



	<b>Dr. Vinay Ch</b> MD (Pathology & Chairman & Cor		Dr. Yugam MD CEO & Consultant	(Pathology)
NAME	: Mrs. PARVISHA			
AGE/ GENDER	: 29 YRS/FEMALE	РАТ	IENT ID	: 1801765
COLLECTED BY	:	REG	. NO./LAB NO.	: 012503220030
REFERRED BY	: LOOMBA HOSPITAL (AMBA	LA CANTT) <b>REG</b>	ISTRATION DATE	: 22/Mar/2025 11:44 AM
BARCODE NO.	: 01527549	COL	LECTION DATE	: 22/Mar/2025 11:57AM
CLIENT CODE.	: KOS DIAGNOSTIC LAB	REP	ORTING DATE	: 22/Mar/2025 12:02PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
<ol> <li>2) Nutritional deficie</li> <li>3) Bone marrow prob</li> <li>4) Suppression by red</li> <li>5) Kidney failure</li> <li>6) Abnormal hemogle</li> <li>POLYCYTHEMIA (INCF</li> <li>1) People in higher a</li> <li>2) Smoking (Secondal</li> <li>3) Dehydration produ</li> </ol>	Imatic injury, surgery, bleeding, ncy (iron, vitamin B12, folate) Iems (replacement of bone marr d blood cell synthesis by chemot obin structure (sickle cell anemi REASED HAEMOGLOBIN): Ititudes (Physiological) ry Polycythemia) uces a falsely rise in hemoglobin	row by cancer) therapy drugs a or thalassemia).		
5) Certain tumors 6) A disorder of the b 7) Abuse of the drug chemically raising th	ease (for example, emphysema) one marrow known as polycythe erythropoetin (Epogen) by athle e production of red blood cells) <b>FED ON EDTA WHOLE BLOOD</b>	tes for blood doping pu	rposes (increasing the	e amount of oxygen available to the body by

## NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD





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







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CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 22/Mar/2025 02:42PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	MBALA CANTT	2	
Test Name		Value	Unit	<b>Biological Reference interval</b>
		BLEEDIN	IG TIME (BT)	
BLEEDING TIME (E	T)	2 MIN. 10	D SEC. MINS	1 - 5



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CLIENT CODE.	: KOS DIAGNOSTIC LAB	<b>REPORTING DATE</b>	: 22/Mar/2025 02:43PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMBALA CAN	ITT	
Test Name	Value	Unit	<b>Biological Reference interval</b>
	CLOTT	TING TIME (CT)	
CLOTTING TIME (C by Capillary tube N		. 33 SEC. MINS	4 - 9



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CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 22/Mar/2025 01:14PM
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD,	AMBALA CANT	Т	
Test Name		Value	Unit	Biological Reference interva
	IMA	ЛІІЛОРАТН		v
			IOLOGY/SEROLOGY (HCV) ANTIBODY: TO	
		<b>TIS C VIRUS</b> 0.11		
by CMIA (CHEMILUMIN HEPATITIS C ANTI RESULT by CMIA (CHEMILUMIN	<b>HEPATI</b> BODY (HCV) TOTAL: SERUM	TIS C VIRUS 0.11 ISSAY) NON - R	(HCV) ANTIBODY: TO	<b>DTAL</b> NEGATIVE: < 1.00
by CMIA (CHEMILUMIN HEPATITIS C ANTI RESULT by CMIA (CHEMILUMIN INTERPRETATION:-	HEPATI BODY (HCV) TOTAL: SERUM NESCENT MICROPARTICLE IMMUNOA BODY (HCV) TOTAL	TIS C VIRUS 0.11 ISSAY) NON - R	<b>(HCV) ANTIBODY: TO</b> S/CO EACTIVE	<b>DTAL</b> NEGATIVE: < 1.00
by CMIA (CHEMILUMIN HEPATITIS C ANTI RESULT by CMIA (CHEMILUMIN INTERPRETATION:-	HEPATI BODY (HCV) TOTAL: SERUM VESCENT MICROPARTICLE IMMUNOA BODY (HCV) TOTAL	TIS C VIRUS 0.11 ISSAY) NON - R	(HCV) ANTIBODY: TO S/CO	DTAL NEGATIVE: < 1.00 POSITIVE: > 1.00

1. Indicator of past or present infection, but does not differentiate between Acute/ Chronic/Resolved Infection. 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-RNA PCR recommended in all reactive results to differentiate between past and present infection.





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CLIENT ADDRESS	: 6349/1, NICHOLSO	N ROAD, AMBALA CANTT		
Test Name		Value	Unit	Biological Reference interval
	MAN IMMUNODEF			Biological Reference interval I (P-24 ANTIGEN DETECTION)
ANTI HUI HIV 1/2 AND P24 /		ICIENCY VIRUS (HI 0.13		
ANTI HUI HIV 1/2 AND P24 <i>I</i> by CMIA (CHEMILUMIN HIV 1/2 AND P24 <i>I</i> by CMIA (CHEMILUMIN	ANTIGEN: SERUM	ICIENCY VIRUS (HI 0.13 IMMUNOASSAY) NON - RE	<b>V) DUO ULTRA WITH</b> S/CO	I (P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00
ANTI HUI HIV 1/2 AND P24 <i>I</i> by CMIA (CHEMILUMIN HIV 1/2 AND P24 <i>I</i> by CMIA (CHEMILUMIN INTERPRETATION:-	ANTIGEN: SERUM IESCENT MICROPARTICLE ANTIGEN RESULT IESCENT MICROPARTICLE	ICIENCY VIRUS (HI 0.13 IMMUNOASSAY) NON - RE	<b>V) DUO ULTRA WITH</b> S/CO ACTIVE	I (P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00
HIV 1/2 AND P24 <i>I</i> by CMIA (CHEMILUMIN HIV 1/2 AND P24 <i>I</i> by CMIA (CHEMILUMIN <u>INTERPRETATION:-</u> RESU	ANTIGEN: SERUM iescent microparticle ANTIGEN RESULT	ICIENCY VIRUS (HI 0.13 IMMUNOASSAY) NON - RE	<b>V) DUO ULTRA WITH</b> S/CO	I (P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00

Non-Reactive result implies that antibodies to HIV 1/2 have not been detected in the sample. This menas that patient has either not been 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:** 1. Results to be clinically correlated

2. Rarely falsenegativity/positivity may occur.





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	Value	Unit	Biological Reference interval
HEPATITIS	B SURFAC	CE ANTIGEN (HBsAg	) ULTRA
5	0.31 say)	S/CO	NEGATIVE: < 1.0 POSITIVE: > 1.0
ç		EACTIVE	
SOLINI MICHOLANNOLO			
		REMARKS	
		NEGATIVE (-ve	
	Chairman & Const : Mrs. PARVISHA : 29 YRS/FEMALE : : LOOMBA HOSPITAL (AMBALA : 01527549 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, A HEPATITIS ACE ANTIGEN (HBSAg): ESCENT MICROPARTICLE IMMUNOASS ACE ANTIGEN (HBSAg)	Chairman & Consultant Patholo : Mrs. PARVISHA : 29 YRS/FEMALE : : LOOMBA HOSPITAL (AMBALA CANTT) : 01527549 : KOS DIAGNOSTIC LAB : 6349/1, NICHOLSON ROAD, AMBALA CANT Value Value KEPATITIS B SURFAC ACE ANTIGEN (HBsAg): 0.31 ESCENT MICROPARTICLE IMMUNOASSAY) ACE ANTIGEN (HBsAg) NON RESERVED THIS AND	CEO & Consult         : Mrs. PARVISHA         : 29 YRS/FEMALE       PATIENT ID         : OPACINAL (AMBALA CANTT)       REG. NO./LAB NO.         : LOOMBA HOSPITAL (AMBALA CANTT)       REGISTRATION DATT         : 01527549       COLLECTION DATE         : KOS DIAGNOSTIC LAB       REPORTING DATE         : 6349/1, NICHOLSON ROAD, AMBALA CANTT       Init         MEPATITIS B SURFACE ANTIGEN (HBsAg         ACE ANTIGEN (HBSAg):       0.31         SCEENT MICROPARTICLE IMMUNOASSAY)       NON REACTIVE         SCEENT MICROPARTICLE IMMUNOASSAY)       NON REACTIVE         SO       NEGATIVE (-v

Hepatitis B virus (HBV) is a member of the Hepatina 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 Name		Value	Unit	<b>Biological Reference interval</b>
VDRL by IMMUNOCHROMA	TOGRAPHY	NON REA	CTIVE	NON REACTIVE
by IMMUNOCHROMAT INTERPRETATION: 1.Does not become   2.High titer (>1:16) - 3.Low titer (<1:8) - b 4.Treatment of prim 5.Rising titer (4X) inc 6.May benonreactiv 7.Reactive and weak SHORTTERM FALSE P 1.Acute viral illnesse 2.M. pneumoniae; C 3.Some immunizatio	positive until 7 - 10 days after app active disease. iological falsepositive test in 90% ary syphillis causes progressive d licates relapse, reinfection, or trea e in early primary, late latent, an ly reactive tests should always be OSITIVE TEST RESULTS (<6 MONTH es (e.g., hepatitis, measles, infection hlamydia; Malaria infection.	bearance ofchancr cases or due to lat ecline tonegative tment failure and d late syphillis (ap confirmedwith FT S DURATION) MAY	e. Te or late latent syphillis. VDRL within 2 years. need for retreatment. prox. 25% ofcases). A-ABS (fluorescent trepone OCCURIN:	
by IMMUNOCHROMAT INTERPRETATION: 1.Does not become [ 2.High titer (>1:16) - 3.Low titer (<1:8) - b 4.Treatment of prim 5.Rising titer (4X) inc 6.May benonreactiv 7.Reactive and weak SHORTTERM FALSE P 1.Acute viral illnesse 2.M. pneumoniae; C 3.Some immunizatio 4.Pregnancy (rare) LONGTERM FALSE PC 1.Serious underlying 2.Intravenous drug u 3.Rheumatoid arthri 4.<10 % of patients o	positive until 7 - 10 days after app active disease. iological falsepositive test in 90% ary syphillis causes progressive d licates relapse, reinfection, or trea e in early primary, late latent, an- ly reactive tests should always be OSITIVE TEST RESULTS (<6 MONTH hlamydia; Malaria infection. ns	bearance ofchancr cases or due to late ecline tonegative itment failure and d late syphillis (ap confirmedwith FT IS DURATION) MAY ous mononucleos S DURATION) MAY iseases, leprosy , l	e. <b>The or late latent syphillis.</b> VDRL within 2 years. need for retreatment. prox. 25% ofcases). <b>A-ABS (fluorescent trepone</b> <b>Y OCCURIN:</b> is) OCCUR IN:	





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