



	Dr. Vinay Chop MD (Pathology & Mi Chairman & Consult	icrobiology)		(Pathology)
NAME AGE/ GENDER COLLECTED BY REFERRED BY BARCODE NO. CLIENT CODE. CLIENT ADDRESS	: Mr. SANJAY SINGH : 49 YRS/MALE : SURJESH : : 01517656 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AM	IBALA CANTT	PATIENT ID REG. NO./LAB NO. REGISTRATION DATE COLLECTION DATE REPORTING DATE	: 1624648 : 012409250008 : 25/Sep/2024 08:36 AM : 25/Sep/2024 08:55AM : 25/Sep/2024 09:15AM
Test Name		Value	Unit	Biological Reference interval
	CO		ELLNESS PANEL: Y DOD COUNT (CBC)	
	BCS) COUNT AND INDICES	1/ 2	ana (all	120, 170
HAEMOGLOBIN (HB) by CALORIMETRIC		16.2	gm/dL	12.0 - 17.0
RED BLOOD CELL (RB	C) COUNT OCUSING, ELECTRICAL IMPEDENCE	5.47 ^H	Millions/c	mm 3.50 - 5.00
PACKED CELL VOLUM	IE (PCV)	50.3	%	40.0 - 54.0
by CALCULATED BY A MEAN CORPUSCULAI	UTOMATED HEMATOLOGY ANALYZER R VOLUME (MCV)	91.9	fL	80.0 - 100.0
by CALCULATED BY A	UTOMATED HEMATOLOGY ANALYZER R HAEMOGLOBIN (MCH)	29.5	20	27.0 - 34.0
by CALCULATED BY A	UTOMATED HEMATOLOGY ANALYZER		pg	
	R HEMOGLOBIN CONC. (MCHC)	32.1	g/dL	32.0 - 36.0
RED CELL DISTRIBUT	ION WIDTH (RDW-CV)	13.9	%	11.00 - 16.00
	UTOMATED HEMATOLOGY ANALYZER ION WIDTH (RDW-SD)	48	fL	35.0 - 56.0
by CALCULATED BY A MENTZERS INDEX by CALCULATED	UTOMATED HEMATOLOGY ANALYZER	16.8	RATIO	BETA THALASSEMIA TRAIT: < 13.0 IRON DEFICIENCY ANEMIA: >13.0
GREEN & KING INDE	X	23.26	RATIO	BETA THALASSEMIA TRAIT:<= 65.0 IRON DEFICIENCY ANEMIA: > 65.0
WHITE BLOOD CELLS	<u>(WBCS)</u>			
TOTAL LEUCOCYTE C	DUNT (TLC) BY SF CUBE & MICROSCOPY	8550	/cmm	4000 - 11000
NUCLEATED RED BLC	OOD CELLS (nRBCS)	NIL		0.00 - 20.00
NUCLEATED RED BLC	UTOMATED HEMATÓLOGY ANALYZER	NIL	%	< 10 %
NEUTROPHILS	BY SF CUBE & MICROSCOPY	67	%	50 - 70





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Dr. Vinay Chopra Dr. Yugam Chopra MD (Pathology & Microbiology) MD (Pathology) Chairman & Consultant Pathologist **CEO & Consultant Pathologist** NAME : Mr. SANJAY SINGH AGE/ GENDER : 49 YRS/MALE **PATIENT ID** :1624648 **COLLECTED BY** : SURJESH :012409250008 REG. NO./LAB NO. **REFERRED BY REGISTRATION DATE** : 25/Sep/2024 08:36 AM : **BARCODE NO.** :01517656 **COLLECTION DATE** : 25/Sep/2024 08:55AM CLIENT CODE. : KOS DIAGNOSTIC LAB **REPORTING DATE** : 25/Sep/2024 09:15AM **CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT Test Name Value Unit **Biological Reference interval** LYMPHOCYTES 20 - 40 24 % by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY EOSINOPHILS 2 % 1 - 6 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY MONOCYTES 7 % 2 - 12 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY BASOPHILS 0 % 0 - 1 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE LEUKOCYTES (WBC) COUNT ABSOLUTE NEUTROPHIL COUNT 5729 /cmm 2000 - 7500 by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY 2052 800 - 4900 ABSOLUTE LYMPHOCYTE COUNT /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE EOSINOPHIL COUNT 171 40 - 440 /cmm by FLOW CYTOMETRY BY SF CUBE & MICROSCOPY ABSOLUTE MONOCYTE COUNT 598 80 - 880 /cmm 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. 150000 - 450000 PLATELET COUNT (PLT) 167000 /cmm by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE 0.10 - 0.36 PLATELETCRIT (PCT) 0.22 % by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE MEAN PLATELET VOLUME (MPV) 13^H fL 6.50 - 12.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET LARGE CELL COUNT (P-LCC) 77000 30000 - 90000 /cmm by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE PLATELET LARGE CELL RATIO (P-LCR) 46^H % 11.0 - 45.0 by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE 15.0 - 17.0 PLATELET DISTRIBUTION WIDTH (PDW) 16.6 %

by HYDRO DYNAMIC FOCUSING, ELECTRICAL IMPEDENCE NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD

Chopra

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	Dr. Vinay Cho MD (Pathology & Chairman & Cons			(Pathology)	
NAME	: Mr. SANJAY SINGH				
AGE/ GENDER	: 49 YRS/MALE		PATIENT ID	: 1624648	
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BARCODE NO.	:01517656		COLLECTION DATE	: 25/Sep/2024 08:55AM	
CLIENT CODE.	: KOS DIAGNOSTIC LAB		REPORTING DATE	: 25/Sep/2024 02:17PM	
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	AMBALA CANTI			
Test Name		Value	Unit	Biological Reference interval	
	GLY	COSYLATED H	AEMOGLOBIN (HBA1C)		
GLYCOSYLATED HAEI WHOLE BLOOD	MOGLOBIN (HbA1c):	5.5	%	4.0 - 6.4	
ESTIMATED AVERAG		111.15	mg/dL	60.00 - 140.00	
	AS PER AMERICAN				
	REFERENCE GROUP	G	GLYCOSYLATED HEMOGLOGIB (HBAIC) in %		
	abetic Adults >= 18 years		<5.7		
	t Risk (Prediabetes)		5.7 – 6.4 >= 6.5		
U	Diagnosing Diabetes		>= 0.5 Age > 19 Years		
		Goal	s of Therapy:	< 7.0	
Therapeut	tic goals for glycemic control		ns Suggested:	>8.0	
			Age < 19 Years		
		Goa	l of therapy:	<7.5	

COMMENTS:

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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CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPO	DRTING DATE	: 25/Sep/2024 09:28AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AM	BALA CANTT		
Test Name		Value	Unit	Biological Reference interval
	ERYTHRO	OCYTE SEDIMEN	TATION RATE (ES	R)
	MENTATION RATE (ESR) GATION BY CAPILLARY PHOTOMETRY	5	mm/1st h	ur 0 - 20
polycythaemia), sign as sickle cells in sickl NOTE: 1. ESR and C - reactive 2. Generally, ESR doe 3. CRP is not affected 4. If the ESR is elevate 5. Women tend to ha 6. Drugs such as dext	N ESR n with conditions that inhibit the no- ificantly high white blood cell coun e cell anaemia) also lower the ESR. e protein (C-RP) are both markers of s not change as rapidly as does CRP by as many other factors as is ESR, m ed, it is typically a result of two type we a higher ESR, and menstruation a	t (leucocytosis) , an inflammation. , either at the start naking it a better m so of proteins, globu ind pregnancy can ca	d some protein abno of inflammation or a: arker of inflammatior lins or fibrinogen. ause temporary eleva	rmalities. Šome changes in red cell shape (si s it resolves. 1.





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		Chopra gy & Microbiology) Consultant Pathologist	Dr. Yugam MD CEO & Consultant	(Pathology)
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CLIENT ADDRESS	: 6349/1, NICHOLSON RO	AD, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
	CI	LINICAL CHEMIST	RY/BIOCHEMISTR	Y
		GLUCOSE	FASTING (F)	
	F): PLASMA	99.88	mg/dL	NORMAL: < 100.0

KOS Diagnostic Lab (A Unit of KOS Healthcare)

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.
 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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CLIENT ADDRESS : 6349/1, NIC	CHOLSON ROAD, AMBALA CANTT	Unit	Piological Deference interval
	Value	Unit	Biological Reference interval
	LIPID PROF	ILE : BASIC	
CHOLESTEROL TOTAL: SERUM by CHOLESTEROL OXIDASE PAP	166.3	mg/dL	OPTIMAL: < 200.0 BORDERLINE HIGH: 200.0 - 239.0 HIGH CHOLESTEROL: > OR = 240.
TRIGLYCERIDES: SERUM by GLYCEROL PHOSPHATE OXIDASE (118.31 enzymatic)	mg/dL	OPTIMAL: < 150.0 BORDERLINE HIGH: 150.0 - 199.0 HIGH: 200.0 - 499.0 VERY HIGH: > OR = 500.0
HDL CHOLESTEROL (DIRECT): SERU by SELECTIVE INHIBITION	M 57.24	mg/dL	LOW HDL: < 30.0 BORDERLINE HIGH HDL: 30.0 - 60.0 HIGH HDL: > OR = 60.0
LDL CHOLESTEROL: SERUM by CALCULATED, SPECTROPHOTOME	85.4 TRY	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 CHOLESTEROL: SERUM by CALCULATED, SPECTROPHOTOME	109.06 TRY	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 CHOLESTEROL: SERUM by CALCULATED, SPECTROPHOTOME	23.66	mg/dL	0.00 - 45.00
TOTAL LIPIDS: SERUM by CALCULATED, SPECTROPHOTOME	450.91	mg/dL	350.00 - 700.00
CHOLESTEROL/HDL RATIO: SERUM by CALCULATED, SPECTROPHOTOME	2.91	RATIO	LOW RISK: 3.30 - 4.40 AVERAGE RISK: 4.50 - 7.0 MODERATE RISK: 7.10 - 11.0 HIGH RISK: > 11.0
LDL/HDL RATIO: SERUM by CALCULATED, SPECTROPHOTOME	1.49 TRY	RATIO	LOW RISK: 0.50 - 3.0 MODERATE RISK: 3.10 - 6.0 HIGH RISK: > 6.0

KOS Diagnostic Lab (A Unit of KOS Healthcare)



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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAI	D, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
TRIGLYCERIDES/HD		2.07 ^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
Test Marine		Value	Unit	biological Reference interval
	LIV	ER FUNCTION	I TEST (COMPLETE)	
BILIRUBIN TOTAL: S by diazotization, s	ERUM PECTROPHOTOMETRY	1.79 ^H	mg/dL	INFANT: 0.20 - 8.00 ADULT: 0.00 - 1.20
	CONJUGATED): SERUM	0.42 ^H	mg/dL	0.00 - 0.40
BILIRUBIN INDIRECT	(UNCONJUGATED): SERUM	1.37 ^H	mg/dL	0.10 - 1.00
SGOT/AST: SERUM	RIDOXAL PHOSPHATE	23.4	U/L	7.00 - 45.00
SGPT/ALT: SERUM	RIDOXAL PHOSPHATE	31.9	U/L	0.00 - 49.00
AST/ALT RATIO: SER	UM	0.73	RATIO	0.00 - 46.00
ALKALINE PHOSPHA		96.22	U/L	40.0 - 130.0
GAMMA GLUTAMYL by SZASZ, SPECTROF	TRANSFERASE (GGT): SERUM	32.15	U/L	0.00 - 55.0
TOTAL PROTEINS: SE	RUM	6.51	gm/dL	6.20 - 8.00
ALBUMIN: SERUM		4.24	gm/dL	3.50 - 5.50
GLOBULIN: SERUM		2.27 ^L	gm/dL	2.30 - 3.50
A : G RATIO: SERUM		1.87	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 Name		Value Unit	Biological 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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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,		LEI OKTING DATE	. 20/3CP/ 2024 10.30AM
Test Name		Value	Unit	Biological Reference interval
	кі	DNEY FUNCTION	N TEST (COMPLETE)	
UREA: SERUM		17.66	mg/dL	10.00 - 50.00
by UREASE - GLUTAN	AATE DEHYDROGENASE (GLDH)			
CREATININE: SERUN		1.04	mg/dL	0.40 - 1.40
		8.25	ma/dl	7.0 - 25.0
BLOOD UREA NITROGEN (BUN): SERUM by CALCULATED, SPECTROPHOTOMETRY		0.25	mg/dL	7:0 - 25:0
BLOOD UREA NITROGEN (BUN)/CREATININE		7.93 ^L	RATIO	10.0 - 20.0
RATIO: SERUM				
		16.98	RATIO	
UREA/CREATININE F	ECTROPHOTOMETRY	10.90	RATIO	
URIC ACID: SERUM		4.89	mg/dL	3.60 - 7.70
by URICASE - OXIDAS	SE PEROXIDASE			
CALCIUM: SERUM		9.28	mg/dL	8.50 - 10.60
by ARSENAZO III, SPE PHOSPHOROUS: SEF		2.59	mg/dL	2.30 - 4.70
	DATE, SPECTROPHOTOMETRY	2.37	ilig/ uL	2.30 - 7.70
ELECTROLYTES				
sodium: serum		138.3	mmol/L	135.0 - 150.0
by ISE (ION SELECTIV				
POTASSIUM: SERUM		4.4	mmol/L	3.50 - 5.00
by ISE (ION SELECTIV CHLORIDE: SERUM	/E ELECTRODE)	103.73	mmol/l	90.0 - 110.0
by ISE (ION SELECTIV	/E ELECTRODE)	103.73	mmol/L	90.0 - 110.0
	RULAR FILTERATION RATE			
	RULAR FILTERATION RATE	88		
(eGFR): SERUM				
by CALCULATED				
INTERPRETATION:				

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.



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	Dr. Vinay ChopraDr. Yugam ChopraMD (Pathology & Microbiology)MD (Pathology)Chairman & Consultant PathologistCEO & Consultant Pathologist			
IAME	: Mr. SANJAY SINGH			
GE/ GENDER	: 49 YRS/MALE	PATIENT ID	: 1624648	
COLLECTED BY	: SURJESH	REG. NO./LAB NO.	: 012409250008	
REFERRED BY		REGISTRATION DA		6 AM
BARCODE NO.	: 01517656	COLLECTION DATE	1	
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPORTING DATE	1	
CLIENT CODE. CLIENT ADDRESS			. 20/ Sep/ 2024 10.50	JAIVI
LIENI ADDRESS	: 6349/1, NICHOLSON ROAD, A	AMDALA CANT I		
Test Name		Value Unit	Biological	Reference interval
 Low protein diet an Severe liver disease Other source of dec 	e. creased urea synthesis.			
5. Repeated dialysis (t 6. Inherited hyperamr 7. SIADH (syndrome o 8. Pregnancy. DECREASED RATIO (<1 1. Phenacimide therap 2. Rhabdomyolysis (re	0:1) WITH INCREASED CREATININE by (accelerates conversion of crea eleases muscle creatinine).	nt in blood). one) due to tubular secretion of urea. E:		
5. Repeated dialysis (t 6. Inherited hyperamr 7. SIADH (syndrome o 8. Pregnancy. DECREASED RATIO (<1) 1. Phenacimide therap 2. Rhabdomyolysis (re 3. Muscular patients to INAPPROPIATE RATIO: 1. Diabetic ketoacidos should produce an inc	nonemias (urea is virtually absen f inappropiate antidiuretic harmo 0:1) WITH INCREASED CREATININE by (accelerates conversion of crea eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false incl creased BUN/creatinine ratio).	nt in blood). one) due to tubular secretion of urea. E: atine to creatinine). crease in creatinine with certain meth		Il ratio when dehydratio
5. Repeated dialysis (t 6. Inherited hyperamr 7. SIADH (syndrome o 8. Pregnancy. DECREASED RATIO (<1) 1. Phenacimide therap 2. Rhabdomyolysis (re 3. Muscular patients to NAPPROPIATE RATIO: 1. Diabetic ketoacidos 5. hould produce an inc 2. Cephalosporin thera ESTIMATED GLOMERU	nonemias (urea is virtually absen f inappropiate antidiuretic harmo o:1) WITH INCREASED CREATININE by (accelerates conversion of crea eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false inco creased BUN/creatinine ratio). apy (interferes with creatinine me LAR FILTERATION RATE:	nt in blood). one) due to tubular secretion of urea. E: atine to creatinine). crease in creatinine with certain meth easurement).	odologies,resulting in norma	Il ratio when dehydratio
5. Repeated dialysis (u 6. Inherited hyperamr 7. SIADH (syndrome o 3. Pregnancy. DECREASED RATIO (<10 1. Phenacimide therap 2. Rhabdomyolysis (re 3. Muscular patients v NAPPROPIATE RATIO: 1. Diabetic ketoacidos should produce an inc 2. Cephalosporin thera ESTIMATED GLOMERU CKD STAGE	nonemias (urea is virtually absen f inappropiate antidiuretic harmo o:1) WITH INCREASED CREATININE by (accelerates conversion of crea eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false incu creased BUN/creatinine ratio). apy (interferes with creatinine me LAR FILTERATION RATE: DESCRIPTION	nt in blood). one) due to tubular secretion of urea. E: atine to creatinine). crease in creatinine with certain meth easurement). GFR (mL/min/1.73m2)	odologies,resulting in norma ASSOCIATED FINDINGS	Il ratio when dehydratio
5. Repeated dialysis (t 6. Inherited hyperamr 7. SIADH (syndrome o 8. Pregnancy. DECREASED RATIO (<10 1. Phenacimide therap 2. Rhabdomyolysis (re 3. Muscular patients to INAPPROPIATE RATIO: 1. Diabetic ketoacidos should produce an inc 2. Cephalosporin thera ESTIMATED GLOMERU CKD STAGE G1	nonemias (urea is virtually absen f inappropiate antidiuretic harmo 0:1) WITH INCREASED CREATININE by (accelerates conversion of crea eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false incl creased BUN/creatinine ratio). apy (interferes with creatinine me LAR FILTERATION RATE: DESCRIPTION Normal kidney functi	nt in blood). one) due to tubular secretion of urea. E: atine to creatinine). crease in creatinine with certain meth easurement). GFR (mL/min/1.73m2) ion >90	odologies,resulting in norma ASSOCIATED FINDINGS No proteinuria_	Il ratio when dehydratio
5. Repeated dialysis (u 6. Inherited hyperamr 7. SIADH (syndrome o 8. Pregnancy. DECREASED RATIO (<10 1. Phenacimide therap 2. Rhabdomyolysis (re 3. Muscular patients v INAPPROPIATE RATIO: 1. Diabetic ketoacidos should produce an inc 2. Cephalosporin thera ESTIMATED GLOMERU CKD STAGE	nonemias (urea is virtually absen f inappropiate antidiuretic harmo o:1) WITH INCREASED CREATININE by (accelerates conversion of crea eleases muscle creatinine). who develop renal failure. sis (acetoacetate causes false incu creased BUN/creatinine ratio). apy (interferes with creatinine me LAR FILTERATION RATE: DESCRIPTION	nt in blood). one) due to tubular secretion of urea. E: atine to creatinine). crease in creatinine with certain meth easurement). GFR (mL/min/1.73m2) ion >90 h >90	odologies,resulting in norma ASSOCIATED FINDINGS	Il ratio when dehydratio

Severe decrease in GFR



G3b

G4

G5

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Moderate decrease in GFR

Kidney failure

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30-59

15-29

<15









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			/
Test Name		Value 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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CLIENT ADDRESS	: 6349/1, NICHOI				. 20/ 500/ 202 1 10.00/10
CLIENT ADDRESS	. 0040/ 1, 10101	LJOIN ROAD, AN	IDALA CANT I		
Test Name			Value	Unit	Biological Reference interval
IRON: SERUM	TROPHOTOMETRY		IRON 93.9	PROFILE μg/dL	59.0 - 158.0
UNSATURATED IROI		TY (UIBC)	148.55 ^L	μg/dL	150.0 - 336.0
:SERUM			110.00	10.	
TOTAL IRON BINDIN :SERUM			242.45	μg/dL	230 - 430
by SPECTROPHOTOM			00 70	04	
%TRANSFERRIN SAT	URATION: SERUM ECTROPHOTOMETERY		38.73	%	15.0 - 50.0
TRANSFERRIN: SERU	UM	(1 21(21)2)	172.14 ^L	mg/dL	200.0 - 350.0
INTERPRETATION:-					
VARIAE SERUM I		Nemia OF CHRO Normal to R		IRON DEFICIENCY ANEMIA Reduced	THALASSEMIA α/β TRAIT Normal
TOTAL IRON BIND		Decrea		Increased	Normal
% TRANSFERRIN S		Decrea		Decreased < 12-15 %	Normal
	DDITIN	NI 11-1		D	

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

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 BÍNDING CAPACITY (TÍBC):

SERUM FERRITIN:

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

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 or Increased





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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AM	MBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		ENDOCE	RINOLOGY	
	ТН	YROID FUNCT	TION TEST: TOTAL	
TRIIODOTHYRONINE		1.018	ng/mL	0.35 - 1.93
THYROXINE (T4): SER	ESCENT MICROPARTICLE IMMUNOASS/ UM ESCENT MICROPARTICLE IMMUNOASS/	9.23	µgm/dL	4.87 - 12.60
THYROID STIMULATI	NG HORMONE (TSH): SERUM ESCENT MICROPARTICLE IMMUNOASS/	0.639	µIU/mL	0.35 - 5.50

trilodothyronine (T3).Failure at any level of regulation of the hypothalamic-pituitary-thyroid axis will result in either underproduction (hypothyroidism) or overproduction(hyperthyroidism) of T4 and/or T3.

 CLINICAL CONDITION
 T3
 T4
 TSH

 Primary Hypothyroidism:
 Reduced
 Increased (Significantly)

 Subclinical Hypothyroidism:
 Normal or Low Normal
 High

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 (eg: phenytoin , salicylates).

3. Serum T4 levles 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 hypothroidism, pregnancy, phenytoin therapy.

TRIIODOTH	YRONINE (T3)	THYROXINE (T4)		THYROID STIMULATING HORMONE (T	
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





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AGE/ GENDER	: 49 YRS/N	IALE		PATIENT ID	: 1624648	
COLLECTED BY	: SURJESH			REG. NO./LAB NO.	: 0124092	50008
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CLIENT CODE.	: KOS DIAO	GNOSTIC LAB		REPORTING DATE	: 25/Sep/20	024 10:29AM
CLIENT ADDRES	S : 6349/1, 1	NICHOLSON ROAD,	AMBALA CANTT			
Test Name			Value	Uni	t Bio	ological Reference interval
6 - 12 Months	0.74 - 2.40	6 - 12 Months	7.10 - 16.16	6 – 12 Months	0.70 - 7.00	
1 - 10 Years	0.92 - 2.28	1 - 10 Years	6.00 - 13.80	1 – 10 Years	0.60 - 5.50	—

0.92 - 2.28	1 - 10 Years	6.00 - 13.80	1 – 10 Years	0.60 - 5.50
0.35 - 1.93	11 - 19 Years	4.87- 13.20	11 – 19 Years	0.50 - 5.50
0.35 - 1.93	> 20 Years (Adults)	4.87 - 12.60	> 20 Years (Adults)	0.35- 5.50
RECOM	VIENDATIONS OF TSH LE	VELS DURING PREGN	IANCY (µIU/mL)	
1st Trimester			0.10 - 2.50	
2nd Trimester		0.20 - 3.00		
3rd Trimester			0.30 - 4.10	
	0.35 - 1.93 0.35 - 1.93 RECOMI 1st Trimester 2nd Trimester	0.35 - 1.93 11 - 19 Years 0.35 - 1.93 > 20 Years (Adults) RECOMMENDATIONS OF TSH LE 1st Trimester 2nd Trimester	0.35 - 1.93 11 - 19 Years 4.87 - 13.20 0.35 - 1.93 > 20 Years (Adults) 4.87 - 12.60 RECOMMENDATIONS OF TSH LEVELS DURING PREGN 1st Trimester 2nd Trimester	0.35 - 1.93 11 - 19 Years 4.87 - 13.20 11 - 19 Years 0.35 - 1.93 > 20 Years (Adults) 4.87 - 12.60 > 20 Years (Adults) RECOMMENDATIONS OF TSH LEVELS DURING PREGNANCY (μIU/mL) 1st Trimester 0.10 - 2.50 2nd Trimester 0.20 - 3.00

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, idonie 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 goitre & Thyroiditis.

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

3. Autonomously functioning Thyroid adenoma

4. Secondary pituatary or hypothalmic 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





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LIENT ADDRESS . 0349/1, NI	CHOLSON ROAD, AMBALA	CANTI	
Test Name	Va	lue Unit	Biological Reference interval
		VITAMINS	
		D/25 HYDROXY VITAMIN D3	
ITAMIN D (25-HYDROXY VITAMII by CLIA (CHEMILUMINESCENCE IMMU		.5 ^L ng/mL	DEFICIENCY: < 20.0
by CEIA (CHEMILOMINESCENCE IMMC	MOASSAT)		INSUFFICIENCY: 20.0 - 30.0
			SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0
NTERPRETATION:			
DEFICIENT:	< 2	0	ng/mL
INSUFFICIENT:	21 -		ng/mL
PREFFERED RANGE: INTOXICATION:	30 - 2		ng/mL ng/mL
conversion of 7- dihvdrocholecalcife 2.25-OHVitamin D represents the f issue and tightly bound by a transp 3. Vitamin D plays a primary role in obosphate reabsorption, skeletal ca 4. Severe deficiency may lead to fail DECREASED: 1. Lack of sunshine exposure. 2. Inadequate intake, malabsorption 3. Depressed Hepatic Vitamin D 25- 4. Secondary to advanced Liver disea 5. Osteoporosis and Secondary Hype 5. Enzyme Inducing drugs: anti-epile NCREASED: 1. Hypervitaminosis D is Rare, and is severe hypercalcemia and hyperpho CAUTION: Replacement therapy in c hypervitaminosis D	erol to Vitamin D3 in the sk main body resevoir and tra- bort protein while in circula the maintenance of calcium licium deposition, calcium ure to mineralize newly for (celiac disease) hydroxylase activity ase erparathroidism (Mild to M ptic drugs like phenytoin, p is seen only after prolonged ophatemia. leficient individuals must b compare to whites, is at high	kin upon Ultraviolet exposure. Insport form of Vitamin D and tran- ation. In homeostatis. It promotes calciu mobilization, mainly regulated by rmed osteoid in bone, resulting in boderate deficiency) obenobarbital and carbamazepine I exposure to extremely high dose e monitored by periodic assessment	nolecalciferol (from animals, Vitamin D3), or by hsport form of Vitamin D, being stored in adipose um absorption, renal calcium absorption and y parathyroid harmone (PTH). h rickets in children and osteomalacia in adults. e, that increases Vitamin D metabolism. es of Vitamin D. When it occurs, it can result in ent of Vitamin D levels in order to prevent ficiency due to excess of melanin pigment which





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Test Name		Value	Unit	Biological Reference interval
/ITAMIN B12/COBA		VITAMIN B12/C	D BALAMIN pg/mL	190.0 - 890.0
by CMIA (CHEMILUMIN INTERPRETATION:-	ESCENT MICROPARTICLE IMMUNO	194	pg/mL	
by CMIA (CHEMILUMIN <u>INTERPRETATION:-</u> INCREAS	ESCENT MICROPARTICLE IMMUNO	194 ASSAY)		
by CMIA (CHEMILUMIN INTERPRETATION:- INCREAS 1.Ingestion of Vitam	ESCENT MICROPARTICLE IMMUNO ED VITAMIN B12 nin C	194 ASSAY) 1.Pregnancy	pg/mL	IB12
by CMIA (CHEMILUMIN INTERPRETATION:- INCREAS 1.Ingestion of Vitam 2.Ingestion of Estrog	ESCENT MICROPARTICLE IMMUNO. ED VITAMIN B12 hin C gen	194 ASSAY) 1.Pregnancy 2.DRUGS:Asp	pg/mL DECREASED VITAMIN irin, Anti-convulsants,	IB12
by CMIA (CHEMILUMIN INTERPRETATION:- INCREAS 1.Ingestion of Vitam	ESCENT MICROPARTICLE IMMUNO ED VITAMIN B12 hin C gen hin A	194 ASSAY) 1.Pregnancy	pg/mL DECREASED VITAMIN irin, Anti-convulsants, stion	IB12
by CMIA (CHEMILUMIN INTERPRETATION:- INCREAS 1.Ingestion of Vitam 2.Ingestion of Estroy 3.Ingestion of Vitam	ESCENT MICROPARTICLE IMMUNO. ED VITAMIN B12 nin C gen nin A jury	194 ASSAY) 1.Pregnancy 2.DRUGS:Asp 3.Ethanol Iges	pg/mL DECREASED VITAMIN irin, Anti-convulsants, stion ive Harmones	IB12

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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Test Name		Value	Unit	Biological Reference interval
		value	Onit	biological Reference interval
	PR		IR MARKER ANTIGEN (PSA) - TOT	AL
	ANTIGEN (PSA) - TOTAL:	0.62	ng/mL	0.0 - 4.0
INTERPRETATION: NOTE: 1. This is a recommen 2. False negative / po 3. PSA levels may app 4. Immediate PSA tes needle biopsy of pros 5. PSA values regardle correlated with clinic 6. Sites of Non-prosta 7. Physiological decre sexual activity 8. The concentration of in assay methods, cal RECOMMENDED TEST 1. Preoperatively (Bas 2. 2-4 Days Post oper 3. Prior to discharge f 4. Monthly Follow Up	sitive results are observed in ear consistently elevated / d ting following digital rectal e tate is not recommended as ess of levels should not be in al findings and results of oth atic PSA production are breas ease in PSA level by 18% has l of PSA in a given specimen, d ibration, and reagent specifi ING INTERVALS seline) atively	n patients receiving n epressed due to the i examination, ejaculat they falsely elevate le terpreted as absolute er investigations st epithelium, salivar been observed in hos etermined with assay city.	nouse monoclonal antibod interference by heterophi ion, prostatic massage, in evels e evidence of the presence y glands, peri-urethral & a spitalized / sedentary patio	tion (DRE) in males above 50 years of age. dies for diagnosis or therapy lic antibodies & nonspecific protein binding dwelling catheterization, ultrasonography and e or absence of disease. All values should be anal glands, cells of male urethra & breast milk ents either due to supine position or suspended turers, may not be comparable due to differences
	1st Year		Every 3 Months	
	2 nd Year		Every 4 Months	
2	rd Year Onwards		Every 6 Months	
CLINICAL USE: 1. An aid in the early of		when used in conjun		xamination in males more than 50 years of age

and in those with two or more affected first degree relatives. 2. Followup and management of Prostate cancer patients.

3. Detect metastatic or persistent disease in patients following surgical or medical treatment of Prostate cancer

KOS Diagnostic Lab (A Unit of KOS Healthcare)

INCREASED LEVEL:

- 1. Prostate cancer
- 2. Benign Prostatic Hyperplasia

3. Prostatitis



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	Dr. Vinay Chopra MD (Pathology & Microbiolog) Chairman & Consultant Pathol		(Pathology)
NAME	: Mr. SANJAY SINGH		
AGE/ GENDER	: 49 YRS/MALE	PATIENT ID	: 1624648
COLLECTED BY	: SURJESH	REG. NO./LAB NO.	: 012409250008
REFERRED BY	:	REGISTRATION DATE	: 25/Sep/2024 08:36 AM
BARCODE NO.	: 01517656	COLLECTION DATE	: 25/Sep/2024 08:55AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REPORTING DATE	: 25/Sep/2024 10:50AM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMBALA CAN	NTT	
			/
Test Name	Value	Unit	Biological Reference interval

4. Genitourinary infections

*** End Of Report ***



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