



	Dr. Vinay Cho MD (Pathology & Chairman & Const	Microbiology)		(Pathology)
NAME	: Mrs. NISHA			
AGE/ GENDER	: 34 YRS/FEMALE		PATIENT ID	: 1622519
COLLECTED BY	:		REG. NO./LAB NO.	: 012409230051
<b>REFERRED BY</b>	: LOOMBA HOSPITAL (AMBALA	A CANTT)	<b>REGISTRATION DATE</b>	: 23/Sep/2024 02:34 PM
BARCODE NO.	:01517560		COLLECTION DATE	: 23/Sep/2024 02:57PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 23/Sep/2024 03:05PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	MBALA CANTI		
Test Name		Value	Unit	Biological Reference interval
HAEMOGLOBIN (HB		12.2	gm/dL	12.0 - 16.0
HAEMOGLOBIN (HB			GLOBIN (HB) am/dL	12.0 - 16.0
by CALORIMETRIC INTERPRETATION:-				
tissues back to the lu A low hemoglobin lev <b>ANEMIA ( DECRESED</b> I 1) Loss of blood (trau 2) Nutritional deficie 3) Bone marrow prob 4) Suppression by red 5) Kidney failure 6) Abnormal hemogle <b>POLYCYTHEMIA (INCF</b>	ngs. vel is referred to as ANEMIA or low	r red blood cour blon cancer or s w by cancer) erapy drugs	nt. stomach ulcer)	odys tissues and returns carbon dioxide from th
<ol> <li>2) Smoking (Seconda</li> <li>3) Dehydration prodution</li> <li>4) Advanced lung dise</li> <li>5) Certain tumors</li> <li>6) A disorder of the b</li> </ol>	ry Polycythemia) uces a falsely rise in hemoglobin d ease (for example, emphysema) one marrow known as polycythen	nia rubra vera,		e amount of oxygen available to the body by

chemically raising the production of red blood cells).





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

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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.

# NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD

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ISO 9001 : 2008 CERTI		KOS Healthcare)	EXCELLENCE IN HEALTHCARE	& DIAGNOSTICS	
		Chopra & Microbiology) onsultant Pathologist	Dr. Yugam MD ( CEO & Consultant	(Pathology)	
NAME AGE/ GENDER COLLECTED BY REFERRED BY BARCODE NO. CLIENT CODE. CLIENT ADDRESS	: Mrs. NISHA : 34 YRS/FEMALE : : LOOMBA HOSPITAL (AMB. : 01517560 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAI	ALA CANTT) REGIST Colle Repor	NT ID O./LAB NO. FRATION DATE CTION DATE STING DATE	: 1622519 <b>: 012409230051</b> : 23/Sep/2024 02:34 PM : 23/Sep/2024 02:57PM : 23/Sep/2024 03:08PM	
Test Name		Value	Unit	Biological Reference interv	al
ABO GROUP by SLIDE AGGLUTINATI RH FACTOR TYPE by SLIDE AGGLUTINATI	ION	D GROUP (ABO) AND F B POSITIVE			
Kos Ceptral Lab: 6349/1	DR.VINAY CHOPRA CONSULTANT PATHOLOGIST MBBS, MD (PATHOLOGY & MICH Nicholson Road, Ambala Cantt - 13		ATHOLOGIST		
	loor Parry Hotel Staff Road Opp		Harvana		

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CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AM	/IBALA CANT	г	
Test Name		Value	Unit	Biological Reference interval
		BLEEDI	NG TIME (BT)	
BLEEDING TIME (BT		1 MIN 45	SEC MINS	1 - 5



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Test Name		Value	Unit	Biological Reference interval
		CLOTTING TIME	(CT)	
CLOTTING TIME (CT) by CAPILLARY TUBE I		5 MIN 20 SEC	MINS	4 - 9



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	: KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD	_	REPORTING DATE	: 23/Sep/2024 04:00PM
CLIENT CODE. CLIENT ADDRESS Test Name		_	REPORTING DATE	Biological Reference interval
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD	, AMBALA CANTT Value		Biological Reference interval
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD	, AMBALA CANTT Value	Unit	Biological Reference interval

(after consumption of 75 gms of glucose) is recommended for all such patients. 3. A random glucose level of above 200 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 Names		Value	Unit	Biological Reference interval
by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN	ENDOCRI YROID STIMULATII A 1.768		
THYROID STIMULAT	ING HORMONE (TSH): SERUN iescent microparticle immung rasensitive	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) µIU/mL	) 0.35 - 5.50
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN iescent microparticle immung rasensitive AGE	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE	) 0.35 - 5.50 (μΙU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUNG RASENSITIVE AGE 0 – 5 DAYS	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE	) 0.35 - 5.50 (μΙU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN iescent microparticle immung rasensitive AGE	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE	) 0.35 - 5.50 (μΙU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUNG RASENSITIVE AGE 0 – 5 DAYS 6 Days – 2 Months	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL <u>REFFERENCE RANGE</u> 0.70 – 15.20 0.70 – 11.00	) 0.35 - 5.50 (µIU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUNG RASENSITIVE AGE 0 – 5 DAYS 6 Days – 2 Months 3 – 11 Months	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE 0.70 – 15.20 0.70 – 11.00 0.70 – 8.40	) 0.35 - 5.50 (µIU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUNO RASENSITIVE AGE 0 – 5 DAYS 6 Days – 2 Months 3 – 11 Months 1 – 5 Years 6 – 10 Years 11 - 15	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE 0.70 – 15.20 0.70 – 11.00 0.70 – 8.40 0.70 – 7.00	) 0.35 - 5.50 (µlU/mL) )
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUNO RASENSITIVE AGE 0 – 5 DAYS 6 Days – 2 Months 3 – 11 Months 1 – 5 Years 6 – 10 Years	ENDOCRI YROID STIMULATII A 1.768 DASSAY)	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE 0.70 – 15.20 0.70 – 11.00 0.70 – 8.40 0.70 – 7.00 0.60 – 5.50	) 0.35 - 5.50 (µlU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUN RASENSITIVE AGE 0 – 5 DAYS 6 Days – 2 Months 3 – 11 Months 1 – 5 Years 6 – 10 Years 11 - 15 > 20 Years (Adults)	ENDOCRI YROID STIMULATII A 1.768	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE 0.70 – 15.20 0.70 – 15.20 0.70 – 11.00 0.70 – 8.40 0.70 – 7.00 0.60 – 5.50 0.50 – 5.50 0.27 – 5.50	) 0.35 - 5.50 (µlU/mL)
THYROID STIMULAT by CMIA (CHEMILUMIN 3rd GENERATION, ULT	ING HORMONE (TSH): SERUN IESCENT MICROPARTICLE IMMUNO RASENSITIVE AGE 0 – 5 DAYS 6 Days – 2 Months 3 – 11 Months 1 – 5 Years 6 – 10 Years 11 - 15	ENDOCRI YROID STIMULATII A 1.768 DASSAY)	NOLOGY NG HORMONE (TSH) μIU/mL REFFERENCE RANGE 0.70 – 15.20 0.70 – 11.00 0.70 – 8.40 0.70 – 7.00 0.60 – 5.50 0.50 – 5.50	) 0.35 - 5.50 (µlU/mL)

1. Primary or untreated hypothyroidism, may vary from 3 times to more than 100 times normal depending on degree of hypofunction.

2.Hypothyroid patients receiving insufficient thyroid replacement therapy.

3.Hashimotos thyroiditis.

4.DRUGS: Amphetamines, Iodine containing agents and dopamine antagonist.

5. Neonatal period, increase in 1st 2-3 days of life due to post-natal surge.

DECREASED 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.



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Test Name	Value	Unit	Biological Reference interval

7.DRUGS: Glucocorticoids, Dopamine, Levodopa, T4 replacement therapy, Anti-thyroid drugs for thyrotoxicosis. 8.Pregnancy: 1st and 2nd Trimester

LIMITATIONS:

1.TSH may be normal in central hypothyroidism, recent rapid correction of hyperthyroidism or hypothyroidism, pregnancy, phenytoin therapy. 2.Autoimmune disorders may produce spurious results.



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NAME	: Mrs. NISHA			
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Test Name		Value	Unit	Biological Reference interval
		PROLACTI	N	
PROLACTIN: SERUM by cmia (chemilumi immunoassay) INTERPRETATION:	NESCENT MICROPARTICLE	129.47 <sup>H</sup>	ng/mL	3 - 25
3.Primary hypothyrol 4.Section compressio 5.Chest wall lesions a 6.Ectopic tumors. 7.DRUGS:- Anti-Dopa receptors, or serotor Opiates, High doses <b>SIGNIFICANCE:</b> 1.In loss of libido, impo from decreased musi 3. In males, prolactin 5.Clear symptoms an 4. Mild to moderately adenoma is present, <b>CAUTION:</b> Prolactin values that	on of the pituitary stalk. and renal failure. minergic drugs like antipsychotic in reuptake (anti-depressants o of estrogen or progesterone,ant lactorrhea, oligomHyperprolacti itence, infertility, and hypogona cle mass and osteoporosis. levels >13 ng/mL are indicative of n levels >27 ng/mL in the absence d signs of hyperprolactinemia ar y increased levels of serum prola 5. Whereas levels >250 ng/mL ar	c drugs, antinausea/antier f all classes, ergot derivat ticonvulsants (valporic aci nemia often results enorr dism in males. Postmenop of pregnancy and postpar- re often absent in patients actin are not a reliable gui re usually associated with y be due to macroprolacti	ives, some illegal dru d), anti-tuberculous hea or amenorrhea, so bausal and premenop tum lactation are indic with serum prolactin de for determining w a prolactin-secreting n (prolactin bound to	and infertility in premenopausal females. bausal women, as well as men, can also suffe cative of hyperprolactinemia. n levels <100 ng/mL. /hether a prolactin-producing pituitary tumor. o immunoglobulin). Macroprolactin should b





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	ANTI M	ULLERIAN H	ORMONE (AMH) GEN	I
	DRMONE (AMH) GEN II: SERUM HEMILUMINESCENCE IMMUNOASSAY)	0.733	ng/mL	0.05 - 11.00
A Correlation of FER	TILITY POTENTIAL and AMH levels ar	re :		
C	OVARIAN FERTILITY POTENTIAL			IES IN (ng/mL)
				4

OVARIAN FERTILITY POTENTIAL	AIVIE VALUES IN (ng/mL)	
 OPTIMAL FERTILITY:	4.00 – 6.80 ng/mL	
SATISFACTORY FERTILITY:	2.20 – 4.00 ng/mL	
LOW FERTILITY:	0.30 – 2.20 ng/mL	
VERY LOW/UNDETECTABLE:	0.00 – 0.30 ng/mL	
HIGH LEVEL:	>6.8 ng/mL (PCOD/GRANULOSA CELL TUMOUR)	

Anti Mullerian Hormone (AMH) is also known as Mullerian Inhibiting Substance provided by sertoli cells of the testis in males and by ovarian granulose cells in females up to antral stage in females.

#### IN MALES:

1.It is used to evaluate testicular presence and function in infants with intersex conditions or ambiguous genitalia, and to distinguish between cryptorchidism and anorchia in males

### IN FEMALES:

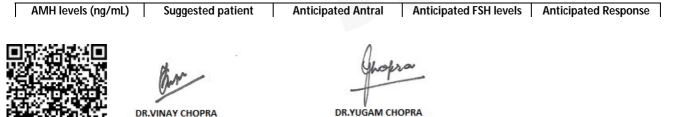
1. During reproductive age, follicular AMH productionbegins during the primary stage, peaks in preantral stage & has influence on follicular sensitivity to FSH which is impoetant in selection for follicular dominance. AMH levels thus represents the pool or number of primordial follicles but not thequality of oocytes. AMH does not vary significantly during menstrual cycle & hence can be measured independently of day of cycle. 2. Polycystic ovarian syndrome can elevate AMH 2 to 5 fold higher than age specific reference range & predict anovulatory, irregular cycles, ovarian tumours like Granulosa cell tumour are often associated with higher AMH levels.

3.Obese women are often associated with diminished ovarian reserve and can have 65% lower mean AMH levels than non-obese women. 4.In females , AMH levels do not change significantly throughout the menstrual cycle and decrease with age.

5.Assess Ovarian Reserve - correlates with the number of antral follicies in the ovaries.

6.Evaluate fertility potential and ovarian response in IVF- Women with low AMG levels are more likely to the poor ovarian responders. 7.Assess the condition of Polycystic Ovary and premature ovarian failure.

A combination of Age, Ultrasound markers-Ovarian Volume and Antral Follicle Count, AMH and FSH levels are useful for optimal assessment of ovarian reserve. Studies in various fertility clinics are ongoing to establish optimal AMH concentretaion for predicting response to invitro fertilization, however, given below is suggested interpretative reference.



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Dr. Yugam Chopra

MD (Pathology)

MD (Pathology & Microbiology) Chairman & Consultant Pathologist **CEO & Consultant Pathologist** NAME : Mrs. NISHA AGE/ GENDER : 34 YRS/FEMALE **PATIENT ID** :1622519 **COLLECTED BY** REG. NO./LAB NO. :012409230051 : **REFERRED BY** : LOOMBA HOSPITAL (AMBALA CANTT) **REGISTRATION DATE** : 23/Sep/2024 02:34 PM **BARCODE NO.** :01517560 **COLLECTION DATE** : 23/Sep/2024 02:57PM CLIENT CODE. : KOS DIAGNOSTIC LAB **REPORTING DATE** : 23/Sep/2024 04:00PM **CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT

Test Name		Value	Unit	Biological Reference interva
	Categorization for fertility based on AMH for age group (20 to 45 yrs)	Follicle counts	(day 3)	to IVF/COH cycle
Below 0.3	Very low	Below 4	Above 20	Negligible/Poor
0.3 to 2.19	Low	4 - 10	Usually 16 - 20	Reduced
2.19 t0 4.00	Satisfactory	11 - 25	Within reference range or between 11 - 15	Safe/Normal
Above 4.00	Optimal	Upto 30 and Above	Within reference range or between 11 – 15 or Above 15	Possibly Excessive

## INCREASED:

1.Polycystic ovarian syndrome (most common)

2. Ovarian Tumour: Granulosa cell tumour

#### DECREASED:

1. Anorchia, Abnormal or absence of testis in males

2.Pseudohermaphroditism

3.Post Menopause

### NOTE:

1.AMH measurement alone is seldom suffcient for diagnosis and results should be interpreted in the light of clinical finding and other relevant test such as ovarian ultrasonography(In fertility applications); abdominal or testicular ultrasound(intersex or testicular function applications); measurement of sex steroids (estradiol, Progesterone, Testosterone), FSH, Inhibin B (For fertility), and Inhibin A and B (for tumour work up). 2.Conversion of AMH grom ng/mL to pmol/L can be performed by using equation 1 ng/mL = 7.14 pmol/L





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AGE/ GENDER	: 34 YRS/FEMALE		PATIENT ID	: 1622519
COLLECTED BY	:		REG. NO./LAB NO.	: 012409230051
REFERRED BY	: LOOMBA HOSPITAL (AM	BALA CANTT)	<b>REGISTRATION DATE</b>	: 23/Sep/2024 02:34 PM
BARCODE NO.	:01517560		COLLECTION DATE	: 23/Sep/2024 02:57PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 23/Sep/2024 04:19PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROA	AD, AMBALA CANT	Т	
Test Name		Value	Unit	Biological Reference interval
		IMMUNOPATI	HOLOGY/SEROLOGY	
			HOLOGY/SEROLOGY 6 (HCV) ANTIBODY: TOTA	AL
		PATITIS C VIRUS		AL NEGATIVE: < 1.00 POSITIVE: > 1.00
	HEI DY (HCV) TOTAL: SERUM ESCENT MICROPARTICLE IMMUN	PATITIS C VIRUS	<b>5 (HCV) ANTIBODY: TOT</b> S/CO	NEGATIVE: < 1.00
by CMIA (CHEMILUMIN HEPATITIS C ANTIBO RESULT	HEI DY (HCV) TOTAL: SERUM ESCENT MICROPARTICLE IMMUN DY (HCV) TOTAL	DATITIS C VIRUS 0.12 NON - R	<b>5 (HCV) ANTIBODY: TOT</b> S/CO	NEGATIVE: < 1.00
by CMIA (CHEMILUMIN HEPATITIS C ANTIBO RESULT by CMIA (CHEMILUMIN	HEI DY (HCV) TOTAL: SERUM ESCENT MICROPARTICLE IMMUN	DATITIS C VIRUS 0.12 NON - R	<b>5 (HCV) ANTIBODY: TOT</b> S/CO	NEGATIVE: < 1.00
by CMIA (CHEMILUMIN HEPATITIS C ANTIBO RESULT by CMIA (CHEMILUMIN INTERPRETATION:-	HEI DY (HCV) TOTAL: SERUM ESCENT MICROPARTICLE IMMUN DY (HCV) TOTAL	DATITIS C VIRUS 0.12 NON - R	<b>5 (HCV) ANTIBODY: TOT</b> S/CO	NEGATIVE: < 1.00
by CMIA (CHEMILUMIN HEPATITIS C ANTIBO RESULT by CMIA (CHEMILUMIN INTERPRETATION:-	HEI DY (HCV) TOTAL: SERUM escent microparticle immun DY (HCV) TOTAL escent microparticle immun	PATITIS C VIRUS 0.12 NON - R IOASSAY)	<b>S (HCV) ANTIBODY: TOT</b> S/CO EACTIVE	NEGATIVE: < 1.00 POSITIVE: > 1.00

2. Routine screening of low and high prevelance population including blood donors.

NOTE:

1. False positive results are seen in Auto-immune disease, Rheumatoid Factor, HYpergammaglobulinemia, Paraproteinemia, Passive antibody transfer, Anti-idiotypes and Anti-superoxide dismutase.

2. False negative results are seen in early Acute infection, Immunosuppression and Immuno-incompetence.

3. HCV-RNĂ PCR recommended in all reactive results to differentiate between past and present infection.





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







	Dr. Vinay Ch MD (Pathology & Chairman & Cor			(Pathology)
NAME	: Mrs. NISHA			
AGE/ GENDER	: 34 YRS/FEMALE		PATIENT ID	: 1622519
COLLECTED BY	:		REG. NO./LAB NO.	: 012409230051
<b>REFERRED BY</b>	: LOOMBA HOSPITAL (AMBA	LA CANTT)	<b>REGISTRATION DATE</b>	: 23/Sep/2024 02:34 PM
BARCODE NO.	:01517560		COLLECTION DATE	: 23/Sep/2024 02:57PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 23/Sep/2024 05:05PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANT		
CLIENT ADDRESS			-	
Test Name		Value	Unit	Biological Reference interval
Test Name		Value	Unit	Biological Reference interval (P-24 ANTIGEN DETECTION)
Test Name ANT HIV 1/2 AND P24 AN	I HUMAN IMMUNODEFICI	Value ENCY VIRUS (H 0.13	Unit	
Test Name ANT HIV 1/2 AND P24 AN by CMIA (CHEMILUMIN HIV 1/2 AND P24 AN	I HUMAN IMMUNODEFICII ITIGEN: SERUM IESCENT MICROPARTICLE IMMUNOA	Value ENCY VIRUS (F 0.13 ISSAY) NON - RE	Unit HIV) DUO ULTRA WITH S/CO	(P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00
Test Name ANT HIV 1/2 AND P24 AN by CMIA (CHEMILUMIN HIV 1/2 AND P24 AN by CMIA (CHEMILUMIN INTERPRETATION:-	I HUMAN IMMUNODEFICII ITIGEN: SERUM IESCENT MICROPARTICLE IMMUNOA	Value ENCY VIRUS (F 0.13 ISSAY) NON - RE	Unit HIV) DUO ULTRA WITH S/CO	(P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00
Test Name ANT HIV 1/2 AND P24 AN by CMIA (CHEMILUMIN HIV 1/2 AND P24 AN by CMIA (CHEMILUMIN INTERPRETATION:- RESUL	I HUMAN IMMUNODEFICI ITIGEN: SERUM IESCENT MICROPARTICLE IMMUNOA ITIGEN RESULT IESCENT MICROPARTICLE IMMUNOA	Value ENCY VIRUS (F 0.13 ISSAY) NON - RE	Unit HIV) DUO ULTRA WITH S/CO	(P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00

exposed to HIV 1/2 infection or the sample has been tested during the "window phase" i.e. before the development of detectable levels of antibodies. Hence a Non Reactive result does not exclude the possibility of exposure or infection with HIV 1/2. **RECOMMENDATIONS:** 

Results to be clinically correlated
 Rarely falsenegativity/positivity may occur.

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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.





		hopra & Microbiology) onsultant Patholo		(Pathology)		
NAME	: Mrs. NISHA					
AGE/ GENDER	: 34 YRS/FEMALE		PATIENT ID	: 1622519		
COLLECTED BY	:		REG. NO./LAB NO.	: 012409230051		
REFERRED BY	: LOOMBA HOSPITAL (AMB	ALA CANTT)	<b>REGISTRATION DATE</b>	: 23/Sep/2024 02:34 PM		
BARCODE NO.	: 01517560		COLLECTION DATE	: 23/Sep/2024 02:57PM		
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 23/Sep/2024 04:00PM		
CLIENT ADDRESS	S : 6349/1, NICHOLSON ROAD, AMBALA CANTT					
Test Name		Value	Unit	Biological Reference interval		
	HEPAT	TITIS B SURFA	CE ANTIGEN (HBsAg) UL	TRA		
SERUM	CE ANTIGEN (HBsAg):	0.03	<b>CE ANTIGEN (HBsAg) UL</b> S/CO	TRA NEGATIVE: < 1.0 POSITIVE: > 1.0		
SERUM by CMIA (CHEMILUMII HEPATITIS B SURFA( RESULT	CE ANTIGEN (HBsAg): NESCENT MICROPARTICLE IMMUNC CE ANTIGEN (HBSAg)	0.03 DASSAY) NON RE		NEGATIVE: < 1.0		
SERUM by CMIA (CHEMILUMII HEPATITIS B SURFA( RESULT by CMIA (CHEMILUMII	CE ANTIGEN (HBsAg):	0.03 DASSAY) NON RE	s/co	NEGATIVE: < 1.0		
SERUM by CMIA (CHEMILUMII HEPATITIS B SURFA) RESULT by CMIA (CHEMILUMII INTERPRETATION:	CE ANTIGEN (HBsAg): NESCENT MICROPARTICLE IMMUNC CE ANTIGEN (HBSAg)	0.03 DASSAY) NON RE	s/co	NEGATIVE: < 1.0		
SERUM by CMIA (CHEMILUMII HEPATITIS B SURFA) RESULT by CMIA (CHEMILUMII INTERPRETATION: RESU	CE ANTIGEN (HBsAg): NESCENT MICROPARTICLE IMMUNC CE ANTIGEN (HBSAg) NESCENT MICROPARTICLE IMMUNC	0.03 DASSAY) NON RE	S/CO	NEGATIVE: < 1.0		

Hepatitis B Virus (HBV) is a member of the Hepadna virus family causing infection of the liver with extremely variable clinical features. Hepatitis B is transmitted primarily by body fluids especially serum and also spread effectively sexually and from mother to baby. In most individuals HBV hepatitis is self limiting, but 1-2 % normal adolescent and adults develop Chronic Hepatitis. Frequency of chronic HBV infection is 5-10% in immunocompromised patients and 80 % neonates. The initial serological marker of acute infection is HBsAg which typically appears 2-3 months after infection and disappears 12-20 weeks after onset of symtoms. Persistence of HBsAg for more than 6 months indicates carrier state or Chronic Liver disease.





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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT





	MD (Patholog	<b>Dr. Vinay Chopra</b> MD (Pathology & Microbiology) Chairman & Consultant Pathologist		n <b>Chopra</b> (Pathology) t Pathologist
NAME	: Mrs. NISHA			
AGE/ GENDER	: 34 YRS/FEMALE	P	ATIENT ID	: 1622519
COLLECTED BY	:	R	EG. NO./LAB NO.	: 012409230051
<b>REFERRED BY</b>	: LOOMBA HOSPITAL (AME	BALA CANTT) <b>R</b>	EGISTRATION DATE	: 23/Sep/2024 02:34 PM
BARCODE NO.	: 01517560	C	<b>DLLECTION DATE</b>	: 23/Sep/2024 02:57PM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	R	EPORTING DATE	: 23/Sep/2024 03:14PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROA	D, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
		VD	RL	
VDRL		NON REACT	IVE	NON REACTIVE
by IMMUNOCHROMAT INTERPRETATION:	<sup>C</sup> OGRAPHY			
1.Does not become p	oositive until 7 - 10 days after a	appearance ofchancre.		
2. High titer (>1:16) -	active disease. iological falsepositive test in 90	1% cases or due to late	or late latent synhillis	
4. Treatment of prima	ary syphillis causes progressive	e decline tonegative VI	ORL within 2 years.	
5.Rising titer (4X) ind	licates relapse, reinfection, or t e in early primary, late latent,	reatment failure and n	eed for retreatment.	
7.Reactive and weak	ly reactive tests should always	be confirmed with FTA	ABS (fluorescent trepon	emal antibody absorptiontest).
	<b>OSITIVE TEST RESULTS (&lt;6 MON</b> s (e.g., hepatitis, measles, infe			
2.M. pneumoniae; Cl	hlamydia; Malaria infection.			
3.Some immunization 4.Pregnancy (rare)	ns			
		_		
LONGTERM FALSE PO	SITIVE TEST RESULTS (>6 MON	THS DURATION) MAY O	CCUR IN:	

- LONGTERM FALSE POSITIVE TEST RESULTS (>6 MONTHS DURATION) MAY OCCUR IN: 1.Serious underlying disease e.g., collagen vascular diseases, leprosy ,malignancy.
- 2.Intravenous drug users.
- 3. Rheumatoid arthritis, thyroiditis, AIDS, Sjogren's syndrome.
- 4.<10 % of patients older thanage 70 years.
- 5.Patients taking some anti-hypertensive drugs.

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





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