



	ME	<b>: Vinay Cho</b> ) (Pathology & airman & Cons			(Pathology)
NAME	: Mrs. VAIBHAV	[			
AGE/ GENDER	: 28 YRS/FEMAL	Ε		PATIENT ID	: 1563942
COLLECTED BY	:			REG. NO./LAB NO.	: 012407290055
<b>REFERRED BY</b>	:			<b>REGISTRATION DATE</b>	: 29/Jul/2024 12:58 PM
BARCODE NO.	:01514079			COLLECTION DATE	: 29/Jul/2024 01:03PM
CLIENT CODE.	: KOS DIAGNOST	IC LAB		<b>REPORTING DATE</b>	: 29/Jul/2024 03:58PM
CLIENT ADDRESS	: 6349/1, NICHO	LSON ROAD, A	MBALA CANTT		
Test Name			Value	Unit	Biological Reference interva
		CLINI		TRY/BIOCHEMISTR' PROFILE	Y
IRON: SERUM			39.8	μg/dL	37.0 - 145.0
by FERROZINE, SPECTROPHOTOMETRY UNSATURATED IRON BINDING CAPACITY (UIBC) :SERUM by FERROZINE, SPECTROPHOTOMETERY		TY (UIBC)	375.91 <sup>H</sup>	µg/dL	150.0 - 336.0
TOTAL IRON BINDIN SERUM	IG CAPACITY (TIBC)		415.71	μg/dL	230 - 430
%TRANSFERRIN SAT			9.57 <sup>L</sup>	%	15.0 - 50.0
TRANSFERRIN: SERUM by SPECTROPHOTOMETERY (FERENE)		295.15	mg/dL	200.0 - 350.0	
INTERPRETATION:-	VILILAT (FERENE)				
VARIAE	BLES A	NEMIA OF CHI	RONIC DISEASE	IRON DEFICIENCY ANEMI	A THALASSEMIA α/β TRAIT
SERUM I	RON:	Normal to	Reduced	Reduced	Normal

VARIABLES	ANEMIA OF CHRONIC DISEASE	IRON DEFICIENCY ANEMIA	THALASSEMIA α/β TRAIT	
SERUM IRON:	Normal to Reduced	Reduced	Normal	
TOTAL IRON BINDING CAPACITY:	Decreased	Increased	Normal	
% TRANSFERRIN SATURATION:	Decreased	Decreased < 12-15 %	Normal	
SERUM FERRITIN:	Normal to Increased	Decreased	Normal or Increased	
IDON				

#### 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 BÍNDING CAPACITY (TÍBC):

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.



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	Dr. Vinay Ch MD (Pathology & Chairman & Cor		Dr. Yugam MD CEO & Consultant	(Pathology)
NAME	: Mrs. VAIBHAVI			
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Test Name		Value	Unit	Biological Reference interval
		FERRI	ΓΙΝ	
FERRITIN: SERUM by CLIA (CHEMILUMIN	ESCENCE IMMUNOASSAY)	8.99	ng/mL	4.63 - 204.0

**KOS Diagnostic Lab** (A Unit of KOS Healthcare)

# 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.

- 2. Hypothyroidism.
   3. Vitamin-C deficiency

### **INCREASED FERRITIN DUE TO IRON OVERLOAD (PRIMARY):**

- 1. Hemochromatosis or hemosiderosis.
- 2. Wilson Disease.

# INCREASED FERRITIN DUE TO IRON OVERLOAD (SECONDARY):

- 1. Transfusion overload
- 2. Excess dietary Iron
- 3. 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.



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CLIENT ADDRESS :	6349/1, NICHOLSON ROAD,	AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
	VI		AMINS YDROXY VITAMIN D3	
VITAMIN D (25-HYDRO by CLIA (CHEMILUMINES	XY VITAMIN D3): SERUM SCENCE IMMUNOASSAY)	21.3 <sup>L</sup>	ng/mL	DEFICIENCY: < 20.0 INSUFFICIENCY: 20.0 - 30.0 SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0
NTERPRETATION:	NIT	20		- ()
DEFICIE INSUFFIC				g/mL g/mL
PREFFERED		<u>30 - 100</u> > 100		g/mL
conversion of 7- dihvdrc 2.25-OHVitamin D reputissue and tiahtly bound 3.Vitamin D plays a prin phosphate reabsorption 4.Severe deficiency may <b>DECREASED:</b> 1.Lack of sunshine expor 2.Inadeguate intake, ma 3.Depressed Hepatic Vit 4.Secondary to advance 5.Osteoporosis and Secc 6.Enzyme Inducing drug <b>INCREASED:</b> 1. Hypervitaminosis D is severe hypercalcemia an <b>CAUTION:</b> Replacement hypervitaminosis D <b>NOTE:</b> -Dark coloured ind	becholecalciferol to Vitamin D3 resents the main body resevo d by a transport protein while hary role in the maintenance n, skeletal calcium deposition l lead to failure to mineralize sure. alabsorption (celiac disease) amin D 25- hydroxylase activ d Liver disease ondary Hyperparathroidism (I s: anti-epileptic drugs like pho- . Rare, and is seen only after p nd hyperphophatemia. therapy in deficient individual <i>lividuals as compare to whites</i> ,	3 in the skin upon ir and transport for in circulation. of calcium homer , calcium mobiliza newly formed ost ity Vild to Moderate enytoin, phenoba prolonged exposu Is must be monite	Ultraviolet exposure. orm of Vitamin D and trans ostatis. It promotes calciun ation, mainly regulated by p teoid in bone, resulting in r e deficiency) rbital and carbamazepine, re to extremely high doses pred by periodic assessmen	lecalciferol (from animals, Vitamin D3), or by port form of Vitamin D, being stored in adipos in absorption, renal calcium absorption and barathyroid harmone (PTH). ickets in children and osteomalacia in adults. that increases Vitamin D metabolism. of Vitamin D. When it occurs, it can result in it of Vitamin D levels in order to prevent <i>iency due to excess of melanin pigment which</i>
hypervitaminosis D	lividuals as compare to whites,			





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est Name		Value	Unit	Biological Reference interval
<u>NTERPRETATION:-</u> INCREAS	IESCENT MICROPARTICLE IMMUNO		DECREASED VITAMIN	N B12
1.Ingestion of Vitan		1.Pregn		
2.Ingestion of Estro 3.Ingestion of Vitan			S:Aspirin, Anti-convulsants ol Igestion	, Colchicine
4.Hepatocellular in			raceptive Harmones	
5.Myeloproliferative disorder			nodialysis	
6.Uremia	amin) is necessary for hemator		ple Myeloma	
In humans, it is ob The body uses its v xcreted. Vitamin B12 deficie eal resection, smal Vitamin B12 deficie proprioception, poor he neurologic defec	tained only from animal protein itamin B12 stores very economi ency may be due to lack of IF set intestinal diseases). ency frequently causes macrocy coordination, and affective bel ts without macrocytic anemia. nic acid and homocysteine leve	ns and requires in cally, reabsorbing cretion by gastric tic anemia, glossi navioral changes. Is are also elevate (IF) is recommend	trinsic factor (IF) for absorp vitamin B12 from the ileun mucosa (eg, gastrectomy, g tis, peripheral neuropathy, These manifestations may o ed in vitamin B12 deficiency ded to identify this potentia	n and returning it to the liver; very little is astric atrophy) or intestinal malabsorption (eg, weakness, hyperreflexia, ataxia, loss of occur in any combination; many patients have





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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD	, AMBALA CANT	Т	
Test Name		Value	Unit	Biological Reference interval
		VITAMIN B9/	FOLIC ACID/FOLATE	
	CID/FOLATE: SERUM escence immunoassay)	4.6	ng/mL	DEFICIENT: < 3.37 INTERMEDIATE: 3.37 - 5.38 NORMAL: > 5.38

# **INTERPRETATION**

RESULT IN ng/mL	REMARKS
0.35 – 3.37	DEFICIENT
3.38 - 5.38	INTERMEDIATE
5.39 - 100.00	NORMAL

# NOTE:

1. Drugs like Methotrexate & Leucovorin interfere with folate measurement

2. To differentiate vitamin B12 & folate deficiency, measurement of Methyl malonic acid in urine & serum Homocysteine level is suggested 3. Risk of toxicity from folic acid is low as it is a water soluble vitamin regularly excreted in urine

#### COMMENTS:

1. Folate plays an important role in the synthesis of purine & pyrimidines in the body and is important for the maturation of erythrocytes.

It is widely available from plants and to a lesser extent organ meats, but more than half the folate content of food is lost during cooking.
 Folate deficiency is commonly prevalent in alcoholic liver disease, pregnancy and the elderly. It may result from poor intestinal absorption, nutrition deficiency, excessive demand as in pregnancy or in malignancy and in response to certain drugs like Methotrexate & anticonvulsants.
 Decreased Levels Megaloblastic anemia, Infantile hyperthyroidism, Alcoholism, Malnutrition, Scurvy, Liver disease, B12 deficiency, dietary amino acid excess, adult Celiac disease, Tropical Sprue, Crohn's disease, Hemolytic anemias, Carcinomas, Myelofibrosis, vitamin B6 deficiency, pregnancy, Whipple's disease, extensive intestinal resection and severe exfoliative dermatitis

# \*\* End Of Report \*\*





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