

TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



	М	<b>Pr. Vinay Chopra</b> D (Pathology & Microbiology) hairman & Consultant Patholog		(Pathology)
NAME	: Mrs. POOJA M	ALHOTRA		
AGE/ GENDER	: 33 YRS/FEMAI	ĿE	PATIENT ID	: 1822248
COLLECTED BY	:		REG. NO./LAB NO.	: 012504080041
<b>REFERRED BY</b>	:		<b>REGISTRATION DATE</b>	: 08/Apr/2025 10:29 AM
BARCODE NO.	:01528594		COLLECTION DATE	: 08/Apr/2025 10:36AM
CLIENT CODE.	: KOS DIAGNOS	TIC LAB	<b>REPORTING DATE</b>	: 08/Apr/2025 11:06AM
CLIENT ADDRESS	: 6349/1, NICH	OLSON ROAD, AMBALA CANT	Т	
Test Name		Value	Unit	<b>Biological Reference interval</b>
	IB)	10.8 <sup>L</sup>	OGLOBIN (HB) gm/dL	12.0 - 16.0
ULE MOCI ODDU (I	IB)	10 eL	gm/dL	12.0 16.0
(		10.0	Surger	12.0 - 10.0
by CALORIMETRIC	_,	10.8	Sun all	12.0 - 10.0
<i>by CALORIMETRIC</i> INTERPRETATION:- Hemoglobin is the pr	otein molecule in			
<i>by CALORIMETRIC</i> INTERPRETATION:- Hemoglobin is the pr tissues back to the lu	rotein molecule in Ings.	red blood cells that carries ox	ygen from the lungs to the b	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lev ANEMIA (DECRESED	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN)</b> :	red blood cells that carries ox	ygen from the lungs to the b unt.	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lev ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ency (iron, vitamin	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate)	ygen from the lungs to the b unt.	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lev ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ncy (iron, vitamin olems (replacemen	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer)	ygen from the lungs to the b unt.	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lev ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by rev 5) Kidney failure	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ency (iron, vitamin blems (replacemen d blood cell synthe	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer) esis by chemotherapy drugs	ygen from the lungs to the b unt. stomach ulcer)	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lev ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by rev 5) Kidney failure 6) Abnormal hemogl	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ncy (iron, vitamin blems (replacemen d blood cell synthe obin structure (sic	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer) esis by chemotherapy drugs kle cell anemia or thalassemia	ygen from the lungs to the b unt. stomach ulcer)	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lew ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by red 5) Kidney failure 6) Abnormal hemogl POLYCYTHEMIA (INCI 1) People in higher a	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg incy (iron, vitamin olems (replacemen d blood cell synthe obin structure (sic <b>REASED HAEMOGL</b> ultitudes (Physiologi	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer) esis by chemotherapy drugs kle cell anemia or thalassemia <b>OBIN):</b>	ygen from the lungs to the b unt. stomach ulcer)	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lew ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by rev 5) Kidney failure 6) Abnormal hemogle POLYCYTHEMIA (INCE 1) People in higher a 2) Smoking (Seconda	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ncy (iron, vitamin blems (replacemen d blood cell synthe obin structure (sic <b>REASED HAEMOGL</b> Ititudes (Physiolog ry Polycythemia)	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer) esis by chemotherapy drugs kle cell anemia or thalassemia <b>OBIN):</b> gical)	ygen from the lungs to the b unt. stomach ulcer) a).	odys tissues and returns carbon dioxide from the
INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lev ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prote 4) Suppression by red 5) Kidney failure 6) Abnormal hemogl POLYCYTHEMIA (INCI 1) People in higher a 2) Smoking (Seconda 3) Dehydration prode 4) Advanced lung dise	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ncy (iron, vitamin blems (replacemen d blood cell synthe obin structure (sic <b>REASED HAEMOGL</b> Ititudes (Physiolog ry Polycythemia) uces a falsely rise i	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer) esis by chemotherapy drugs kle cell anemia or thalassemic <b>OBIN):</b> gical) n hemoglobin due to increase	ygen from the lungs to the b unt. stomach ulcer) a).	
by CALORIMETRIC INTERPRETATION:- Hemoglobin is the pr tissues back to the lu A low hemoglobin lew ANEMIA (DECRESED 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by rev 5) Kidney failure 6) Abnormal hemogl POLYCYTHEMIA (INCI 1) People in higher a 2) Smoking (Seconda 3) Dehydration produ 4) Advanced lung diss 5) Certain tumors	rotein molecule in ings. vel is referred to as <b>HAEMOGLOBIN):</b> umatic injury, surg ency (iron, vitamin blems (replacemen d blood cell synthe obin structure (sic <b>REASED HAEMOGL</b> iltitudes (Physiolog ry Polycythemia) uces a falsely rise i ease (for example,	red blood cells that carries ox s ANEMIA or low red blood cou ery, bleeding, colon cancer or B12, folate) t of bone marrow by cancer) esis by chemotherapy drugs kle cell anemia or thalassemic <b>OBIN):</b> gical) n hemoglobin due to increase	ygen from the lungs to the b unt. stomach ulcer) a).	

NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD





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







		C <b>hopra</b> & Microbiology) onsultant Pathologis		(Pathology)	
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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAI	), AMBALA CANTT			
Test Name		Value	Unit	<b>Biological Reference interval</b>	
	CLINIC		STRY/BIOCHEMIS	TRY	
		URI	C ACID		
URIC ACID: SERUM		5.3	mg/dL	2.50 - 6.80	
by URICASE - OXIDASI INTERPRETATION:-	E PEROXIDASE				
1.GOUT occurs when	high levels of Uric Acid in the	blood cause crystal	s to form & accumulate arc	ound a joint.	
2.Uric Acid is the end intestinal tract by mid		. Uric acid is excret	ed to a large degree by the	kidneys and to a smaller degree in the	
INCREASED:-	crobial degradation.				
(A).DUE TO INCREASE 1.Idiopathic primary					
2.Excessive dietary pu	irines (organ meats, legumes, a	nchovies, etc).			
3.Cytolytic treatment	of malignancies especially leu & myeloid metaplasia.	ikemais & lymphom	nas.		
5.Psoriasis.	a myelola metaplasia.				
6.Sickle cell anaemia	etc. DEXCREATION (BY KIDNEYS)				
1. Alcohol ingestion.	DEACREATION (BT KIDNETS)				
2.Thiazide diuretics. 3.Lactic acidosis.					
	ess than 2 grams per day ).				
5.Diabetic ketoacidos 6.Renal failure due to	sis or starvation.				
DECREASED:-					
(A).DUE TO DIETARY D					
2.Fanconi syndrome	of Zinc, Iron and molybdenum. & Wilsons disease.				
3. Multiple sclerosis.	opriate antidiuretic hormone (	SIADU) cocration 8	low puripo dist sto		
(B).DUE TO INCREASE	DEXCREATION				
1.Drugs:-Probenecid,	, sulphinpyrazone, aspirin dose	es (more than 4 gra	ms per day), corticosterroio	ds and ACTH, anti-coagulants and estrogens etc	2.
o section of the sect	2		1		

KOS Diagnostic Lab (A Unit of KOS Healthcare)





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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD	), AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		ENDOCR	RINOLOGY	
	TH	IYROID FUNCT	TION TEST: TOTAL	
TRIIODOTHYRON by CMIA (CHEMILUMIN	INE (T3): SERUM IESCENT MICROPARTICLE IMMUNO	0.954 ASSAY)	ng/mL	0.35 - 1.93
THYROXINE (T4): by CMIA (CHEMILUMIN	SERUM IESCENT MICROPARTICLE IMMUNO	10.6 ASSAY)	µgm/dL	4.87 - 12.60
	ATING HORMONE (TSH): S		µIU/mL	0.35 - 5.50
3rd GENERATION, ULT <u>INTERPRETATION</u> :	RASENSITIVE			
day has influence on the triiodothyronine (T3).Fai	measured serum TSH concentrations.	TSH stimulates the prod	duction and secretion of the m	m. The variation is of the order of 50%.Hence time of t etabolically active hormones, thyroxine (T4)and er underproduction (hypothyroidism) or
CLINICAL CONDITION	Т3		T4	TSH

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 (	
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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Test Name			Value	Uni	t	<b>Biological Reference interval</b>
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	
11- 19 Years	0.35 - 1.93	11 - 19 Years	4.87-13.20	11 – 19 Years	0.50 - 5.50	
> 20 years (Adults)	0.35 - 1.93	> 20 Years (Adults)	4.87 - 12.60	> 20 Years (Adults)	0.35-5.50	
	RECOM	MENDATIONS OF TSH LE	VELS DURING PREC	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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Test Name		Value	Unit	<b>Biological Reference interval</b>
	LUT	EINISING H	ORMONE (LH)	
	RMONE (LH): SERUM NESCENT MICROPARTICLE IMMUNOASSAY	10.38 7	mIU/mL	MALES: 0.57 - 12.07 FOLLICULAR PHASE: 1.80 - 11.78 MID-CYCLE PEAK: 7.59 - 89.08 LUTEAL PHASE: 0.56 - 14.0 POST MENOPAUSAL WITHOUT HRT: 5.16 - 61.99
hormone from the hy	pothalamus controls the secretion of	the gonadotrop	oins, FSH and LH, from th	nits (alpha and beta). Gonadotropin-releasing e anterior pituitary. ivided by a mid cycle surge of both LH and FSH

- into a follicular phase and a luteal phase.
- 3. This "LH surge" triggers ovulation thereby not only releasing the egg, but also initiating the conversion of the residual follicle into a corpus luteum that, in turn, produces progesterone to prepare the endometrium for a possible implantation.
  4. LH supports thecal cells in the ovary that provide androgens and hormonal precursors for estradiol production. LH in males acts on testicular interstitial cells of Leydig to cause increased synthesis of testosterone.
  The test is useful in the following situations:
  1. An ediumetin the output in ediumeting interstitian.

- 1. An adjunctin the evaluation of menstrual irregularities.
- 2. Evaluating patients with suspected hypogonadism
- 3. Predicting ovulation & Evaluating infertility
- Diagnosing pituitary disorders
- 5. In both males and females, primary hypogonadism results in an elevation of basal follicle-stimulating hormone and luteinizing hormone levels

### FSH AND LH ELEVTED IN:

- 1. Primary gonadal failure
- Complete testicular feminization syndrome
   Precocious puberty (either idiopathic or secondary to a central nervous system lesion)
- 4. Menopause
- Finary ovarian hypo dysfunction in females
   Polycystic ovary disease in females
   Primary hypogonadism in males
   LH IS DECREASED IN:

- 1.Primary ovarian hyper function in females
- 2. Primary hypergonadism in males
- NOTE

1 .FSH and LH are both decreased in failure of the pituitary or hypothalamus.



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Test Name		Value	Unit	<b>Biological Reference interval</b>
	FOLL	ICLE STIMULATIN	G HORMONE (FS	H)
	LATING HORMONE (FSH): iescence immunoassay)	SERUM 4.68	mIU/mL	FEMALE FOLLICULAR PHASE: 3.03 - 8.08 FEMALE MID-CYCLE PEAK: 2.55 - 16.69 FEAMLE LUTEAL PHASE: 1.38 5.47 FEMALE POST-MENOPAUSAL

An adjunct in the evaluation of menstrual irregularities.

Evaluating patients with suspected hypogonadism.

3. Predicting ovulation

4. Evaluating infertility

 Diagnosing pituitary disorders
 In both males and females, primary hypogonadism results in an elevation of basal follicle-stimulating hormone (FSH) and luteinizing hormone (LH) levels

# **FSH and LH LEVELS ELEVATED IN:**

1. Primary gonadal failure

2. Complete testicular feminization syndrome.

Precocious puberty (either idiopathic or secondary to a central nervous system lesion)
 Menopause (postmenopausal FSH levels are generally >40 IU/L)
 Primary ovarian hypofunction in females

- 6. Primary hypogonadism in males

### NOTE:

1. Normal or decreased FSH is seen in polycystic ovarian disease in females 2. FSH and LH are both decreased in failure of the pituitary or hypothalamus.



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Test Name		Value	Unit	<b>Biological Reference interval</b>	1	
rest i tunic		value	Cint	biological Reference Interval	]	
		PRO	LACTIN			
PROLACTIN: SERU by CMIA (CHEMILUMINE INTERPRETATION:	M ESCENT MICROPARTICLE IMMUNOASSAY)	25.26 <sup>H</sup>	ng/mL	3 - 25		
3.Physiological functi physiologic stimuli su newborn infant. <b>INCREASED (HYPERPRO</b> 1.Prolactin-secreting J 2.Functional and orga 3.Primary hypothyroi 4.Section compression 5.Chest wall lesions a 6.Ectopic tumors. 7.DRUGS:- Anti-Dopar receptors, or seroton Opiates, High doses of <b>SIGNIFICANCE</b> :	INCREASED (HYPERPROLACTEMIA): 1.Prolactin-secreting pituitary adenoma (prolactinoma, which is 5 times more frequent in females than males). 2.Functional and organic disease of the hypothalamus. 3.Primary hypothyroidism. 4.Section compression of the pituitary stalk. 5.Chest wall lesions and renal failure. 6.Ectopic tumors. 7.DRUGS:- Anti-Dopaminergic drugs like antipsychotic drugs, antinausea/antiemetic drugs, Drugs that affect CNS serotonin metabolism, serotonin receptors, or serotonin reuptake (anti-depressants of all classes, ergot derivatives, some illegal drugs such as cannabis), Antihypertensive drugs ,Opiates, High doses of estrogen or progesterone,anticonvulsants (valporic acid), anti-tuberculous medications (Isoniazid).					
from decreased musc 3. In males, prolactin le 4. In women, prolactin 5. Clear symptoms and 4. Mild to moderately	le mass and osteoporosis. evels >13 ng/mL are indicative of hyperp levels >27 ng/mL in the absence of preg l signs of hyperprolactinemia are ofter	prolactinemi gnancy and n absent in re not a reli	ia. postpartum lactation are in patients with serum prolac able guide for determining	<i>dicative of hyperprolactinemia.</i> tin levels <100 ng/mL. whether a prolactin-producing pituitary		
Prolactin values that e	exceed the reference values may be du symptoms of hyperprolactinemia are	ue to macro absent, or	pprolactin (prolactin bound pituitary imaging studies a	to immunoglobulin). Macroprolactin should be re not informative.		





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AGE/ GENDER	: 33 YRS/FEMALE		PATIENT ID	: 1822248	
COLLECTED BY	:		REG. NO./LAB NO.	: 012504080041	
<b>REFERRED BY</b>	:		<b>REGISTRATION DATE</b>	: 08/Apr/2025 10:29 AM	
BARCODE NO.	: 01528594		COLLECTION DATE	: 08/Apr/2025 10:36AM	
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 08/Apr/2025 02:52PM	
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AME	BALA CANTT			
Test Name		Value	Unit	Pielogiael Deference interval	1
Test Name		value	Unit	<b>Biological Reference interval</b>	
	T	ESTOSTE	CRONE: TOTAL		
TESTOSTERONE - ' by CMIA (CHEMILUMINI INTERPRETATION:	TOTAL: SERUM ESCENT MICROPARTICLE IMMUNOASSAY	0.31	ng/mL	0.0 - 0.80	
testosterone is in the i 3.The bioavailable fra- and bound to cortisol 4.The total testostero <b>CLINIC USE:</b> 1.Assesment of testic	free form. ction includes the free form and that binding globulin (CBG). It is the mos ne bound to SHBG fluctuates since S ular functions in males sutism and virilization in females (Males) e Hyperplasia disease ales)	weakly bo	und" to albumin (40% of the	globulin (ŠHBG). Less than 1% of the total a total in men and 20% of the total in women) le. disease, sex steroids and insulin.	





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

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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.



	<b>Dr. Vinay Chopra</b> MD (Pathology & Microl Chairman & Consultant		Dr. Yugam MD CEO & Consultant	(Pathology)
NAME AGE/ GENDER COLLECTED BY REFERRED BY	<b>: Mrs. POOJA MALHOTRA</b> : 33 YRS/FEMALE : :	RI	TIENT ID C. NO./LAB NO. CGISTRATION DATE	: 1822248 <b>: 012504080041</b> : 08/Apr/2025 10:29 AM
BARCODE NO. CLIENT CODE. CLIENT ADDRESS	: 01528594 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AMBAI	RI	DELECTION DATE	: 08/Apr/2025 10:36AM : 08/Apr/2025 04:40PM
Test Name		Value	Unit	<b>Biological Reference interval</b>
	DEHYDROEPIAN	DROSTER	<b>ΟΝΕ SULPHATE</b> ( μg/dL	<b>DHEA-S</b> ) 23.00 - 266.00
INTERPRETATION:- CLINICAL USE: 1. Marker for Adrena 2. Differential diagno INCREASED LEVELS: 1. Adrenogenital synd 2. Congenital Adrena 3. Adrenal Carcinoma 4. Virilizing tumor of 5. Cushing's disease, 6. Hirsutism 7. Polycystic ovarian DECREASED LEVELS: 1. Addison's disease 2. Adrenal Hypoplasi 3. Hyperlipidaemia 4. Psychoses 5. Psoriasis 6. Increasing age. NOTE:	a adrenal gland. pituitary dependent. Syndrome (PCOD)	vdrogenase, 2	1-hydroxylase and 11 be	
	DR.VINAY CHOPRA CONSULTANT PATHOLOGIST MBBS, MD (PATHOLOGY & MICROBIOLOG		chopra NT PATHOLOGIST (PATHOLOGY)	







	<b>Dr. Vinay Chopr</b> MD (Pathology & Mic Chairman & Consulta	robiology)		(Pathology)
NAME	: Mrs. POOJA MALHOTRA			
AGE/ GENDER	: 33 YRS/FEMALE		PATIENT ID	: 1822248
COLLECTED BY	:		REG. NO./LAB NO.	: 012504080041
<b>REFERRED BY</b>	:		<b>REGISTRATION DATE</b>	: 08/Apr/2025 10:31 AM
BARCODE NO.	:01528594		COLLECTION DATE	: 08/Apr/2025 10:36AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	:08/Apr/202504:40PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AME	BALA CANTT		
Test Name		Value	Unit	<b>Biological Reference interval</b>
		VIT	AMINS	
	VITAMI	N D/25 HY	DROXY VITAMIN D	93
	DROXY VITAMIN D3): SERUM escence immunoassay)	26.4 <sup>L</sup>	ng/mL	DEFICIENCY: < 20.0 INSUFFICIENCY: 20.0 - 30.0 SUFFICIENCY: 30.0 - 100.0 TOXICITY: > 100.0
INTERPRETATION:		20		

DEFICIENT:	< 20	ng/mL			
INSUFFICIENT:	21 - 29	ng/mL			
PREFFERED RANGE:	30 - 100	ng/mL			
INTOXICATION:	> 100	ng/mL	ĺ		

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-OH--Vitamin D represents the main body resevoir and transport form of Vitamin D and transport form of Vitamin D, being stored in adipose tissue 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 phosphate 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. DECREASED:

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)

6.Enzyme Inducing drugs: anti-epileptic drugs like phenytoin, phenobarbital and carbamazepine, that increases Vitamin D metabolism.

INCREASED:

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.

CAUTION: Replacement therapy in deficient individuals must be monitored by periodic assessment of Vitamin D levels in order to prevent hypervitaminosis D

NOTE:-Dark coloured individuals as compare to whites, is at higher risk of developing Vitamin D deficiency due to excess of melanin pigment which interefere with Vitamin D absorption.

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



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