

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



	Dr. Vinay Chopra MD (Pathology & Micro Chairman & Consultan	obiology)		(Pathology)
IAME : M	r. MOHINDER SINGH			
GE/ GENDER : 64	YRS/MALE		PATIENT ID	: 1708179
COLLECTED BY :			REG. NO./LAB NO.	: 012412250001
EFERRED BY :			REGISTRATION DATE	: 25/Dec/2024 07:18 AM
	522950		COLLECTION DATE	: 25/Dec/2024 07:25AM
	DS DIAGNOSTIC LAB 849/1, NICHOLSON ROAD, AMBA	LA CANTT	REPORTING DATE	: 25/Dec/2024 11:14AM
Fest Name		Value	Unit	Biological Reference interval
	SWASTH	IYA WE	LLNESS PANEL: 1.	5
	COMP	LETE BL	OOD COUNT (CBC)	
RED BLOOD CELLS (RB	CS) COUNT AND INDICES			
HAEMOGLOBIN (HB)		13.6	gm/dL	12.0 - 17.0
RED BLOOD CELL (RBC)	COUNT ING, ELECTRICAL IMPEDENCE	5.3 ^H	Millions	/cmm 3.50 - 5.00
ACKED CELL VOLUME		43.9	%	40.0 - 54.0
AEAN CORPUSCULAR V	OLUME (MCV) MATED HEMATOLOGY ANALYZER	82.9	fL	80.0 - 100.0
MEAN CORPUSCULAR H by calculated by autom	AEMOGLOBIN (MCH) MATED HEMATOLOGY ANALYZER	25.6 ^L	pg	27.0 - 34.0
	EMOGLOBIN CONC. (MCHC)	30.9 ^L	g/dL	32.0 - 36.0
RED CELL DISTRIBUTIO	N WIDTH (RDW-CV) MATED HEMATOLOGY ANALYZER	15.5	%	11.00 - 16.00
RED CELL DISTRIBUTIO	N WIDTH (RDW-SD) MATED HEMATOLOGY ANALYZER	48.1	fL	35.0 - 56.0
MENTZERS INDEX by CALCULATED		15.64	RATIO	BETA THALASSEMIA TRAIT: < 13.0 IRON DEFICIENCY ANEMIA: >13.0
GREEN & KING INDEX by calculated WHITE BLOOD CELLS (WRCS)	24.19	RATIO	BETA THALASSEMIA TRAIT:< 65.0 IRON DEFICIENCY ANEMIA: > 65.0
FOTAL LEUCOCYTE COU	INT (TLC)	6040	/cmm	4000 - 11000
by FLOW CYTOMETRY BY S NUCLEATED RED BLOO by AUTOMATED 6 PART HEI	D CELLS (nRBCS)	NIL		0.00 - 20.00
SY AUTOWATED UPANT HEI		NIL	%	< 10 %





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Dr. Yugam Chopra

MD (Pathology)

Chairman & Consultant Pathologist **CEO & Consultant Pathologist** : Mr. MOHINDER SINGH NAME AGE/ GENDER : 64 YRS/MALE **PATIENT ID** :1708179 **COLLECTED BY** :012412250001 REG. NO./LAB NO. **REFERRED BY REGISTRATION DATE** : 25/Dec/2024 07:18 AM **BARCODE NO.** :01522950 **COLLECTION DATE** : 25/Dec/2024 07:25AM CLIENT CODE. : KOS DIAGNOSTIC LAB **REPORTING DATE** : 25/Dec/2024 11:14AM **CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT Test Name Value Unit **Biological Reference interval DIFFERENTIAL LEUCOCYTE COUNT (DLC) NEUTROPHILS** 64 % 50 - 70 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY LYMPHOCYTES 23 % 20 - 40 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY EOSINOPHILS 5 % 1 - 6 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY MONOCYTES 8 % 2 - 12by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY 0 BASOPHILS % 0 - 1 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY **IMMATURE GRANULOCTE (IG) %** 0 % 0 - 5.0 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY **ABSOLUTE LEUKOCYTES (WBC) COUNT** ABSOLUTE NEUTROPHIL COUNT 3866 2000 - 7500 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE LYMPHOCYTE COUNT 800 - 4900 1389 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE EOSINOPHIL COUNT 302 40 - 440 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE MONOCYTE COUNT 483 /cmm 80 - 880 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE BASOPHIL COUNT 0 - 110 0 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY PLATELETS AND OTHER PLATELET PREDICTIVE MARKERS. PLATELET COUNT (PLT) 180000 /cmm 150000 - 450000 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELETCRIT (PCT) 0.22 % 0.10 - 0.36 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE MEAN PLATELET VOLUME (MPV) 12^H fL 6.50 - 12.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET LARGE CELL COUNT (P-LCC) 79000 /cmm 30000 - 90000 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET LARGE CELL RATIO (P-LCR) 43.7 % 11.0 - 45.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET DISTRIBUTION WIDTH (PDW) 16.6% 15.0 - 17.0

Dr. Vinay Chopra

MD (Pathology & Microbiology)

by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE



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







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

NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD



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	Dr. Vinay Cho MD (Pathology & Chairman & Cons	Microbiology)	Dr. Yugam MD (CEO & Consultant	(Pathology)
NAME	: Mr. MOHINDER SINGH			
AGE/ GENDER	: 64 YRS/MALE	PATI	ENT ID	: 1708179
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CLIENT CODE.	: KOS DIAGNOSTIC LAB		RTING DATE	: 25/Dec/2024 02:48PM
			KIING DATE	. 23/ Dec/ 2024 02.481 M
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	MBALA CANTI		
Test Name		Value	Unit	Biological Reference interval
WHOLE BLOOD	EMOGLOBIN (HbA1c):	OSYLATED HAEMO 6.5 ^H	%	4.0 - 6.4
ESTIMATED AVERA	GE PLASMA GLUCOSE RMANCE LIQUID CHROMATOGRAPHY)	139.85	mg/dL	60.00 - 140.00
INTERPRETATION:				
		DIABETES ASSOCIATION		
	REFERENCE GROUP	GLYCOSY	LATED HEMOGLOGIB ((HBAIC) in %
	abetic Adults >= 18 years	/	<5.7	
	t Risk (Prediabetes)		5.7 – 6.4	
D	iagnosing Diabetes		>= 6.5	
			Age > 19 Years	
Thoropout	is goals for alyzomia control	Goals of The	1.3	< 7.0
inerapeut	ic goals for glycemic control	Actions Sugge		>8.0
		Goal of ther	Age < 19 Years	<7.5

KOS Diagnostic Lab

(A Unit of KOS Healthcare)

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 appropiate.

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 faisely 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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Test Name		Value	Unit	Biological Reference interval
2. An ESR can be affe as C-reactive protein		inflammation. For this r	eason, the ESR is typ	ically used in conjunction with other test such





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Test Name		Value	Unit	Biological Reference interval
	CLIN	ICAL CHEMISTRY/	BIOCHEMIST	RY
		GLUCOSE FAST	'ING (F)	

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. 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	Biological Reference interval	
			FILE : BASIC		
CHOLESTEROL TO	TAL SEDIM	96.72		OPTIMAL: < 200.0	
by CHOLESTEROL O		90.72	mg/dL	BORDERLINE HIGH: 200.0 - 239.0 HIGH CHOLESTEROL: > OR = 240.0	
TRIGLYCERIDES: S by GLYCEROL PHOSE	ERUM HATE OXIDASE (ENZYMATIC)	71.75	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 ION	41.66	mg/dL	LOW HDL: < 30.0 BORDERLINE HIGH HDL: 30.0 60.0 HIGH HDL: > OR = 60.0	
LDL CHOLESTEROI by CALCULATED, SPE		40.71	mg/dL	OPTIMAL: < 100.0 ABOVE OPTIMAL: 100.0 - 129.0 BORDERLINE HIGH: 130.0 - 159.0 HIGH: 160.0 - 189.0 VERY HIGH: > OR = 190.0	
NON HDL CHOLEST by calculated, spe		55.06	mg/dL	OPTIMAL: < 130.0 ABOVE OPTIMAL: 130.0 - 159.0 BORDERLINE HIGH: 160.0 - 189.0 HIGH: 190.0 - 219.0 VERY HIGH: > OR = 220.0	
VLDL CHOLESTER		14.35	mg/dL	0.00 - 45.00	
TOTAL LIPIDS: SER	CUM	265.19 ^L	mg/dL	350.00 - 700.00	
CHOLESTEROL/HE by CALCULATED, SPE	DL RATIO: SERUM	2.32	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		0.98	RATIO	LOW RISK: 0.50 - 3.0 MODERATE RISK: 3.10 - 6.0 HIGH RISK: > 6.0
TRIGLYCERIDES/H by CALCULATED, SPE		1.72 ^L	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
	LIVER	FUNCTION	N TEST (COMPLETE)	
BILIRUBIN TOTAL: by DIAZOTIZATION, SF	SERUM PECTROPHOTOMETRY	0.53	mg/dL	INFANT: 0.20 - 8.00 ADULT: 0.00 - 1.20
	(CONJUGATED): SERUM	0.22	mg/dL	0.00 - 0.40
BILIRUBIN INDIRE	CT (UNCONJUGATED): SERUM CTROPHOTOMETRY	0.31	mg/dL	0.10 - 1.00
SGOT/AST: SERUM by IFCC, WITHOUT PY		23.2	U/L	7.00 - 45.00
SGPT/ALT: SERUM by IFCC, WITHOUT PY		19.1	U/L	0.00 - 49.00
AST/ALT RATIO: SI by CALCULATED, SPE	ERUM	1.21	RATIO	0.00 - 46.00
ALKALINE PHOSPH by Para Nitropheny Propanol	IATASE: SERUM YL PHOSPHATASE BY AMINO METHYL	75.28	U/L	40.0 - 130.0
GAMMA GLUTAMY by SZASZ, SPECTROF	L TRANSFERASE (GGT): SERUM	17.31	U/L	0.00 - 55.0
TOTAL PROTEINS: by BIURET, SPECTRON		7.22	gm/dL	6.20 - 8.00
ALBUMIN: SERUM by BROMOCRESOL GI	REEN	4.83	gm/dL	3.50 - 5.50
GLOBULIN: SERUM	[2.39	gm/dL	2.30 - 3.50
A : G RATIO: SERUN	Л	2.02 ^H	RATIO	1.00 - 2.00

by CALCULATED, SPECTROPHOTOMETRY

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:

> 2
> 2 (Highly Suggestive)
1.4 - 2.0
> 1.5
> 1.3 (Slightly Increased)



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INTERPRETATION





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NAME	: Mr. MOHINDER SINGH			
	MD (Pathology 8	& Microbiology)		(Pathology)
	Dr. Vinay Cł	nopra	Dr. Yugan	n Chopra

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).

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



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	KIDNI	EY FUNCTION T	TEST (COMPLETE)	
UREA: SERUM	IATE DEHYDROGENASE (GLDH)	40.09	mg/dL	10.00 - 50.00
CREATININE: SER	UM	1.25	mg/dL	0.40 - 1.40
BLOOD UREA NITH	BLOOD UREA NITROGEN (BUN): SERUM by CALCULATED, SPECTROPHOTOMETRY		mg/dL	7.0 - 25.0
	ROGEN (BUN)/CREATININE	14.98	RATIO	10.0 - 20.0
by CALCULATED, SPE	ECTROPHOTOMETRY			
UREA/CREATININ	E RATIO: SERUM	32.07	RATIO	
URIC ACID: SERUM	1	4.69	mg/dL	3.60 - 7.70
CALCIUM: SERUM by ARSENAZO III, SPE		9	mg/dL	8.50 - 10.60
PHOSPHOROUS: SI		3.8	mg/dL	2.30 - 4.70
ELECTROLYTES				
SODIUM: SERUM		139.2	mmol/L	135.0 - 150.0
by ISE (ION SELECTIV POTASSIUM: SERU by ISE (ION SELECTIV	M	4.1	mmol/L	3.50 - 5.00
CHLORIDE: SERUN by ISE (ION SELECTIV	1	104.4	mmol/L	90.0 - 110.0
	IERULAR FILTERATION RATE			
ESTIMATED GLOM (eGFR): SERUM by CALCULATED INTERPRETATION:	IERULAR FILTERATION RATE	64.3		

INTERPRETATION:

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.



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



Page 11 of 20

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.





Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologist		Microbiology)	Dr. Yugam Chopra MD (Pathology) st CEO & Consultant Pathologist		
NAME	: Mr. MOHINDER SINGH				
AGE/ GENDER	: 64 YRS/MALE	PATIENT ID	: 1708179		
COLLECTED BY	:	REG. NO./LAB NO	. : 01241225(0001	
REFERRED BY		REGISTRATION I			
BARCODE NO.	: 01522950	COLLECTION DAT			
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPORTING DAT	E : 25/Dec/202	4 11:14AM	
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	AMBALA CANTT			
Test Name		Value U	nit Biol	ogical Reference interval	
8. Reduced muscle m 9. Certain drugs (e.g. INCREASED RATIO (>2 1. Postrenal azotemia 2. Prerenal azotemia DECREASED RATIO (<	superimposed on renal disease. 10:1) WITH DECREASED BUN :		e uropathy).		
8. Reduced muscle m 9. Certain drugs (e.g. INCREASED RATIO (>2 1. Postrenal azotemia 2. Prerenal azotemia DECREASED RATIO (< 1. Acute tubular necr 2. Low protein diet al 3. Severe liver diseas 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 INAPPROPIATE RATIO 1. Diabetic ketoacido should produce an in 2. Cephalosporin thei ESTIMATED GLOMERI OKD STAGE	ass (subnormal creatinine productetracycline, glucocorticoids) 20:1) WITH ELEVATED CREATININE a (BUN rises disproportionately m superimposed on renal disease. 10:1) WITH DECREASED BUN : osis. ad starvation. e. creased urea synthesis. (urea rather than creatinine diffu- monemias (urea is virtually absen- of inappropiate antidiuretic harmon 10:1) WITH INCREASED CREATININ py (accelerates conversion of cre- eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false inco- creased BUN/creatinine ratio). rapy (interferes with creatinine m JLAR FILTERATION RATE: DESCRIPTION	LEVELS: ore than creatinine) (e.g. obstructive ses out of extracellular fluid). nt in blood). one) due to tubular secretion of ure E: atine to creatinine). crease in creatinine with certain me easurement).	a. thodologies,resulting in ASSOCIATED FINDIN		
A. Reduced muscle m Certain drugs (e.g. NCREASED RATIO (>2 Postrenal azotemia Prerenal azotemia DECREASED RATIO (< Acute tubular necr Low protein diet al Severe liver diseas Other causes of de Repeated dialysis Inherited hyperam SIADH (syndrome of Pregnancy. DECREASED RATIO (< Nuscular patients NAPPROPIATE RATIO Diabetic ketoacido hould produce an in Cephalosporin ther STIMATED GLOMERI CKD STAGE G1	ass (subnormal creatinine productetracycline, glucocorticoids) 20:1) WITH ELEVATED CREATININE a (BUN rises disproportionately m superimposed on renal disease. 10:1) WITH DECREASED BUN : osis. ad starvation. e. creased urea synthesis. (urea rather than creatinine diffu- monemias (urea is virtually absen- of inappropiate antidiuretic harmon 10:1) WITH INCREASED CREATININ py (accelerates conversion of cre- eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false inco- creased BUN/creatinine ratio). rapy (interferes with creatinine m JLAR FILTERATION RATE: 	LEVELS: ore than creatinine) (e.g. obstructive ses out of extracellular fluid). nt in blood). one) due to tubular secretion of ure E: atine to creatinine). crease in creatinine with certain me easurement). GFR (mL/min/1.73m2) ion	a. thodologies,resulting in ASSOCIATED FINDIN No proteinuria	IGS	
A Reduced muscle m Certain drugs (e.g. NCREASED RATIO (>2 Prerenal azotemia Prerenal azotemia DECREASED RATIO (< Acute tubular necr Low protein diet al Severe liver diseas Other causes of de Repeated dialysis Inherited hyperam SIADH (syndrome of Pregnancy. DECREASED RATIO (< Nuscular patients Nuscular patients Muscular patients Muscular patients Mappropiate RATIO Diabetic ketoacido should produce an in CEphalosporin their ESTIMATED GLOMERI OKD STAGE	ass (subnormal creatinine productetracycline, glucocorticoids) 20:1) WITH ELEVATED CREATININE a (BUN rises disproportionately m superimposed on renal disease. 10:1) WITH DECREASED BUN : osis. ad starvation. e. creased urea synthesis. (urea rather than creatinine diffu- monemias (urea is virtually absen- of inappropiate antidiuretic harmon 10:1) WITH INCREASED CREATININ py (accelerates conversion of cre- eleases muscle creatinine). who develop renal failure. : sis (acetoacetate causes false inco- creased BUN/creatinine ratio). rapy (interferes with creatinine m JLAR FILTERATION RATE: DESCRIPTION Normal kidney funct Kidney damage wit	LEVELS: ore than creatinine) (e.g. obstructive ses out of extracellular fluid). nt in blood). one) due to tubular secretion of ure E: atine to creatinine). crease in creatinine with certain me easurement).	a. thodologies,resulting in ASSOCIATED FINDIN No proteinuria Presence of Proteir	IGS	
A. Reduced muscle m Certain drugs (e.g. NCREASED RATIO (>2 Prerenal azotemia Prerenal azotemia DECREASED RATIO (< Acute tubular necr Low protein diet al Severe liver diseas Other causes of de Repeated dialysis Inherited hyperam SIADH (syndrome of Pregnancy. DECREASED RATIO (< Nuscular patients NAPPROPIATE RATIO Diabetic ketoacido hould produce an in Cephalosporin ther STIMATED GLOMERI CKD STAGE G1 G2	ass (subnormal creatinine producterracycline, glucocorticoids) 20:1) WITH ELEVATED CREATININE a (BUN rises disproportionately m superimposed on renal disease. 10:1) WITH DECREASED BUN : osis. ad starvation. e. creased urea synthesis. (urea rather than creatinine diffu- monemias (urea is virtually absen- of inappropiate antidiuretic harmon 10:1) WITH INCREASED CREATININ py (accelerates conversion of cre- eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false inco- creased BUN/creatinine ratio). rapy (interferes with creatinine m JLAR FILTERATION RATE: DESCRIPTION Normal kidney funct Kidney damage wit normal or high GFI	LEVELS: ore than creatinine) (e.g. obstructive ses out of extracellular fluid). nt in blood). one) due to tubular secretion of ure E: atine to creatinine). crease in creatinine with certain me easurement). Image: the secret of the secret o	a. thodologies,resulting in ASSOCIATED FINDIN No proteinuria	IGS	
B. Reduced muscle m Certain drugs (e.g. INCREASED RATIO (>2 I. Postrenal azotemia DECREASED RATIO (< I. Acute tubular necr Low protein diet an Severe liver diseas Other causes of de Severe liver diseas Other causes of de Severe liver diseas Other causes of de Repeated dialysis Inherited hyperam SIADH (syndrome of SIADH (syndrome of Severe liver diseas Regnancy. DECREASED RATIO (< I. Phenacimide thera Rhabdomyolysis (r S. Muscular patients INAPPROPIATE RATIO Loiabetic ketoacido should produce an in CEphalosporin there STIMATED GLOMERI G1 G2 G3a	ass (subnormal creatinine producterracycline, glucocorticoids) 20:1) WITH ELEVATED CREATININE a (BUN rises disproportionately m superimposed on renal disease. 10:1) WITH DECREASED BUN : osis. ad starvation. e. creased urea synthesis. (urea rather than creatinine diffu- monemias (urea is virtually abser- of inappropiate antidiuretic harmon 10:1) WITH INCREASED CREATININ py (accelerates conversion of cre- eleases muscle creatinine). who develop renal failure. : sis (acetoacetate causes false inco- creased BUN/creatinine ratio). rapy (interferes with creatinine m JLAR FILTERATION RATE: DESCRIPTION Normal kidney funct- Kidney damage wit- normal or high GFf Mild decrease in GF	LEVELS: ore than creatinine) (e.g. obstructive ses out of extracellular fluid). nt in blood). one) due to tubular secretion of ure E: atine to creatinine). crease in creatinine with certain me easurement). ion >90 h >90 R 60 -89	a. thodologies,resulting in ASSOCIATED FINDIN No proteinuria Presence of Proteir	IGS	
B. Reduced muscle m Certain drugs (e.g. INCREASED RATIO (>2 I. Postrenal azotemia DECREASED RATIO (< 1. Acute tubular necr Low protein diet al Severe liver diseas Other causes of de S. Repeated dialysis Inherited hyperam SIADH (syndrome of SIADH (syndrome of Regnancy. DECREASED RATIO (< I. Phenacimide thera Rhabdomyolysis (r S. Muscular patients INAPPROPIATE RATIO Loiabetic ketoacido should produce an in CED STAGE G1 G2	ass (subnormal creatinine producterracycline, glucocorticoids) 20:1) WITH ELEVATED CREATININE a (BUN rises disproportionately m superimposed on renal disease. 10:1) WITH DECREASED BUN : osis. ad starvation. e. creased urea synthesis. (urea rather than creatinine diffu- monemias (urea is virtually absen- of inappropiate antidiuretic harmon 10:1) WITH INCREASED CREATININ py (accelerates conversion of cre- eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false inco- creased BUN/creatinine ratio). rapy (interferes with creatinine m JLAR FILTERATION RATE: DESCRIPTION Normal kidney funct Kidney damage wit normal or high GFI	LEVELS: ore than creatinine) (e.g. obstructive ses out of extracellular fluid). nt in blood). one) due to tubular secretion of ure E: atine to creatinine). crease in creatinine with certain me easurement). ion >90 h >90 R 60 -89 GFR 30-59	a. thodologies,resulting in ASSOCIATED FINDIN No proteinuria Presence of Proteir	IGS	





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

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









: KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AMBAL	REPORTING DATE A CANTT	: 25/Dec/2024 11:14AM
		: 25/Dec/2024 11:14AM
: KOS DIAGNOSTIC LAB	REPORTING DATE	: 25/Dec/2024 11:14AM
: 01522950	COLLECTION DATE	: 25/Dec/2024 07:25AM
:	REGISTRATION DATE	: 25/Dec/2024 07:18 AM
:	REG. NO./LAB NO.	: 012412250001
: 64 YRS/MALE	PATIENT ID	: 1708179
: Mr. MOHINDER SINGH		
		0 (Pathology) t Pathologist
Dr. Vinay Chopra		n Chopra
	MD (Pathology & Microt Chairman & Consultant : Mr. MOHINDER SINGH : 64 YRS/MALE :	MD (Pathology & Microbiology) Chairman & Consultant Pathologist CEO & Consultant : Mr. MOHINDER SINGH : 64 YRS/MALE PATIENT ID : REG. NO./LAB NO. : REGISTRATION DATE

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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	1	D r. Vinay Chopi MD (Pathology & Mic Chairman & Consulta	robiology)		(Pathology)
NAME	: Mr. MOHINI	DER SINGH			
AGE/ GENDER	: 64 YRS/MALI	3		PATIENT ID	: 1708179
COLLECTED BY	:			REG. NO./LAB NO.	: 012412250001
REFERRED BY	:			REGISTRATION DATE	: 25/Dec/2024 07:18 AM
BARCODE NO.	:01522950			COLLECTION DATE	: 25/Dec/2024 07:25AM
CLIENT CODE.	: KOS DIAGNO	STIC LAB		REPORTING DATE	: 25/Dec/2024 12:13PM
CLIENT ADDRESS : 6349/1, NICHOLSON ROAD, AMBALA CANTT					
Test Name			Value	Unit	Biological Reference interva
			IRON	PROFILE	
IRON: SERUM	TROPHOTOMETRY		57.81 ^L	µg/dL	59.0 - 158.0
UNSATURATED IRON BINDING CAPACITY (UIBC) SERUM by FERROZINE, SPECTROPHOTOMETERY		241.69	µg/dL	150.0 - 336.0	
		299.5	µg/dL	230 - 430	
		19.3	%	15.0 - 50.0	
TRANSFERRIN: SEI by SPECTROPHOTOM	RUM		212.65	mg/dL	200.0 - 350.0
INTERPRETATION:-					
VARIAB SERUM IF		ANEMIA OF CHRON Normal to Re		IRON DEFICIENCY ANEMIA Reduced	A THALASSEMIA α/β TRAIT Normal

TOTAL IRON BINDING CAPACITY: Normal Decreased Increased % TRANSFERRIN SATURATION: Decreased Decreased < 12-15 % Normal **SERUM FERRITIN:** Normal to Increased Decreased Normal or Increased

IRON:

TEST PERFORMED AT KOS DIAGNOSTIC LAB. AMBALA CANTT

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. 2. 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):**

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

% 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.





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







	Dr. Vinay Ch MD (Pathology & Chairman & Con		Dr. Yugam Ch MD (Path CEO & Consultant Path	ology)	
NAME	: Mr. MOHINDER SINGH				
AGE/ GENDER	: 64 YRS/MALE	PATIE	NT ID : 1	708179	
COLLECTED BY	:	REG. N	0./LAB NO. : (12412250001	
REFERRED BY	:	REGIST	FRATION DATE : 2	5/Dec/2024 07:18 AM	
BARCODE NO.	: 01522950	COLLE	CTION DATE : 2	5/Dec/2024 07:25AM	
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPOF	TING DATE : 2	5/Dec/2024 11:39AM	
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANTT			
Test Name		Value	Unit	Biological Reference inter	rval
		ENDOCRINO	LOGY		
	ТН	YROID FUNCTION	TEST: TOTAL		
TRIIODOTHYRONI	NE (T3): SERUM	0.916 ssay)	ng/mL	0.35 - 1.93	
		8.04 SSAY)	µgm/dL	4.87 - 12.60	
by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN	ATING HORMONE (TSH): SERU		µIU/mL	0.35 - 5.50	
by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ATING HORMONE (TSH): SERU		µIU/mL	0.35 - 5.50	
by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT <u>INTERPRETATION</u> : TSH levels are subject to day has influence on the trilodothyronine (T3).Fai	ATING HORMONE (TSH): SERU VESCENT MICROPARTICLE IMMUNOAS RASENSITIVE	SSAY) 5 between 2-4 a.m and at a mir 5H stimulates the production a	nimum between 6-10 pm. The and secretion of the metabo	<i>variation is of the order of 50%.Hence tim</i> ically active hormones, thyroxine (T4)and	
by CMIA (CHEMILUMIN THYROID STIMULA by CMIA (CHEMILUMIN 3rd GENERATION, ULT <u>INTERPRETATION:</u> TSH levels are subject to day has influence on the triiodothyronine (T3).Fai	ATING HORMONE (TSH): SERU VESCENT MICROPARTICLE IMMUNOAS RASENSITIVE circadian variation, reaching peak levels measured serum TSH concentrations. TS ilure at any level of regulation of the hy	SSAY) 5 between 2-4 a.m and at a mir 5H stimulates the production a	nimum between 6-10 pm. The and secretion of the metabo	<i>variation is of the order of 50%.Hence tim</i> ically active hormones, thyroxine (T4)and	

CLINICAL CONDITION	T3	T4	TSH
Primary Hypothyroidism:	Reduced	Reduced	Increased (Significantly)
Subclinical Hypothyroidism:	Normal or Low Normal	Normal or Low Normal	High
Primary Hyperthyroidism:	Increased	Increased	Reduced (at times undetectable)
Subclinical Hyperthyroidism:	Normal or High Normal	Normal or High Normal	Reduced

LIMITATIONS:-

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.

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)	THYROXINE (T4)		THYROID STIMULATING HORMONE (TSH)		
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	





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	Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologis		(Pathology)
NAME	: Mr. MOHINDER SINGH		
AGE/ GENDER	: 64 YRS/MALE	PATIENT ID	: 1708179
COLLECTED BY	:	REG. NO./LAB NO.	: 012412250001
REFERRED BY	:	REGISTRATION DATE	: 25/Dec/2024 07:18 AM
BARCODE NO.	: 01522950	COLLECTION DATE	: 25/Dec/2024 07:25AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPORTING DATE	: 25/Dec/2024 11:39AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMBALA CANTT		
Test Name	Value	Unit	Biological Reference interval

Test Name			Value	Unit	t	Biological Reference interva
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	IMENDATIONS OF TSH LI	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) DR.YUGAM CHOPRA CONSULTANT PATHOLOGIST MBBS , MD (PATHOLOGY)





TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



NAME : Mr. MOHINDER SINGH AGE/ GENDER : 64 YRS/MALE PATIENT ID : 1708179 COLLECTED BY :: REG.NO./LAB NO. : 012412250001 REFERED BY :: REGISTRATION DATE : 25/Dec/2024 07:15A M BARCODE NO. : 01522950 COLLECTION DATE : 25/Dec/2024 07:25A M CLIENT CODE : KOS DIAGNOSTIC LAB REPORTING DATE : 25/Dec/2024 07:25A M CLIENT ADDRESS : 6349/1, NICHOLSON ROAD, AMBALA CANTT Estop : 25/Dec/2024 07:25A M VITAMIND D/25 HYDROXY VITAMIN D/25 HYDROXY VITAMIN D3 USTAMIND //25 HYDROXY VITAMIN D/25 HYDROXY VITAMIN D3 INSUFFICIENCY: < 20.0 VITAMIN D (25-HYDROXY VITAMIND 3): SERUM 22.9 ^L ng/mL DEFICIENCY: < 20.0 by CLIA (CHEMILIAMNESCENCE MAMINOASSAY) Z2.9 ^L ng/mL INSUFFICIENCY: < 20.0 NUTERPRETATION: - :20 ng/mL INSUFFICIENCY: < 20.0 NUTERPRETATION: - :20 ng/mL INSUFFICIENCY: < 20.0 INSUFFICIENT: - :20 ng/mL INSUFFICIENCY: < 20.0 INSUFFICIENT: - :20 ng/mL INSUFFICIENCY: < 20.0 INSUFFICIENT			Chopra ogy & Microbiology) Consultant Pathologist		(Pathology)
VITAMINS VITAMIN D/25 HYDROXY VITAMIN D3 by CLIA (CHEMILUMINESCENCE IMMUNOASSAY) DEFICIENCY: < 20.0	AGE/ GENDER COLLECTED BY REFERRED BY BARCODE NO. CLIENT CODE.	: 64 YRS/MALE : : : 01522950 : KOS DIAGNOSTIC LAB		REG. NO./LAB NO. REGISTRATION DATE COLLECTION DATE	: 012412250001 : 25/Dec/2024 07:18 AM : 25/Dec/2024 07:25AM
UTAMIN D/25 HYDROXY VITAMIN D3): SERUM by CLIA (CHEMILLUMINESCENCE IMMUNOASSAY) 22.9 ^L ng/mL DEFICIENCY: < 20.0 DEFICIENCY: 20.0 - 30.0 SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0	Test Name		Value	Unit	Biological Reference interval
SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0 INTERPRETATION: DEFICIENT: 21 - 29 ng/mL INSUFFICIENT: 21 - 29 ng/mL INTOXICATION: > 100 ng/mL NOT Colspan="2">Not Market in the start of the st		DROXY VITAMIN D3): SEI	ITAMIN D/25 HY	(DROXY VITAMIN D	DEFICIENCY: < 20.0
PREFFERED RANGE: 30 - 100 ng/mL INTOXICATION: > 100 ng/mL .Vitamin D compounds are derived from dietary eraocalciferol (from plants. Vitamin D2), or cholecalciferol (from animals, Vitamin D3), or by conversion of 7- dihydrocholecalciferol to Vitamin D3 in the skin upon Ultraviolet exposure. .25-OHVitamin D represents the main body resevoir and transport form of Vitamin D and transport form of Vitamin D, being stored in adipolisue and tightly bound by a transport protein while in circulation. .Vitamin D plays a primary role in the maintenance of calcium momeostatis. It promotes calcium absorption, renal calcium absorption and phosobate reabsorption, skeletal calcium deposition, calcium mobilization. mainly regulated by parathyroid harmone (PTH). .Severe deficiency may lead to failure to mineralize newly formed osteoid in bone, resulting in rickets in children and osteomalacia in adults SECREASED: .Lack of sunshine exposure. .Lack of sunshine is exposure. .Lack of sunshine exposure. .Lack of sunshine exposure. .Lack of sunshine exposure. .Lack of sunshine exposure.	DEFIC				SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0 g/mL
INTOXICATION: > 100 ng/mL 1. Vitamin D compounds are derived from dietary ergocalciferol (from plants, Vitamin D2), or cholecalciferol (from animals, Vitamin D3), or by conversion of 7- dihydrocholecalciferol to Vitamin D3 in the skin upon Ultraviolet exposure. 2.25-OHVitamin D represents the main body resevoir and transport form of Vitamin D and transport form of Vitamin D, being stored in adipot issue and tightly bound by a transport protein while in circulation. 3. Vitamin D plays a primary role in the maintenance of calcium homeostatis. It promotes calcium absorption, renal calcium absorption and obsophate reabsorption, skeletal calcium deposition, calcium mobilization, mainly regulated by parathyroid harmone (PTH). 4. Severe deficiency may lead to failure to mineralize newly formed osteoid in bone, resulting in rickets in children and osteomalacia in adults DCREASED: 1. Lack of sunshine exposure. 2. Inadequate intake, malabsorption (celiac disease) 3. Depressed Hepatic Vitamin D 25- hydroxylase activity 4. Secondary to advanced Liver disease 5. Osteoporosis and Secondary Hyperparathroidism (Mild to Moderate deficiency) 5. Enzyme Inducing drugs: anti-epileptic drugs like phenytoin, phenobarbital and carbamazepine, that increases Vitamin D metabolism. NCREASED: 1. Hypervitaminosis D is Rare, and is seen only after prolonged exposure to extremely high doses of Vitamin D. When it occurs, it can result in severe hypercalcemia and hyperphophatemia. AUTION: Replacement therapy in deficient individuals must be monito					5
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	conversion of 7- dihyc 2.25-OHVitamin D re- tissue and tiahtly bou 3.Vitamin D plays a pr phosphate reabsorpti 4.Severe deficiency m DECREASED: 1.Lack of sunshine ext 2.Inadequate intake, 1 3.Depressed Hepatic V 4.Secondary to advant 5.Osteoporosis and Se 6.Enzyme Inducing dru INCREASED: 1. Hypervitaminosis D severe hypercalcemia CAUTION: Replacemer hypervitaminosis D NOTE: -Dark coloured ii	Irocholecalciferol to Vitam presents the main body re- nd by a transport protein w imary role in the maintena on, skeletal calcium deposi ay lead to failure to minera bosure. malabsorption (celiac disea /itamin D 25- hydroxylase a condary Hyperparathroidi: ugs: anti-epileptic drugs lik is Rare, and is seen only af and hyperphophatemia. ht therapy in deficient indiv <i>ndividuals as compare to wh</i>	in D3 in the skin upon sevoir and transport for vhile in circulation. ince of calcium homeo tion, calcium mobiliza alize newly formed ost ase) activity sm (Mild to Moderate e phenytoin, phenobar ter prolonged exposur riduals must be monito	Ultraviolet exposure. form of Vitamin D and trans ostatis. It promotes calciur tion, mainly regulated by r eoid in bone, resulting in r deficiency) rbital and carbamazepine, re to extremely high doses pred by periodic assessmer	port form of Vitamin D, being stored in adipose n absorption, renal calcium absorption and barathyroid 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

KOS Diagnostic Lab (A Unit of KOS Healthcare)





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NAME : Mr. MC AGE/ GENDER : 64 YRS/ COLLECTED BY : REFERRED BY :	DHINDER SINGH /MALE	PATIE			
COLLECTED BY :	/MALE	PATIE			
			NT ID :	1708179	
REFERRED BY :		REG. N	O./LAB NO. :	012412250001	
		REGIS	RATION DATE :	25/Dec/2024 07:18 AM	
BARCODE NO. : 015229	950	COLLE	CTION DATE :	25/Dec/2024 07:25AM	
	AGNOSTIC LAB			25/Dec/2024 11:39AM	
	I, NICHOLSON ROAD, AME				
	r, menolson kond, nim	men on with			
Test Name		Value	Unit	Biological Reference interva	
by CMIA (CHEMILUMINESCENT MIC INTERPRETATION:-	CROPARTICLE IMMUNOASSAY	315.3	pg/mL	190.0 - 830	
INCREASED VITAMI	IN B12	C	ECREASED VITAMIN B1	2	
1.Ingestion of Vitamin C		1.Pregnancy			
	2.DRUGS:Aspirin, Anti-convulsants, Colchicine				
2.Ingestion of Estrogen				ICHICITE	
3.Ingestion of Vitamin A		3.Ethanol Igestic	on		
3.Ingestion of Vitamin A 4.Hepatocellular injury		3.Ethanol Igestic 4. Contraceptive	harmones		
3.Ingestion of Vitamin A		3.Ethanol Igestic	n Harmones S		

5. Vitamin B12 deficiency frequently causes macrocytic anemia, glossitis, peripheral neuropathy, weakness, hyperreflexia, ataxia, loss of proprioception, poor coordination, and affective behavioral changes. These manifestations may occur in any combination; many patients have the neurologic defects without macrocytic anemia.

6.Serum methylmalonic acid and homocysteine levels are also elevated in vitamin B12 deficiency states.

7.Follow-up testing for antibodies to intrinsic factor (IF) is recommended to identify this potential cause of vitamin B12 malabsorption. **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.





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	Dr. Vinay Cho MD (Pathology & N Chairman & Consu	licrobiology)	Dr. Yugam MD O & Consultant	(Pathology)
NAME	: Mr. MOHINDER SINGH			
AGE/ GENDER	: 64 YRS/MALE	PATIENT	ID	: 1708179
COLLECTED BY	:	REG. NO./	LAB NO.	: 012412250001
REFERRED BY	:	REGISTRA	TION DATE	: 25/Dec/2024 07:18 AM
BARCODE NO.	: 01522950	COLLECTI	ON DATE	: 25/Dec/2024 07:25AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPORTI	NG DATE	: 25/Dec/2024 11:14AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AN	MBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		CLINICAL PATHO	LOGY	
		TINE & MICROSCOP		ATION
PHYSICAL EXAMIN	NATION			
QUANTITY RECIEV	ED TANCE SPECTROPHOTOMETRY	10	ml	
COLOUR	TANCE SPECTROPHOTOMETRY	PALE YELLOW		PALE YELLOW
TRANSPARANCY	TANCE SPECTROPHOTOMETRY	CLEAR		CLEAR
SPECIFIC GRAVITY		>=1.030		1.002 - 1.030
CHEMICAL EXAMI				
REACTION	TANCE SPECTROPHOTOMETRY	ACIDIC		
PROTEIN		Negative		NEGATIVE (-ve)
SUGAR	TANCE SPECTROPHOTOMETRY	4+		NEGATIVE (-ve)
pH	TANCE SPECTROPHOTOMETRY	<=5.0		5.0 - 7.5
BILIRUBIN	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
NITRITE	TANCE SPECTROPHOTOMETRY.	Negative		NEGATIVE (-ve)
UROBILINOGEN	TANCE SPECTROPHOTOMETRY	Normal	EU/dL	0.2 - 1.0
KETONE BODIES	TANCE SPECTROPHOTOMETRY	Negative		NEGATIVE (-ve)
BLOOD	TANCE SPECTROPHOTOMETRY	TRACE		NEGATIVE (-ve)
ASCORBIC ACID	TANCE SPECTROPHOTOMETRY	NEGATIVE (-ve)		NEGATIVE (-ve)
RED BLOOD CELLS		2-3	/HPF	0 - 3

Chopre

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Page 19 of 20

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EXCELLENCE IN HEALTHCARE & DIAGNOSTICS

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

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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMBALA CANTT				
			/		
Test Name	Value	Unit	Biological Reference interval		

rest Name	value	UIIIt	Diviogical Meler ence inter var
PUS CELLS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	1-2	/HPF	0 - 5
EPITHELIAL CELLS by MICROSCOPY ON CENTRIFUGED URINARY SEDIMENT	0-1	/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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