

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



	Dr. Vinay Chopr: MD (Pathology & Micr Chairman & Consultar	robiology)		(Pathology)
NAME	: Mr. SAGAR BISHT			
AGE/ GENDER	: 31 YRS/MALE		PATIENT ID	: 1698841
COLLECTED BY	:		REG. NO./LAB NO.	:012412140014
REFERRED BY	:		REGISTRATION DATE	: 14/Dec/2024 09:59 AM
BARCODE NO.	: 01522430		COLLECTION DATE	: 14/Dec/2024 10:06AM
CLIENT CODE. CLIENT ADDRESS	: KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AMB.		REPORTING DATE	: 14/Dec/2024 10:34AM
CLIENT ADDRESS	. 0349/ I, NICHOLSON KOAD, AMD	ALA CANTI		
Test Name		Value	Unit	Biological Reference interval
RED BLOOD CELL			ELLNESS PANEL: 1.5 .00D COUNT (CBC)	5
HAEMOGLOBIN (H		15	gm/dL	12.0 - 17.0
by CALORIMETRIC			^o	
RED BLOOD CELL	(RBC) COUNT FOCUSING, ELECTRICAL IMPEDENCE	5.15 ^H	Millions/	(cmm 3.50 - 5.00
PACKED CELL VOL		47.9	%	40.0 - 54.0
MEAN CORPUSCUL	automated hematology analyzer .AR VOLUME (MCV)	93	fL	80.0 - 100.0
MEAN CORPUSCUI	AUTOMATED HEMATOLOGY ANALYZER LAR HAEMOGLOBIN (MCH)	29.1	pg	27.0 - 34.0
	AUTOMATED HEMATOLOGY ANALYZER	31.3 ^L	g/dL	32.0 - 36.0
by CALCULATED BY A	AUTOMATED HEMATOLOGY ANALYZER		_	
	BUTION WIDTH (RDW-CV) AUTOMATED HEMATOLOGY ANALYZER	14.2	%	11.00 - 16.00
	BUTION WIDTH (RDW-SD) AUTOMATED HEMATOLOGY ANALYZER	49.2	fL	35.0 - 56.0
MENTZERS INDEX		18.06	RATIO	BETA THALASSEMIA TRAIT: < 13.0
				IRON DEFICIENCY ANEMIA:
GREEN & KING INI	DFX	25.62	RATIO	>13.0 BETA THALASSEMIA TRAIT:<;
by CALCULATED		20.02	imito	65.0
				IRON DEFICIENCY ANEMIA: > 65.0
WHITE BLOOD CE	ELLS (WBCS)			00.0
TOTAL LEUCOCYT		6910	/cmm	4000 - 11000
	y by sf cube & microscopy BLOOD CELLS (nRBCS)	NIL		0.00 - 20.00
by AUTOMATED 6 PA	RT HEMATOLOGY ANALYZER		0/	~ 10.9/
	BLOOD CELLS (nRBCS) % automated hematology analyzer	NIL	%	< 10 %





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

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







Dr. Yugam Chopra

MD (Pathology & Microbiology) MD (Pathology) Chairman & Consultant Pathologist **CEO & Consultant Pathologist** NAME : Mr. SAGAR BISHT AGE/ GENDER : 31 YRS/MALE **PATIENT ID** :1698841 **COLLECTED BY** REG. NO./LAB NO. :012412140014 **REFERRED BY REGISTRATION DATE** : 14/Dec/2024 09:59 AM **BARCODE NO.** :01522430 **COLLECTION DATE** :14/Dec/2024 10:06AM CLIENT CODE. : KOS DIAGNOSTIC LAB **REPORTING DATE** :14/Dec/2024 10:34AM **CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT Test Name Value Unit **Biological Reference interval DIFFERENTIAL LEUCOCYTE COUNT (DLC)** NEUTROPHILS 49^L % 50 - 70 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY 41^H LYMPHOCYTES % 20 - 40 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY EOSINOPHILS 4 % 1 - 6 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY MONOCYTES 6 % 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 3386 2000 - 7500 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE LYMPHOCYTE COUNT 2833 800 - 4900 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE EOSINOPHIL COUNT 276/cmm 40 - 440 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE MONOCYTE COUNT 415 /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 298000 /cmm by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELETCRIT (PCT) 0.33 % 0.10 - 0.36 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE MEAN PLATELET VOLUME (MPV) fL 11 6.50 - 12.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE 30000 - 90000 PLATELET LARGE CELL COUNT (P-LCC) /cmm 101000^H by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE % PLATELET LARGE CELL RATIO (P-LCR) 33.9 11.0 - 45.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET DISTRIBUTION WIDTH (PDW) 15.0 - 17.0 16.6% by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD

Dr. Vinay Chopra

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







	Dr. Vinay Chopra MD (Pathology & Micro Chairman & Consultan	obiology) ME	m Chopra D (Pathology) ht Pathologist
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Test Name		Value Unit	Biological Reference interval



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







Dr. Vinay Cho MD (Pathology & Chairman & Cons		icrobiology)	Dr. Yugan MD CEO & Consultant	(Pathology)
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CLIENT CODE.	: KOS DIAGNOSTIC LAB	REP	ORTING DATE	: 14/Dec/2024 02:03PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AM	IBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
GLYCOSYLATED HA WHOLE BLOOD	GLYCOS EMOGLOBIN (HbA1c):	YLATED HAEM 5.4	DGLOBIN (HBA1) %	C) 4.0 - 6.4
WHOLE BLOOD by HPLC (HIGH PERFORMANCE LIQUID CHROMATOGRAPHY) ESTIMATED AVERAGE PLASMA GLUCOSE by HPLC (HIGH PERFORMANCE LIQUID CHROMATOGRAPHY)		108.28	mg/dL	60.00 - 140.00
INTERPRETATION:	AS PER AMERICAN DI			
	REFERENCE GROUP		YLATED HEMOGLOGIB	(HBAIC) in %
	abetic Adults >= 18 years	<5.7		
	t Risk (Prediabetes)	5.7 – 6.4		
Diagnosing Diabetes			>= 6.5	
D	Diagnosing Diabetes		A 40 V	
D		Goals of Th	Age > 19 Years erapy:	< 7.0
	ic goals for glycemic control	Goals of Th Actions Sugg	erapy: gested:	< 7.0 >8.0
			erapy: jested: Age < 19 Years	

KOS Diagnostic Lab

(A Unit of KOS Healthcare)

COMMENTS:

TEST PERFORMED AT KOS DIAGNOSTIC LAB. AMBALA CANTT

1.Glycosylated hemoglobin (HbA1c) test is three monthly monitoring done to assess compliace with therapeutic regimen in diabetic patients. 2.Since Hb1c reflects long term fluctuations in blood glucose concentration, a diabetic patient who has recently under good control may still have high concentration of HbAlc. Converse is true for a diabetic previously under good control but now poorly controlled.

3. Target goals of < 7.0 % may be beneficial in patients with short duration of diabetes, long life expectancy and no significant cardiovascular disease. In patients with significant complications of diabetes, limited life expectancy or extensive co-morbid conditions, targetting a goal of < 7.0% may not be appropriate.

4.High HbA1c (>9.0 -9.5 %) is strongly associated with risk of development and rapid progression of microvascular and nerve complications 5.Any condition that shorten RBC life span like acute blood loss, hemolytic anemia falsely lower HbA1c results.

6.HbA1c results from patients with HbSS,HbSC and HbD must be interpreted with caution, given the pathological processes including anemia, increased red cell turnover, and transfusion requirement that adversely impact HbA1c as a marker of long-term gycemic control.

7.Specimens from patients with polycythemia or post-splenctomy may exhibit increse in HbA1c values due to a somewhat longer life span of the red cells.



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	Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologist		Dr. Yugam Chopra MD (Pathology) st CEO & Consultant Pathologist	
AME	: Mr. SAGAR BISH	ſ		
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LIENT CODE.	: KOS DIAGNOSTIC	LAB	REPORTING DATE	: 14/Dec/2024 10:47AM
LIENT ADDRESS	: 6349/1, NICHOLS	ON ROAD, AMBALA CANTT		
est Name		Value	Unit	Biological Reference interval
by RED CELL AGGRE NTERPRETATION: . ESR is a non-specif mune disease, but . An ESR can be affe s C-reactive protein . This test may also ystemic lupus eryth ONDITION WITH LO	does not tell the hea cted by other conditi be used to monitor d ematosus W ESR	E (ESR) 25 ^H wated result often indicates th practitioner exactly wher ons besides inflammation. For sease activity and response	e the inflammation is in the or this reason, the ESR is ty to therapy in both of the a	hr 0 - 20 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
by RED CELL AGGRE NTERPRETATION: . ESR is a non-specif mmune disease, but . An ESR can be affe s C-reactive protein . This test may also ystemic lupus eryth ONDITION WITH LO . low ESR can be see bolycythaemia), sigr s sickle cells in sickl IOTE: . ESR and C - reactiv . Generally, ESR doe . CRP is not affected . If the ESR is elevat	GATION BY CAPILLARY ic test because an ele does not tell the hea cted by other conditi be used to monitor d ematosus W ESR n with conditions than ificantly high white b e cell anaemia) also e protein (C-RP) are b by as many other fac ed, it is typically a res	E (ESR) 25 ^H wated result often indicates th practitioner exactly wher ons besides inflammation. For sease activity and response t inhibit the normal sedimer lood cell count (leucocytosi	mm/1st the presence of inflammat e the inflammation is in the or this reason, the ESR is ty to therapy in both of the a natation of red blood cells, s s), and some protein abno s, and some protein abno start of inflammation or as tter marker of inflammation globulins or fibrinogen.	hr 0 - 20 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 (such s it resolves.





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	M		opra Microbiology) sultant Pathologist		(Pathology)
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CLIENT ADDRESS	: 6349/1, NICHC	OLSON ROAD,	AMBALA CANTT		
Test Name			Value	Unit	Biological Reference interval
		CLINIC	CAL CHEMIST	RY/BIOCHEMIST	'RY
			GLUCOSE	FASTING (F)	
GLUCOSE FASTING	G (F): PLASMA Se - peroxidase (go	D-POD)	104.01 ^H	mg/dL	NORMAL: < 100.0 PREDIABETIC: 100.0 - 125.0

KOS Diagnostic Lab (A Unit of KOS Healthcare)

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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DR.YUGAM CHOPRA CONSULTANT PATHOLOGIST

MBBS, MD (PATHOLOGY)

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		hopra & Microbiology) nsultant Pathologist	Dr. Yugam MD CEO & Consultant	(Pathology)
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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD	, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		LIPID PROFIL	E : BASIC	
CHOLESTEROL TO by CHOLESTEROL OX		207.62 ^H	mg/dL	OPTIMAL: < 200.0 BORDERLINE HIGH: 200.0 - 239.0
				HIGH CHOLESTEROL: > OR = 240.0
TRIGLYCERIDES: S by GLYCEROL PHOSF	ERUM PHATE OXIDASE (ENZYMATIC)	178.51 ^H	mg/dL	OPTIMAL: < 150.0 BORDERLINE HIGH: 150.0 - 199.0
				HIGH: 200.0 - 499.0 VERY HIGH: > OR = 500.0
HDL CHOLESTERO	L (DIRECT): SERUM 10N	32	mg/dL	LOW HDL: < 30.0 BORDERLINE HIGH HDL: 30.0 60.0
LDL CHOLESTEROI by CALCULATED, SPE		139.92 ^H	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
NON HDL CHOLEST by CALCULATED, SPE		175.62 ^H	mg/dL	VERY HIGH: > OR = 190.0 OPTIMAL: < 130.0 ABOVE OPTIMAL: 130.0 - 159.0 BORDERLINE HIGH: 160.0 - 189.0
		25.7	m e /dI	HIGH: 190.0 - 219.0 VERY HIGH: > OR = 220.0
VLDL CHOLESTER(by calculated, spe TOTAL LIPIDS: SER	ECTROPHOTOMETRY	35.7 593.75	mg/dL mg/dL	0.00 - 45.00 350.00 - 700.00
by CALCULATED, SPE CHOLESTEROL/HE by CALCULATED, SPE	ECTROPHOTOMETRY DL RATIO: SERUM	6.49 ^H	RATIO	LOW RISK: 3.30 - 4.40 AVERAGE RISK: 4.50 - 7.0 MODERATE RISK: 7.10 - 11.0 HIGH RISK: > 11.0

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

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Test Name		Value	Unit	Biological Reference interval
LDL/HDL RATIO: S by CALCULATED, SPE		4.37 ^H	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	5.58 ^H	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 DIRECT by DIAZO MODIFIED, S BILIRUBIN INDIRE	PECTROPHOTOMETRY Γ (CONJUGATED): SERUM SPECTROPHOTOMETRY ECT (UNCONJUGATED): SERUM	0.41 0.12 0.29	mg/dL mg/dL mg/dL	INFANT: 0.20 - 8.00 ADULT: 0.00 - 1.20 0.00 - 0.40 0.10 - 1.00
SGOT/AST: SERUM	ECTROPHOTOMETRY [/RIDOXAL PHOSPHATE	49.2 ^H	U/L	7.00 - 45.00
SGPT/ALT: SERUM		87.8 ^H	U/L	0.00 - 49.00
AST/ALT RATIO: S		0.56	RATIO	0.00 - 46.00
ALKALINE PHOSPI		76.82	U/L	40.0 - 130.0
GAMMA GLUTAMY by SZASZ, SPECTRO	L TRANSFERASE (GGT): SERUM	41.06	U/L	0.00 - 55.0
TOTAL PROTEINS: by BIURET, SPECTRO	SERUM	8.09 ^H	gm/dL	6.20 - 8.00
ALBUMIN: SERUM by BROMOCRESOL G		4.41	gm/dL	3.50 - 5.50
GLOBULIN: SERUN by CALCULATED, SPE	I ECTROPHOTOMETRY	3.68 ^H	gm/dL	2.30 - 3.50
A : G RATIO: SERUI	Μ	1.2	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)
	> 1.5 (Sirginity Increased)





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Tost Namo		Value Unit	Biological Potoronco interval

0	Test Name Value Unit Biological Reference inte
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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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: 6349/1, NICHOLSON ROAD, AM	MBALA CANTT		
	Value	Unit	Biological Reference interva
KIDNI	EY FUNCTIO	N TEST (COMPLETE))
	31.55	mg/dL	10.00 - 50.00
	1 97	ma/dI	0.40 - 1.40
ROPHOTOMETERY	1.27	liig/ uL	0.40 - 1.40
DGEN (BUN): SERUM	14.74	mg/dL	7.0 - 25.0
	11.61	RATIO	10.0 - 20.0
	24.84	RATIO	
CTROPHOTOMETRY			
PEROXIDASE	6.07	mg/dL	3.60 - 7.70
	10.25	mg/dL	8.50 - 10.60
	2 92	mg/dI.	2.30 - 4.70
ATE, SPECTROPHOTOMETRY	2.02	ing, ui	2.00 1.70
ELECTRODE)	142.7	mmol/L	135.0 - 150.0
1	3.99	mmol/L	3.50 - 5.00
ELECTRODE)	107.03	mmol/I	90.0 - 110.0
ELECTRODE)	107.03	IIIII01/L	30.0 - 110.0
ERULAR FILTERATION RATE			
RULAR FILTERATION RATE	77.5		
	MD (Pathology & N Chairman & Consu : Mr. SAGAR BISHT : 31 YRS/MALE : : : 01522430 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, A KIDNI M ROPHOTOMETERY DGEN (BUN): SERUM CTROPHOTOMETERY DGEN (BUN)/CREATININE CTROPHOTOMETRY DGEN (BUN)/CREATININE CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY RATIO: SERUM CTROPHOTOMETRY COMPACTANE COMP	: Mr. SAGAR BISHT: 31 YRS/MALE:: 01522430: KOS DIAGNOSTIC LAB: 6349/1, NICHOLSON ROAD, AMBALA CANTTKIDNEY FUNCTIOAlueKIDNEY FUNCTIOAlueKIDNEY FUNCTIOAlueKIDNEY FUNCTIOAlueKIDNEY FUNCTIOAlueKIDNEY FUNCTIOAlueCOLENTION ROAD, AMBALA CANTTKIDNEY FUNCTIOAlue <tr< td=""><td>MD (Pathology & Microbiology) Chairman & Consultant Pathologist MD (Zet & Consultant Rec No./LAB NO. ::::::::::::::::::::::::::::::::::::</td></tr<>	MD (Pathology & Microbiology) Chairman & Consultant Pathologist MD (Zet & Consultant Rec No./LAB NO. ::::::::::::::::::::::::::::::::::::

To differentiate between pre- and post renal azotemia.

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.



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

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

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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.





М		Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologist		Dr. Yugam Chopra MD (Pathology) t CEO & Consultant Pathologist					
IAME	: Mr. SAGAR B	ISHT							
GE/ GENDER	: 31 YRS/MALE]	P	PATIENT ID	:1	698841			
COLLECTED BY	:		R	REG. NO./LAB NO.	:0	1241214001	14		
REFERRED BY			R	REGISTRATION D		4/Dec/2024 0			
BARCODE NO.	: 01522430			COLLECTION DAT		4/Dec/2024 1			
CLIENT CODE.	: KOS DIAGNO	STIC I AB		REPORTING DATH		4/Dec/2024 1			
CLIENT ADDRESS		IOLSON ROAD, AME				1/ DCC/ 2024 1	1.20/101		
Test Name			Value	Uni	it	Biolog	ical Refer	ence inter	val
8. Reduced muscle m 9. Certain drugs (e.g. INCREASED RATIO (>2 1. Postrenal azotemia 2. Prerenal azotemia DECREASED RATIO (<1 1. Acute tubular necr	tetracycline, glu 0:1) WITH ELEVA (BUN rises dispr superimposed or 0:1) WITH DECRE psis.	cocorticoids) TED CREATININE LEV oportionately more n renal disease.	ELS:	e) (e.g. obstructive	e uropathy).				
 Certain drugs (e.g. NCREASED RATIO (>2 Postrenal azotemia Prerenal azotemia Prerenal azotemia CECREASED RATIO (<1 Acute tubular necr Low protein diet ar Severe liver disease Other causes of de Repeated dialysis (SIADH (syndrome c Pregnancy. PECREASED RATIO (<1 Phenacimide thera Rhabdomyolysis (rolling) Muscular patients NAPPROPIATE RATIO Diabetic ketoacido cephalosporin there STIMATED GLOMERL CKD STAGE 	tetracycline, glue 0:1) WITH ELEVA (BUN rises dispr superimposed of 0:1) WITH DECRE osis. Id starvation. a. creased urea syn urea rather than monemias (urea f inappropiate a 0:1) WITH INCRE oy (accelerates c eleases muscle c who develop rer sis (acetoacetate creased BUN/cre apy (interferes w LAR FILTERATION	cocorticoids) TED CREATININE LEV oportionately more a renal disease. CASED BUN : thesis. creatinine diffuses is virtually absent ir ntidiuretic harmone) ASED CREATININE: onversion of creatin reatinine). al failure. causes false increatin atinine ratio). vith creatinine measu I RATE: DESCRIPTION	ELS: than creatining but of extracel blood). due to tubula e to creatining se in creatining urement).	Ilular fluid). r secretion of urea e). e with certain met ./min/1.73m2)	i. hodologies,r ASSOCIA	TED FINDINGS		when dehyd	drati
Certain drugs (e.g. NCREASED RATIO (>2 Postrenal azotemia Prerenal azotemia DECREASED RATIO (<1 Acute tubular necr Low protein diet ar Severe liver disease Other causes of de Nepeated dialysis (Diherited hyperam SIADH (syndrome c SIADH (syndrome c Repaancy. DECREASED RATIO (<1 Phenacimide thera Rhabdomyolysis (n Diabetic ketoacido hould produce an in SETIMATED GLOMERL CKD STAGE	tetracycline, glui 0:1) WITH ELEVA (BUN rises dispr superimposed of 0:1) WITH DECRE osis. Id starvation. creased urea syn urea rather than monemias (urea f inappropiate a 0:1) WITH INCRE oy (accelerates c eleases muscle c who develop rer sis (acetoacetate creased BUN/cre apy (interferes w LAR FILTERATION Norn	cocorticoids) TED CREATININE LEV oportionately more a renal disease. CASED BUN : thesis. creatinine diffuses is virtually absent ir ntidiuretic harmone) ASED CREATININE: onversion of creatin reatinine). al failure. causes false increase atinine ratio). vith creatinine mease IRATE: DESCRIPTION nal kidney function	ELS: than creatining but of extracel blood). due to tubula e to creatining se in creatining urement).	Ilular fluid). r secretion of urea e). e with certain met <u>/min/1.73m2)</u> >90	hodologies,r ASSOCIA No p	TED FINDINGS roteinuria		when dehyd	drati
Certain drugs (e.g. NCREASED RATIO (>2 Postrenal azotemia Prerenal azotemia CEREASED RATIO (<1 Acute tubular necr Low protein diet ar Severe liver disease Other causes of de Nepeated dialysis (Inherited hyperam SIADH (syndrome c Pregnancy. Pregnancy. Phenacimide thera Rhabdomyolysis (r Napproplate RATIO Diabetic ketoacido hould produce an in- SETIMATED GLOMERL CKD STAGE	tetracycline, glui 0:1) WITH ELEVA (BUN rises dispr superimposed of 0:1) WITH DECRE osis. Id starvation. creased urea syn urea rather than monemias (urea f inappropiate a 0:1) WITH INCRE oy (accelerates c eleases muscle c who develop rer sis (acetoacetate creased BUN/cre apy (interferes w LAR FILTERATION Norn Kic	cocorticoids) TED CREATININE LEV oportionately more a renal disease. CASED BUN : thesis. creatinine diffuses is virtually absent ir ntidiuretic harmone) ASED CREATININE: onversion of creatin reatinine). al failure. causes false increatin atinine ratio). vith creatinine measu I RATE: DESCRIPTION	ELS: than creatining but of extracel blood). due to tubula e to creatining se in creatining urement).	Ilular fluid). r secretion of urea e). e with certain met ./min/1.73m2)	hodologies,r ASSOCIA No p Presenc	TED FINDINGS		when dehy	drati
Certain drugs (e.g. NCREASED RATIO (>2 Postrenal azotemia Prerenal azotemia DECREASED RATIO (<1 Acute tubular necr Low protein diet ar Severe liver disease Other causes of de Repeated dialysis (Inherited hyperam SIADH (syndrome c Pregnancy. DECREASED RATIO (<1 Phenacimide thera Rhabdomyolysis (r Muscular patients NAPPROPIATE RATIO Diabetic ketoacido hould produce an in Cephalosporin ther STIMATED GLOMERL G1 G2 G3a	tetracycline, glui 0:1) WITH ELEVA (BUN rises dispr superimposed of 0:1) WITH DECRE osis. Id starvation. a. creased urea syn urea rather than monemias (urea f inappropiate a 0:1) WITH INCRE oy (accelerates c eleases muscle c who develop rer sis (acetoacetate creased BUN/crea apy (interferes w LAR FILTERATION Norr Kic nc	cocorticoids) TED CREATININE LEV oportionately more a renal disease. FASED BUN : thesis. creatinine diffuses is virtually absent ir ntidiuretic harmone) ASED CREATININE: onversion of creatin reatinine). al failure. causes false increase atinine ratio). vith creatinine measu IRATE: DESCRIPTION nal kidney function Iney damage with rmal or high GFR d decrease in GFR	ELS: than creatining blood). due to tubula e to creatining se in creatining urement).	Ilular fluid). r secretion of urea e). e with certain met <u>/min/1.73m2) >90 >90 60 -89</u>	hodologies,r ASSOCIA No p Presenc	TED FINDINGS roteinuria e of Protein ,		when dehy	drati
 P. Certain drugs (e.g. NCREASED RATIO (>2 Postrenal azotemia Prerenal azotemia PCREASED RATIO (<1 Acute tubular necr Low protein diet ar Severe liver disease Other causes of de Repeated dialysis (SIADH (syndrome c Pregnancy. PCREASED RATIO (<1 Phenacimide thera Rabdomyolysis (r- Muscular patients NAPPROPIATE RATIO Diabetic ketoacido hould produce an in Cephalosporin ther STIMATED GLOMERL G1 G2 	tetracycline, glui 0:1) WITH ELEVA (BUN rises dispr superimposed of 0:1) WITH DECRE osis. Id starvation. a. creased urea syn urea rather than monemias (urea f inappropiate a 0:1) WITH INCRE py (accelerates c eleases muscle c who develop rer sis (acetoacetate creased BUN/crea apy (interferes w LAR FILTERATION Norr Kic nc Mode	cocorticoids) TED CREATININE LEV oportionately more a renal disease. FASED BUN : thesis. creatinine diffuses is virtually absent ir ntidiuretic harmone) ASED CREATININE: onversion of creatin reatinine). al failure. causes false increatin reatinine ratio). vith creatinine measu IRATE: DESCRIPTION nal kidney function liney damage with rmal or high GFR.	ELS: than creatining blood). due to tubula e to creatining se in creatining urement).	Ilular fluid). r secretion of urea e). e with certain met <u>/min/1.73m2) >90 >90</u>	hodologies,r ASSOCIA No p Presenc	TED FINDINGS roteinuria e of Protein ,		when dehyd	drati





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









	Dr. Vinay Chopra MD (Pathology & Microb Chairman & Consultant I	iology) ME	m Chopra D (Pathology) ht Pathologist
NAME	: Mr. SAGAR BISHT		
AGE/ GENDER	: 31 YRS/MALE	PATIENT ID	: 1698841
COLLECTED BY	:	REG. NO./LAB NO.	: 012412140014
REFERRED BY	:	REGISTRATION DATE	: 14/Dec/2024 09:59 AM
BARCODE NO.	: 01522430	COLLECTION DATE	: 14/Dec/2024 10:06AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPORTING DATE	: 14/Dec/2024 11:29AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMBAL	A CANTT	
Test Name	V	alue Unit	Biological Reference interval

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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		Dr. Vinay Chop MD (Pathology & Mi Chairman & Consult	crobiology)		Pathology)
NAME	: Mr. SAGAR H	BISHT			
AGE/ GENDER	: 31 YRS/MAL	E		PATIENT ID	: 1698841
COLLECTED BY	:			REG. NO./LAB NO.	:012412140014
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CLIENT ADDRESS	: 6349/1, NICI	HOLSON ROAD, AM	BALA CANTT		
Test Name			Value	Unit	Biological Reference interval
			IRON	PROFILE	
IRON: SERUM by FERROZINE, SPEC	TROPHOTOMETRY	,	83.2	µg/dL	59.0 - 158.0
UNSATURATED IR SERUM by FERROZINE, SPEC			236.5	µg/dL	150.0 - 336.0
TOTAL IRON BIND SERUM		(TIBC)	319.7	µg/dL	230 - 430
%TRANSFERRIN S.			26.02	%	15.0 - 50.0
TRANSFERRIN: SE	RUM	. ,	226.99	mg/dL	200.0 - 350.0
INTERPRETATION:-					
VARIAB		ANEMIA OF CHRO		IRON DEFICIENCY ANEMIA	
SERUM I		Normal to Re	educed	Reduced	Normal
TOTAL IRON BIND	NG CAPACITY:	Decrease	ed	Increased	Normal

IRON:

1.Serum iron studies is recommended for differential diagnosis of microcytic hypochromic anemia.i.e iron deficiency anemia, zinc deficiency anemia, anemia of chronic disease and thalassemia syndromes.

Decreased < 12-15 %

Decreased

It is essential to isolate iron deficiency anemia from Beta thalassemia syndromes because during iron replacement which is therapeutic for iron deficiency anemia, is severely contra-indicated in Thalassemia.
 TOTAL IRON BINDING CAPACITY (TIBC):

% TRANSFERRIN SATURATION:

SERUM FERRITIN:

1.It is a direct measure of protein transferrin which transports iron from the gut to storage sites in the bone marrow.

Decreased

Normal to Increased

% TRANSFERRIN SATURATION:

1. Occurs in idiopathic hemochromatosis and transfusional hemosiderosis where no unsaturated iron binding capacity is available for iron mobilization. Similar condition is seen in congenital deficiency of transferrin.



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Normal

Normal or Increased

TEST PERFORMED AT KOS DIAGNOSTIC LAB. AMBALA CANTT





	Dr. Vinay Chopr: MD (Pathology & Micr Chairman & Consultar	robiology)	M	m Chopra D (Pathology) It Pathologist
NAME	: Mr. SAGAR BISHT			
AGE/ GENDER	: 31 YRS/MALE		PATIENT ID	: 1698841
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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMB.	ALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		ENDOCI	RINOLOGY	
	THYR	DID FUNC	TION TEST: TOTAL	
TRIIODOTHYRONI	NE (T3): SERUM iescent microparticle immunoassay)	1.36	ng/mL	0.35 - 1.93
by CMIA (CHEMILUMIN			ugm /dl	107 1000
THYROXINE (T4): S	SERUM iescent microparticle immunoassay)	9.68	µgm/dl	4.87 - 12.60
THYROXINE (T4): S by CMIA (CHEMILUMIN THYROID STIMULA	iescent microparticle immunoassay) ATING HORMONE (TSH): SERUM) 1.018	μgii/di μIU/mI	
THYROXINE (T4): S by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN	iescent microparticle immunoassay) ATING HORMONE (TSH): SERUM iescent microparticle immunoassay)) 1.018		
THYROXINE (T4): S by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT	iescent microparticle immunoassay) ATING HORMONE (TSH): SERUM iescent microparticle immunoassay)) 1.018		
THYROXINE (T4): S by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT INTERPRETATION: TSH levels are subject to a day has influence on the i	IESCENT MICROPARTICLE IMMUNOASSAY) ATING HORMONE (TSH): SERUM IESCENT MICROPARTICLE IMMUNOASSAY) RASENSITIVE circadian variation, reaching peak levels betw. measured serum TSH concentrations. TSH stin) 1.018) reen 2-4 a.m and nulates the pro	µIU/mI d at a minimum between 6-10 iduction and secretion of the	0.35 - 5.50
THYROXINE (T4): S by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT INTERPRETATION: TSH levels are subject to a day has influence on the r triiodothyronine (T3).Fail overproduction(hyperthy	IESCENT MICROPARTICLE IMMUNOASSAY) ATING HORMONE (TSH): SERUM IESCENT MICROPARTICLE IMMUNOASSAY) RASENSITIVE circadian variation, reaching peak levels betwin measured serum TSH concentrations. TSH stin lure at any level of regulation of the hypotha rroidism) of T4 and/or T3.) 1.018) reen 2-4 a.m and nulates the pro	µIU/mI d at a minimum between 6-10 iduction and secretion of the y-thyroid axis will result in eith	. 0.35 - 5.50 om. The variation is of the order of 50%.Hence time of the netabolically active hormones, thyroxine (T4)and her underproduction (hypothyroidism) or
THYROXINE (T4): S by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT <u>INTERPRETATION</u> : TSH levels are subject to a day has influence on the r trilodothyronine (T3).Fail	IESCENT MICROPARTICLE IMMUNOASSAY) ATING HORMONE (TSH): SERUM IESCENT MICROPARTICLE IMMUNOASSAY) RASENSITIVE circadian variation, reaching peak levels betwin measured serum TSH concentrations. TSH stin lure at any level of regulation of the hypotha rooidism) of T4 and/or T3.) 1.018) reen 2-4 a.m and nulates the pro	µIU/mI d at a minimum between 6-10 iduction and secretion of the y-thyroid axis will result in eith	0.35 - 5.50

LIMITATIONS:-

Primary Hyperthyroidism:

Subclinical Hyperthyroidism:

1. T3 and T4 circulates in reversibly bound form with Thyroid binding globulins (TBG), and to a lesser extent albumin and Thyroid binding Pre Albumin so conditions in which TBG and protein levels alter such as pregnancy, excess estrogens, androgens, anabolic steroids and glucocorticoids may falsely affect the T3 and T4 levels and may cause false thyroid values for thyroid function tests.

Increased

Normal or High Normal

Reduced (at times undetectable)

Reduced

2. Normal levels of T4 can also be seen in Hyperthyroid patients with :T3 Thyrotoxicosis, Decreased binding capacity due to hypoproteinemia or ingestion of certain drugs (e.g.: phenytoin , salicylates).

3. Serum T4 levels in neonates and infants are higher than values in the normal adult , due to the increased concentration of TBG in neonate serum.

4. TSH may be normal in central hypothyroidism , recent rapid correction of hyperthyroidism or hypothyroidism , pregnancy , phenytoin therapy.

TRIIODOTH	YRONINE (T3)	THYROX	INE (T4)	THYROID STIMULATING HORMONE (TSF		
Age	Refferance Range (ng/mL)	Age	Refferance Range (µg/dL)	Age	Reference Range (µIU/mL)	
0 - 7 Days	0.20 - 2.65	0 - 7 Days	5.90 - 18.58	0 - 7 Days	2.43 - 24.3	
7 Days - 3 Months	0.36 - 2.59	7 Days - 3 Months	6.39 - 17.66	7 Days - 3 Months	0.58 - 11.00	
3 - 6 Months	0.51 - 2.52	3 - 6 Months	6.75 - 17.04	3 Days – 6 Months	0.70 - 8.40	
6 - 12 Months	0.74 - 2.40	6 - 12 Months	7.10 - 16.16	6 – 12 Months	0.70 - 7.00	

Increased

Normal or High Normal





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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT





		Dr. Vinay Ch MD (Pathology & Chairman & Cor		۲	am Chopra 1D (Pathology) ant Pathologist
NAME	: Mr. SAGAR	BISHT			
AGE/ GENDER	: 31 YRS/MA	LE		PATIENT ID	: 1698841
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CLIENT ADDRESS	: 6349/1, NI	CHOLSON ROAD,	AMBALA CANTT		
Test Name			Value	Unit	Biological Reference interval
1 10 Voars	0 02 2 28	1 10 Voars	6.00 13.80	1 10 Veers	0.40 E E0

1 - 10 Years	0.92 - 2.28	1 - 10 Years	6.00 - 13.80	1 – 10 Years	0.60 - 5.50	
11- 19 Years	0.35 - 1.93	11 - 19 Years	4.87-13.20	11 – 19 Years	0.50 - 5.50	
> 20 years (Adults)	0.35 - 1.93	> 20 Years (Adults)	4.87 - 12.60	> 20 Years (Adults)	0.35- 5.50	
	RECOM	MENDATIONS OF TSH L	EVELS DURING PRE	GNANCY (µIU/mL)		
	1st Trimester			0.10 - 2.50		
	2nd Trimester			0.20 - 3.00		
	3rd Trimester			0.30 - 4.10		

INCREASED TSH LEVELS:

1. Primary or untreated hypothyroidism may vary from 3 times to more than 100 times normal depending upon degree of hypofunction.

2. Hypothyroid patients receiving insufficient thyroid replacement therapy.

3.Hashimotos thyroiditis

4.DRUGS: Amphetamines, iodine containing agents & dopamine antagonist.

5.Neonatal period, increase in 1st 2-3 days of life due to post-natal surge

DECREASED TSH LEVELS:

1.Toxic multi-nodular goiter & Thyroiditis.

2. Over replacement of thyroid hormone in treatment of hypothyroidism.

3. Autonomously functioning Thyroid adenoma

4. Secondary pituitary or hypothalamic hypothyroidism

5. Acute psychiatric illness

6.Severe dehydration.

7.DRUGS: Glucocorticoids, Dopamine, Levodopa, T4 replacement therapy, Anti-thyroid drugs for thyrotoxicosis.

8.Pregnancy: 1st and 2nd Trimester





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





TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



	MD (Patho	y Chopra logy & Microbiology) & Consultant Pathologist	Dr. Yugam MD CEO & Consultant	(Pathology)
NAME	: Mr. SAGAR BISHT			
AGE/ GENDER	: 31 YRS/MALE	PATI	INT ID	: 1698841
COLLECTED BY	:	REG. I	IO./LAB NO.	:012412140014
REFERRED BY	:	REGIS	TRATION DATE	: 14/Dec/2024 09:59 AM
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CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPO	RTING DATE	: 14/Dec/2024 11:53AM
LIENT ADDRESS	: 6349/1, NICHOLSON R	OAD, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		VITAMI	VS	
		VITAMIN D/25 HYDRO	XY VITAMIN D	3
by CLIA (CHEMILUMIN	DROXY VITAMIN D3): SE rescence immunoassay)	ERUM 8.183^L	ng/mL	DEFICIENCY: < 20.0 INSUFFICIENCY: 20.0 - 30.0 SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0
<u>NTERPRETATION:</u> DFFI	CIENT:	< 20	n	g/mL
	FICIENT:	21 - 29		g/mL
PREFFER	ED RANGE:	30 - 100	ng	g/mL
1.Vitamin D compou	ICATION: nds are derived from dieta	> 100 rv ergocalciferol (from plants, nin D3 in the skin upon Ultrav	Vitamin D2), or cho	g/mL lecalciferol (from animals, Vitamin D3), or by

KOS Diagnostic Lab (A Unit of KOS Healthcare)





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

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







		chopra v & Microbiology) onsultant Pathologist	Dr. Yugan MD CEO & Consultant	(Pathology)
NAME	: Mr. SAGAR BISHT			
AGE/ GENDER	: 31 YRS/MALE	PATI	ENT ID	: 1698841
COLLECTED BY	:	REG. 1	NO./LAB NO.	: 012412140014
REFERRED BY			STRATION DATE	: 14/Dec/2024 09:59 AM
BARCODE NO.	: 01522430		ECTION DATE	: 14/Dec/2024 10:06AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		RTING DATE	: 14/Dec/2024 12:13PM
LIENT ADDRESS	: 6349/1, NICHOLSON ROA	D, MUDALA OMITT		
Test Name		Value	Unit	Biological Reference interval
<u>NTERPRETATION:-</u> Increas	SED VITAMIN B12		DECREASED VITAMI	N B12
1.Ingestion of Vitan	nin C	1.Pregnancy		
2.Ingestion of Estro			in, Anti-convulsants	, Colchicine
3.Ingestion of Vitan		3.Ethanol Igest		
4.Hepatocellular in		4. Contraceptiv		
5.Myeloproliferativ 6.Uremia	e aisorder	5.Haemodialys		
	lamin) is necessary for hemato	6. Multiple My		
2.In humans, it is ob	tained only from animal prote	ins and requires intrinsic f	actor (IF) for absorp	
	itamin B12 stores very econon	nically, reabsorbing vitami	n B12 from the ileun	n and returning it to the liver; very little is
excreted. 1 Vitamin B12 doficio	ancy may be due to lack of IE s	perotion by asstric mucos	log asstroctomy a	astric atrophy) or intestinal malabsorption (eq
	l intestinal diseases).	ecietion by gastric mucose	r (eg, gastrectority, g	
5.Vitamin B12 deficie	ency frequently causes macroo	ytic anemia, glossitis, peri	pheral neuropathy,	weakness, hyperreflexia, ataxia, loss of
proprioception, poor	coordination, and affective be	ehavioral changes. These r	nanifestations may	occur in any combination; many patients have
the neurologic defec 6 Serum methylmalo	ts without macrocytic anemia. nic acid and homocysteine lev	ols are also elevated in vit	amin B12 deficiency	states
				al cause of vitamin B12 malabsorption.
NOTE: A normal serur	m concentration of vitamin B12	2 does not rule out tissue o	leficiency of vitamin	B12. The most sensitive test for vitamin B12
				surement of MMA and homocysteine should k

NOTE: A normal serum concentration of vitamin B12 does not rule out tissue deficiency of vitamin B12. The most sensitive test for vitamin B12 deficiency at the cellular level is the assay for MMA. If clinical symptoms suggest deficiency, measurement of MMA and homocysteine should be considered, even if serum vitamin B12 concentrations are normal.





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







Dr. Vinay Cho MD (Pathology & Chairman & Cons				
NAME	: Mr. SAGAR BISHT			
AGE/ GENDER	: 31 YRS/MALE	PATI	ENT ID	: 1698841
COLLECTED BY	:		NO./LAB NO.	: 012412140014
REFERRED BY	:		STRATION DATE	: 14/Dec/2024 09:59 AM
BARCODE NO.	: 01522430		ECTION DATE	: 14/Dec/2024 10:06AM
CLIENT CODE. CLIENT ADDRESS	: KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD,		ORTING DATE	: 14/Dec/2024 11:11AM
Test Name		Value	Unit	Biological Reference interval
		CLINICAL PAT	HOLOGY	
	URINE RO	UTINE & MICROS	COPIC EXAMINA	ATION
PHYSICAL EXAMIN	ATION			
QUANTITY RECIEV	ED TANCE SPECTROPHOTOMETRY	10	ml	
COLOUR	TANCE SPECTROPHOTOMETRY	PALE YELLOW		PALE YELLOW
TRANSPARANCY		CLEAR		CLEAR
SPECIFIC GRAVITY	TANCE SPECTROPHOTOMETRY	1.02		1.002 - 1.030
CHEMICAL EXAMI				
REACTION		ACIDIC		
PROTEIN	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
SUGAR		Negative		NEGATIVE (-ve)
pH	TANCE SPECTROPHOTOMETRY	6		5.0 - 7.5
BILIRUBIN		Negative		NEGATIVE (-ve)
NITRITE	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
UROBILINOGEN	TANCE SPECTROPHOTOMETRY. TANCE SPECTROPHOTOMETRY	Normal	EU/dL	0.2 - 1.0
KETONE BODIES	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
BLOOD		Negative		NEGATIVE (-ve)
ASCORBIC ACID	TANCE SPECTROPHOTOMETRY TANCE SPECTROPHOTOMETRY MINATION	NEGATIVE (-ve)	NEGATIVE (-ve)
RED BLOOD CELLS		NEGATIVE (-ve) /HPF	0 - 3



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

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

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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



CACAD DICHT

NANCE



Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologist

Dr. Yugam Chopra MD (Pathology) CEO & Consultant Pathologist

NAME	: Mr. SAGAR BISHT			
AGE/ GENDER	: 31 YRS/MALE	Р	ATIENT ID	: 1698841
COLLECTED BY	:	R	EG. NO./LAB NO.	:012412140014
REFERRED BY	:	R	EGISTRATION DATE	: 14/Dec/2024 09:59 AM
BARCODE NO.	:01522430 CO		OLLECTION DATE	: 14/Dec/2024 10:06AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	R	EPORTING DATE	: 14/Dec/2024 11:11AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAI), AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval

PUS CELLS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	2-3	/HPF	0 - 5
EPITHELIAL CELLS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	1-2	/HPF	ABSENT
CRYSTALS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
CASTS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
BACTERIA by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
OTHERS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	NEGATIVE (-ve)		NEGATIVE (-ve)
TRICHOMONAS VAGINALIS (PROTOZOA) by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	ABSENT		ABSENT

** End Of Report ***



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