM antibody to IgG immunogiobulin. This autoantibody (RF) is diagnostically useful although it may not be etiologically related to RA.         3. Inflammatory Markers such as ESR & C-Reactive protein (CRP) are normal in about 60 % of patients with positive RA.         4. The titer of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.         State of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.         5. The titer of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.         I. The diagnosis of RA is primarily based on clinical, radiological a tenty phase.         1. Relegnation of RA factor.         2. Non rheumatoid arthritis, sa it is often present in healthy individuals with other autoimmune diseases and chronic infection.         2. Non rheumatoid arthritis, sas it is often present in healthy individuals with other autoimmune	rest name		value	UIII	biological kelerence interval
by NEPHLOMETRY       POSITIVE: > 25.0         INTERPRETATION:- RHEUMATOID FACTOR (RA):       .         1. Rheumatoid factors (RF) are antibodies that are directed against the Fc fragment of IgG altered in its tertiary structure.       .         2. Over 75% of patients with rheumatoid arthritis (RA) have an IgM antibody to IgG immunoglobulin. This autoantibody (RF) is diagnostically useful although it may not be etiologically related to RA.       .         3. Inflammatory Markers such as ESR & C-Reactive protein (CRP) are normal in about 60 % of patients with positive RA.       .         4. The titer of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.       .         5. The test is useful for diagnosis and prognosis of rheumatoid arthritis.       RHEUMATOID ARTHIRITIS:         1. Rheumatoid Arthrititis is a systemic autoimmune disease that is multi-functional in origin and is characterized by chronic inflammation of the membrane lining (synovium) joints which ledas to progressive joint destruction and in most cases to disability and reduction of quality life.         2. The diagnosis of RA is primarily based on clinical, radiological & immunological features. The most frequent serological test is the measurement of RA factor.         CAUTION (FALSE POSITVE):       .         1. RA factor is not specific for Rheumatoid arthritis, as it is often present in healthy individuals with other autoimmune diseases and chronic infection.         2. Non rheumatoid and rheumatoid arthritis (RA) populations are not clearly separate with regard to the presence of rheumatoid f		RHEUMATOI	) FACTOR (	RA): QUANTITATIVE	- <b>SERUM</b> NEGATIVE: < 18.0
<ul> <li>INTERPRETATION:- RHEUMATOID FACTOR (RA):         <ol> <li>Rheumatoid factors (RF) are antibodies that are directed against the Fc fragment of IgG altered in its tertiary structure.</li> <li>Over 75% of patients with rheumatoid arthritis (RA) have an IgM antibody to IgG immunoglobulin. This autoantibody (RF) is diagnostically useful although it may not be etiologically related to RA.</li> <li>Inflammatory Markers such as ESR &amp; C-Reactive protein (CRP) are normal in about 60% of patients with positive RA.</li> <li>The titer of RF correlates poorly with disease activity, but those patients with high titers tend to have more severe disease course.</li> <li>The test is useful for diagnosis and prognosis of rheumatoid arthritis.</li> </ol></li></ul> <li>RHEUMATOID ARTHIRTIS:         <ul> <li>Rheumatoid Arthritis is a systemic autoimmune disease that is multi-functional in origin and is characterized by chronic inflammation of the membrane lining (synovium) joints which ledas to progressive joint destruction and in most cases to disability and reduction of quality life.</li> <li>The diagnosis of RA is primarily based on clinical, radiological &amp; immunological features. The most frequent serological test is the measurement of RA factor.</li> <li>CAUTION (FALSE POSTIVE):             <ul> <li>A factor is not specific for Rheumatoid arthritis, as it is often present in healthy individuals with other autoimmune diseases and chronic infection.</li> <li>Nor theumatoid and rheumatoid arthritis (RA) populations are not clearly separate with regard to the presence of rheumatoid factor (RF) (15% of RA patients have a nonreactive titer and 8% of nonrheumatoid patients have a positive titer).</li> <li>Patients with various nonrheumatoid diseases, characterized by chronic inflammation may have positive tests for RF. These diseases include systemic lupus erythematosus, polymyosi</li></ul></li></ul></li>					
	1. Rheumatoid factors 2. Over 75% of patient useful although it may 3. Inflammatory Mark 4. The titer of RF correct 5. The test is useful for <b>RHEUMATOID ARTHIRI</b> 1. Rheumatoid Arthiri membrane lining (syn 2. The diagnosis of RA measurement of RA fac <b>CAUTION (FALSE POST</b> 1. RA factor is not spect 2. Non rheumatoid and RA patients have a nor 3. Patients with variou: lupus erythematosus, p 4. Anti-CCP have been specific (98%) than RA 5. Upto 30 % of patient	(RF) are antibodies that are direct ts with rheumatoid arthritis (RA) r not be etiologically related to RA ers such as ESR & C-Reactive prot elates poorly with disease activity, or diagnosis and prognosis of rheu <b>TIS:</b> tis is a systemic autoimmune dise ovium) joints which ledas to prog s from small to large joints, with o to sprimarily based on clinical, ra- ctor. <b>IVE):</b> - fific for Rheumatoid arthiritis, as it d rheumatoid arthritis (RA) populat preactive titer and 8% of nonrheum s nonrheumatoid diseases, characte polymyositis, tuberculosis, syphilis, discovered in joints of patients with factor. ts with Seronegative Rheumatoid a	have an IgM ar A. ein (CRP) are n , but those pati umatoid arthrif ease that is mu greatest damag diological & im is often present tions are not cle hatoid patients erized by chroni viral hepatitis, h RA, but not in arthiritis also sh	ntibody to IgG immunoglobu ormal in about 60 % of patie ents with high titers tend to tis. Ilti-functional in origin and is estruction and in most case je in early phase. munological features. The m the healthy individuals with or parly separate with regard to have a positive titer). c inflammation may have pos infectious mononucleosis, an other form of joint disease. A now Anti-CCP antibodies.	lin. This autoantibody (RF) is diagnostically ents with positive RA. have more severe disease course. s characterized by chronic inflammation of the s to disability and reduction of quality life. host frequent serological test is the 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

KOS Diagnostic Lab (A Unit of KOS Healthcare)

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



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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



	MD (Pathology & N Chairman & Consu		st CEO & Consultant	(Pathology) Pathologist
IAME AGE/ GENDER COLLECTED BY REFERRED BY BARCODE NO. CLIENT CODE. CLIENT ADDRESS	: Mr. RAJESH BABU : 62 YRS/MALE : : : 01521078 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AN	MBALA CANT	PATIENT ID REG. NO./LAB NO. REGISTRATION DATE COLLECTION DATE REPORTING DATE	: 1675875 <b>: 012411190026</b> : 19/Nov/2024 10:44 AM : 19/Nov/2024 10:47AM : 19/Nov/2024 12:45PM
Fest Name		Value	Unit	Biological Reference interval
		¥ 7 ¥7		
			<b>FAMINS</b>	
			YDROXY VITAMIN D	
	DROXY VITAMIN D3): SERUM ESCENCE IMMUNOASSAY)	31.7	ng/mL	DEFICIENCY: < 20.0 INSUFFICIENCY: 20.0 - 30.0 SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0
ITERPRETATION:				
	CIENT: FICIENT:	< 20 21 - 29		g/mLg/mL
	D RANGE:	30 - 100		g/mL
2.25-OHVitamin D re- issue and tightly bou- Nitamin D plays a p shosphate reabsorpt .Severe deficiency m <b>BCREASED:</b> .Lack of sunshine ex .Inadequate intake, .Depressed Hepatic .Secondary to advan .Osteoporosis and S .Enzyme Inducing dr <b>NCREASED:</b> . Hypervitaminosis E evere hypercalcemia	and by a transport protein while in rimary role in the maintenance of ion, skeletal calcium deposition, ca hay lead to failure to mineralize ne posure. malabsorption (celiac disease) Vitamin D 25- hydroxylase activity need Liver disease econdary Hyperparathroidism (Mi rugs: anti-epileptic drugs like phen D is Rare, and is seen only after pro- a and hyperphophatemia. In therapy in deficient individuals	and transport in circulation. calcium home alcium mobiliz ewly formed or ld to Moderat ytoin, phenob plonged expose must be monif	form of Vitamin D and trans costatis. It promotes calciur ation, mainly regulated by p steoid in bone, resulting in r e deficiency) arbital and carbamazepine, ure to extremely high doses tored by periodic assessmer	port form of Vitamin D, being stored in adipose n absorption, renal calcium absorption and parathyroid harmone (PTH). rickets in children and osteomalacia in adults. that increases Vitamin D metabolism. of Vitamin D. When it occurs, it can result in at of Vitamin D levels in order to prevent <i>iency due to excess of melanin pigment which</i>
vpervitaminosis D				

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

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