

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



M	Pr. Vinay Chopra D (Pathology & Microbiolog hairman & Consultant Patho	y) (v	am Chopra 1D (Pathology) ant Pathologist	
NAME : Mr. RAHUL				
AGE/ GENDER : 28 YRS/MALE		PATIENT ID	: 1779228	
COLLECTED BY :		REG. NO./LAB NO.	: 012503050021	
REFERRED BY		REGISTRATION DATE	E : 05/Mar/2025 10:45 AM	
BARCODE NO. : 01526495		COLLECTION DATE	:05/Mar/2025 11:21AM	
CLIENT CODE. : KOS DIAGNOS		REPORTING DATE	:05/Mar/2025 11:20AM	
CLIENT ADDRESS : 6349/1, NICH	OLSON ROAD, AMBALA CA	NTT		
Test Name	Value	Unit	Biological Reference i	nterval
	HAI	EMATOLOGY		
	COMPLETE	BLOOD COUNT (CBC)		
RED BLOOD CELLS (RBCS) COUNT	AND INDICES			
HAEMOGLOBIN (HB) by CALORIMETRIC	14.8	gm/dl	L 12.0 - 17.0	
RED BLOOD CELL (RBC) COUNT by hydro dynamic focusing, electri	4.4	Million	ns/cmm 3.50 - 5.00	
PACKED CELL VOLUME (PCV) by CALCULATED BY AUTOMATED HEMAT	44.4	%	40.0 - 54.0	
MEAN CORPUSCULAR VOLUME (MO	CV) 100. 9	H fL	80.0 - 100.0	
MEAN CORPUSCULAR HAEMOGLO		pg	27.0 - 34.0	
MEAN CORPUSCULAR HEMOGLOB by CALCULATED BY AUTOMATED HEMAT	N CONC. (MCHC) 33.3	g/dL	32.0 - 36.0	
RED CELL DISTRIBUTION WIDTH (by Calculated by automated hemat	OLOGY ANALYZER	%	11.00 - 16.00	
RED CELL DISTRIBUTION WIDTH (by CALCULATED BY AUTOMATED HEMAT		fL	35.0 - 56.0	
MENTZERS INDEX by CALCULATED	22.93	3 RATIC) BETA THALASSEMIA 13.0 IRON DEFICIENCY AN >13.0	
GREEN & KING INDEX by CALCULATED	32.3	RATIC) BETA THALASSEMIA 65.0 IRON DEFICIENCY AN 65.0	
<u>WHITE BLOOD CELLS (WBCS)</u>				
TOTAL LEUCOCYTE COUNT (TLC) by FLOW CYTOMETRY BY SF CUBE & MIC	1050 Roscopy	0 /cmm	4000 - 11000	
NUCLEATED RED BLOOD CELLS (n by AUTOMATED 6 PART HEMATOLOGY A	RBCS) NIL		0.00 - 20.00	
NUCLEATED RED BLOOD CELLS (n by CALCULATED BY AUTOMATED HEMAT		%	< 10 %	





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

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

KOS Central Lab: 6349/1, Nicholson Road, Ambala Cantt -133 001, Haryana KOS Molecular Lab: IInd Floor, Parry Hotel, Staff Road, Opp. GPO, Ambala Cantt - 133 001, Haryana 0171-2643898, +91 99910 43898 | care@koshealthcare.com | www.koshealthcare.com







Dr. Vinay Chopra



MD (Pathology & Microbiology) Chairman & Consultant Pathologist

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

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Test Name		Value	Unit	Biological Reference interval
DIFFERENTIAL LEUCOCY	TE COUNT (DLC)			
NEUTROPHILS by FLOW CYTOMETRY BY SF C		59	%	50 - 70
LYMPHOCYTES by FLOW CYTOMETRY BY SF C		30	%	20 - 40
EOSINOPHILS by FLOW CYTOMETRY BY SF C	CUBE & MICROSCOPY	6	%	1 - 6
MONOCYTES by FLOW CYTOMETRY BY SF C	CUBE & MICROSCOPY	5	%	2 - 12
BASOPHILS by FLOW CYTOMETRY BY SF C	CUBE & MICROSCOPY	0	%	0 - 1
ABSOLUTE LEUKOCYTES	<u>(WBC) COUNT</u>			
ABSOLUTE NEUTROPHIL by FLOW CYTOMETRY BY SF C		6195	/cmm	2000 - 7500
ABSOLUTE LYMPHOCYTE by FLOW CYTOMETRY BY SF C		3150	/cmm	800 - 4900
ABSOLUTE EOSINOPHIL C by FLOW CYTOMETRY BY SF C	OUNT	630 ^H	/cmm	40 - 440
ABSOLUTE MONOCYTE CO		525	/cmm	80 - 880
ABSOLUTE BASOPHIL COU by FLOW CYTOMETRY BY SF C		0	/cmm	0 - 110
PLATELETS AND OTHER	PLATELET PREDICTIV	E MARKERS.		
PLATELET COUNT (PLT) by HYDRO DYNAMIC FOCUSING	G, ELECTRICAL IMPEDENCE	355000	/cmm	150000 - 450000
PLATELETCRIT (PCT) by HYDRO DYNAMIC FOCUSING	G. ELECTRICAL IMPEDENCE	0.3	%	0.10 - 0.36
MEAN PLATELET VOLUME by HYDRO DYNAMIC FOCUSING	E (MPV)	8	fL	6.50 - 12.0
PLATELET LARGE CELL CO by HYDRO DYNAMIC FOCUSING		58000	/cmm	30000 - 90000
PLATELET LARGE CELL R. by HYDRO DYNAMIC FOCUSING		16.4	%	11.0 - 45.0
PLATELET DISTRIBUTION by hydro dynamic focusing NOTE: TEST CONDUCTED C	G, ELECTRICAL IMPEDENCE	15.8	%	15.0 - 17.0





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

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Tost Namo	Valuo	Imit	Biological Poforanco interval







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 0171-2643898, +91 99910 43898
 care@koshealthcare.com
 www.koshealthcare.com





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Test Name	T T	Value Unit	Biological Reference interval
IRON: SERUM		HEMISTRY/BIOCHEMIS IRON 110 μg/dL	
	DR.VINAY CHOPRA CONSULTANT PATHOLOGIST MBBS, MD (PATHOLOGY & MICROBIOLOGY Nicholson Road, Ambala Cantt -133 001, Ha		

KOS Central Lab: 6349/1, Nicholson Road, Ambala Cantt -133 001, Haryana KOS Molecular Lab: IInd Floor, Parry Hotel, Staff Road, Opp. GPO, Ambala Cantt -133 001, Haryana 0171-2643898, +91 99910 43898 | care@koshealthcare.com | www.koshealthcare.com







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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		FERR	RITIN	
FERRITIN: SERUM	ESCENCE IMMUNOASSAY)	123.83	ng/mL	21.81 - 274.66

INTERPRETATION:

TEST PERFORMED AT KOS DIAGNOSTIC LAB. AMBALA CANTT

Serum ferritin appears to be in equilibrium with tissue ferritin and is a good indicator of storage iron in normal subjects and in most disorders. In patients with some hepatocellular diseases, malignancies and inflammatory diseases, serum ferritin is a disproportionately high estimate of storage iron because serum ferritin is an acute phase reactant. In such disorders iron deficiency anemia may exist with a normal serum ferritin concentration. In the presence of inflammation, persons with low serum ferritin are likely to respond to iron therapy. DECREASED:

1. Iron depletion appears to be the only condition associated with reduced serum ferritin concentrations.

- Hypothyroidism.
 Vitamin-C deficiency

INCREASED FERRITIN DUE TO IRON OVERLOAD (PRIMARY):

1. Hemochromatosis or hemosiderosis.

Wilson Disease.

INCREASED FERRITIN DUE TO IRON OVERLOAD (SECONDARY):

- 1. Transfusion overload
- Excess dietary Iron
 Porphyria Cutanea tada

4. Ineffective erythropoiesis. INCREASED FERRITIN WITHOUT IRON OVERLOAD:

- 1. Liver disorders (NASH) or viral hepatitis (B/C)
- 2. Inflammatory conditions (Ferritin is a acute phase reactant) both acute and chronic.
- 3. Leukaemia, hodgkin's disease.
- 4. Alcohol excess.

5. Other malignancies in which increases probably reflect the escape of ferritin from damaged liver cells, impaired clearance from the plasma, synthesis of ferritin by tumour cells.

6. Ferritin levels below 10 ng/ml have been reported as indicative of iron deficiency anemia.

NOTE: 1. As Ferritin is an acute phase reactant, it is often raised in both acute and chronic inflammatory condition of the body such as infections leading to false positive results. It can thererfore mask a diagnostically low result. In such Cases serum ferritin levels should always be correlated with C-Reactive proteins to rule out any inflammatory conditions.

2. Patients with iron deficiency anaemia may occasionally have elevated or normal ferritin levels. This is usually seen in patients already receiving iron therapy or in patients with concomitant hepatocellular injury.



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

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

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CLIENT ADDRESS	: 6349/1, NICHOLSON ROA	D, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		MAGNES	IUM	
MAGNESIUM: SERU		2.16	mg/dL	1.6 - 2.6

KOS Diagnostic Lab (A Unit of KOS Healthcare)

INTERPRETATION:-

1. Magnesium along with potassium is a major intracellular cation.

2.Magnesium is a cofactor of many enzyme systems. All adenosine triphosphate (ATP)-dependent enzymatic reactions require magnesium as a cofactor. 3.Approximately 70% of magnesium ions are stored in bone. The remainder is involved in intermediary metabolic processes; about 70% is present in free form while the other 30% is bound to proteins (especially albumin), citrates, phosphate, and other complex formers. The serum magnesium level is kept constant within very narrow limits. Regulation takes place mainly via the kidneys, primarily via the ascending loop of Henle.

INCREASD (HYPERMAGNESIA):-Conditions that interfere with glomerular filtration result in retention of magnesium and hence elevation of serum concentrations.

1. Acute and chronic renal failure.

2.magnesium overload.

3. Magnesium release from the intracellular space.

4.Mild-to-moderate hypermagnesemia may prolong atrioventricular conduction time. Magnesium toxicity may result in central nervous system (CNS) depression, cardiac arrest, and respiratory arrest.

DECREASED (HYPOMAGNESIA):-

- 1.Chronic alcoholism.
- 2.Childhood malnutrition.
- 3. Malabsorption.
- 4. Acute pancreatitis.
- 5.Hypothyroidism.
- 6.Chronic glomerulonephritis.
- 7.Aldosteronism.
- 8. Prolonged intravenous feeding.

NOTE:-

Numerous studies have shown a correlation between magnesium deficiency and changes in calcium-, potassium-, and phosphate-homeostasis which are associated with cardiac disorders such as ventricular arrhythmias that cannot be treated by conventional therapy, increased sensitivity to digoxin, coronary artery spasms, and sudden death. Additional concurrent symptoms include neuromuscular and neuropsychiatric disorders.





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

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Test Name		Value	Unit	Biological Reference interv
			RINOLOGY	
			TION TEST: TOTA	
TRIIODOTHYRONI	NE (13): SERUM IESCENT MICROPARTICLE IMMUN	0.937 OASSAY)	ng/ml	0.35 - 1.93
THYROXINE (T4): S	SERUM IESCENT MICROPARTICLE IMMUN	6.28 OASSAY)	μgm/o	1L 4.87 - 12.60
	TING HORMONE (TSH): SE		µIU/m	L 0.35 - 5.50
3rd GENERATION, ULT		043347)		
INTERPRETATION:				
day has influence on the triiodothyronine (T3).Fai	measured serum TSH concentrations	s. TSH stimulates the pr	oduction and secretion of the	0 pm. The variation is of the order of 50%.Hence time o e metabolically active hormones, thyroxine (T4)and ther underproduction (hypothyroidism) or
CLINICAL CONDITION	T3		T4	TSH
Primary Hypothyroidis		d	Reduced	Increased (Significantly)
Subclinical Hypothyroi	dism: Normal or I	ow Normal	Normal or Low Normal	High

LIMITATIONS:-

Primary Hyperthyroidism:

Subclinical Hyperthyroidism:

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT

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)	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	

Increased

Normal or High Normal





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Test Name			Value	Unit	t	Biological Reference interval
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	
	RECON	IMENDATIONS OF TSH LE	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





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	٢	Dr. Vinay Chopra 1D (Pathology & Microbiolog Chairman & Consultant Patho	y) M	am Chopra ID (Pathology) ant Pathologist
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Test Name		Value	Unit	Biological Reference interval
	CIENT: FICIENT:	< 20 21 - 29		SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0 ng/mL
1.Vitamin D compour conversion of 7- dihy 2.25-OHVitamin D r tissue and tiahtly bou 3.Vitamin D plays a p ohosphate reabsorpt 4.Severe deficiency n DECREASED: 1.Lack of sunshine ex 2.Inadequate intake, 3.Depressed Hepatic 4.Secondary to advar 5.Osteoporosis and S 6.Enzyme Inducing di INCREASED: 1. Hypervitaminosis I severe hypercalcemia CAUTION: Replacemen hypervitaminosis D NOTE:-Dark coloured	rdrocholecalciferce epresents the ma und by a transpor primary role in the ion, skeletal calci may lead to failure posure. malabsorption (or Vitamin D 25- hy need Liver disease econdary Hyperp rugs: anti-epilepti D is Rare, and is se a and hyperphoph ent therapy in def <i>individuals as corr</i>	b) to Vitamin D3 in the skin u in body resevoir and transpo- rt protein while in circulation e maintenance of calcium ho ium deposition, calcium mot e to mineralize newly formed celiac disease) droxylase activity e parathroidism (Mild to Mode ic drugs like phenytoin, pher een only after prolonged exp natemia. icient individuals must be mo	upon Ultraviolet exposure. ort form of Vitamin D and train. omeostatis. It promotes calci- pilization, mainly regulated by d osteoid in bone, resulting in erate deficiency) hobarbital and carbamazepine posure to extremely high dose onitored by periodic assessm	ng/mL holecalciferol (from animals, Vitamin D3), or by nsport form of Vitamin D, being stored in adipos ium absorption, renal calcium absorption and y parathyroid harmone (PTH). n rickets in children and osteomalacia in adults. e, that increases Vitamin D metabolism. es of Vitamin D. When it occurs, it can result in thent of Vitamin D levels in order to prevent efficiency due to excess of melanin pigment which
CAUTION : Replacement hypervitaminosis D	ent therapy in def individuals as com	icient individuals must be m	31	





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







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LIENI ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANT I			
lest Name		Value	Unit	Biological Reference interval	
NTERPRETATION:-					
INCREASED VITAMIN B12				1.510	
			ECREASED VITAMIN	I B12	
1.Ingestion of Vitan	nin C	1.Pregnancy			
1.Ingestion of Vitan 2.Ingestion of Estro	nin C gen	1.Pregnancy 2.DRUGS:Aspirir	n, Anti-convulsants		
1.Ingestion of Vitan 2.Ingestion of Estro 3.Ingestion of Vitan	nin C gen nin A	1.Pregnancy 2.DRUGS:Aspirir 3.Ethanol Igestic	n, Anti-convulsants on		
1.Ingestion of Vitan 2.Ingestion of Estro 3.Ingestion of Vitan 4.Hepatocellular in	nin C gen nin A jury	1.Pregnancy 2.DRUGS:Aspirir	n, Anti-convulsants on Harmones		
1.Ingestion of Vitan 2.Ingestion of Estro 3.Ingestion of Vitan 4.Hepatocellular in 5.Myeloproliferativ 6.Uremia	nin C gen nin A jury e disorder	1.Pregnancy 2.DRUGS:Aspirir 3.Ethanol lgestic 4. Contraceptive 5.Haemodialysis 6. Multiple Myel	n, Anti-convulsants on Harmones s oma		
1.Ingestion of Vitan 2.Ingestion of Estro 3.Ingestion of Vitan 4.Hepatocellular in 5.Myeloproliferativ 6.Uremia .Vitamin B12 (cobal .In humans, it is ob .The body uses its v .xcreted. .Vitamin B12 deficie	nin C gen nin A jury re disorder lamin) is necessary for hematopo tained only from animal proteins itamin B12 stores very economic	1.Pregnancy 2.DRUGS:Aspirir 3.Ethanol Igestic 4. Contraceptive 5.Haemodialysis 6. Multiple Myel biesis and normal neuron and requires intrinsic faally, reabsorbing vitamin	n, Anti-convulsants on Harmones s oma al function. ctor (IF) for absorp B12 from the ileun	, Colchicine	

*** End Of Report ***

deficiency at the cellular level is the assay for MMA. If clinical symptoms suggest deficiency, measurement of MMA and homocysteine should be





0171-2643898, +91 99910 43898 | care@koshealthcare.com | www.koshealthcare.com

considered, even if serum vitamin B12 concentrations are normal.

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