



<b>5. PARVEEN WALIA</b> /RS/FEMALE /JESH		PATIENT ID			
		PATIENT ID			
UESH			: 1685584		
		REG. NO./LAB NO.	: 012411290026		
		<b>REGISTRATION DATE</b>	: 29/Nov/2024 10:29 AM		
21669		COLLECTION DATE	: 29/Nov/2024 10:52AM		
S DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 29/Nov/2024 12:14PM		
: 6349/1, NICHOLSON ROAD, AMBALA CANTT					
	Value	Unit	Biological Reference interval		
(HCV) TOTAL: SERUM	0.13	S/CO	NEGATIVE: < 1.00 POSITIVE: > 1.00		
		ACTIVE			
		DEWVDK2			
<b>RESULT (INDEX)</b> < 1.00		NON - REACTIVE/NOT - DETECTED			
		YMPTOMATIC/INFECTIVE STATE/CARRIER STATE.			
	19/1, NICHOLSON ROAD, VIRA HEPATT (HCV) TOTAL: SERUM MICROPARTICLE IMMUNOA (HCV) TOTAL T MICROPARTICLE IMMUNOA (HCV) TOTAL T MICROPARTICLE IMMUNOA (INDEX) 00 00 virus of Favivirus group t	19/1, NICHOLSON ROAD, AMBALA CANTT  Value  VIRAL MARKERS  HEPATITIS C VIRUS (1 (HCV) TOTAL: SERUM 0.13 rmicroparticle immunoassay) (HCV) TOTAL NON - RE  rmicroparticle immunoassay)  NON - RE  MICROPARTICLE immunoassay	19/1, NICHOLSON ROAD, AMBALA CANTT          Value       Unit         VIRAL MARKERS COMBO PANEL: 2         HEPATITIS C VIRUS (HCV) ANTIBODY: TO         (HCV) TOTAL: SERUM       0.13       S/CO         (HCV) TOTAL: SERUM       0.13       S/CO         (HCV) TOTAL: SERUM       0.13       S/CO         IMICROPARTICLE IMMUNOASSAY)         (HCV) TOTAL       NON - REACTIVE         INDEX)       REMARKS         OO       NON - REACTIVE/NOT - DET		

Indicator of past or present infection, but does not differentiate between Acute/ Chronic/Resolved Infection.
 Routine screening of low and high prevelance population including blood donors.

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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.





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







	Dr. Vinay Ch MD (Pathology & Chairman & Cons	Microbiology)	Dr. Yugam Chopra MD (Pathology) t CEO & Consultant Pathologist		
NAME	: Mrs. PARVEEN WALIA				
AGE/ GENDER	: 57 YRS/FEMALE		PATIENT ID	: 1685584	
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REFERRED BY	:		<b>REGISTRATION DATE</b>	: 29/Nov/2024 10:29 AM	
BARCODE NO.	: 01521669		COLLECTION DATE	: 29/Nov/2024 10:52AM	
CLIENT CODE.	: KOS DIAGNOSTIC LAB		<b>REPORTING DATE</b>	: 29/Nov/2024 12:14PM	
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	AMBALA CANTT			
Test Name		Value	Unit	Biological Reference interval	
	MAN IMMUNODEFICIENC			Biological Reference interval I (P-24 ANTIGEN DETECTION)	
ANTI HUN HIV 1/2 AND P24 /		Y VIRUS (HIV			
ANTI HUI HIV 1/2 AND P24 A by CMIA (CHEMILUMIN HIV 1/2 AND P24 A by CMIA (CHEMILUMIN	ANTIGEN: SERUM	Y VIRUS (HIV 0.08 ssay) NON - RE	<b>) DUO ULTRA WITH</b> S/CO	I (P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00	
ANTI HUN HIV 1/2 AND P24 A by CMIA (CHEMILUMIN HIV 1/2 AND P24 A by CMIA (CHEMILUMIN INTERPRETATION:-	ANTIGEN: SERUM iescent microparticle immunoas ANTIGEN RESULT	Y VIRUS (HIV 0.08 ssay) NON - RE	<b>) DUO ULTRA WITH</b> S/CO	I (P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00	
ANTI HUN HIV 1/2 AND P24 A by CMIA (CHEMILUMIN HIV 1/2 AND P24 A by CMIA (CHEMILUMIN INTERPRETATION:- RESUL	ANTIGEN: SERUM IESCENT MICROPARTICLE IMMUNOAS ANTIGEN RESULT IESCENT MICROPARTICLE IMMUNOAS	Y VIRUS (HIV 0.08 ssay) NON - RE	7) DUO ULTRA WITH S/CO ACTIVE	I (P-24 ANTIGEN DETECTION) NEGATIVE: < 1.00	

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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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REFERRED BY	:				
BARCODE NO.	:01521669				
CLIENT CODE.	: KOS DIAGNOSTIC LAB				
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, A	MBALA CANTT			
		Value	Unit	Biological Reference interval	
Test Name		value	UIIIt	biological kelerence interval	
Test Name	HEPATITIS	B SURFACE ANTIGEN (H			
HEPATITIS B SURI SERUM	FACE ANTIGEN (HBsAg):	<b>B SURFACE ANTIGEN (H</b> 0.01			
HEPATITIS B SURF SERUM by CMIA (CHEMILUMIN HEPATITIS B SURF RESULT		<b>B SURFACE ANTIGEN (H</b> 0.01 SAY) NON REACTIVE	(BsAg) (	J <b>LTRA</b> NEGATIVE: < 1.0	
HEPATITIS B SURF SERUM by CMIA (CHEMILUMII HEPATITIS B SURF RESULT by CMIA (CHEMILUMII INTERPRETATION:	FACE ANTIGEN (HBsAg): IESCENT MICROPARTICLE IMMUNOAS FACE ANTIGEN (HBsAg) IESCENT MICROPARTICLE IMMUNOAS	S B SURFACE ANTIGEN (H 0.01 SAY) NON REACTIVE SAY)	( <b>BsAg) (</b> s/co	J <b>LTRA</b> NEGATIVE: < 1.0	
HEPATITIS B SURF SERUM by CMIA (CHEMILUMII HEPATITIS B SURF RESULT by CMIA (CHEMILUMII INTERPRETATION: RESUL	FACE ANTIGEN (HBsAg): IESCENT MICROPARTICLE IMMUNOAS: FACE ANTIGEN (HBsAg)	S B SURFACE ANTIGEN (H 0.01 SAY) NON REACTIVE SAY) REMA	( <b>BsAg) (</b> s/co	J <b>LTRA</b> NEGATIVE: < 1.0	

KOS Diagnostic Lab (A Unit of KOS Healthcare)

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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BIQS 150 9001 : 2008 CERT	KOS Diagno (A Unit of KOS		CELLENCE IN HEALTHCARE				
	<b>Dr. Vinay Chop</b> MD (Pathology & Mic Chairman & Consulta	crobiology)	Dr. Yugam MD EO & Consultant	(Pathology)			
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BARCODE NO.	:01521669		TON DATE	: 29/Nov/2024 10:52AM			
CLIENT CODE.	: KOS DIAGNOSTIC LAB		ING DATE	: 29/Nov/2024 11:30AM			
CLIENT ADDRESS	: 6349/1, NICHOLSON ROAD, AMI	BALA CANTT					
Test Name		Value	Unit	<b>Biological Reference interval</b>	1   		
		VDBI					
VDRL       NON REACTIVE       NON REACTIVE         by IMMUNOCHROMATOGRAPHY       NON REACTIVE       NON REACTIVE         by IMMUNOCHROMATOGRAPHY       NUTERPRETATIONE       NON REACTIVE         1. Does not become positive until 7 - 10 days after appearance ofchancre.       2.High itter (-1:6) - active disease.       3.Low titer (-1:8) - biological falsepositive test in 90% cases or due to late or late latent syphills.         3. Low titer (-1:8) - biological falsepositive test in 90% cases or due to late or late latent syphills.       4.Treatment of primary syphills causes progressive decline tonegative VDRL within 2 years.         5. Rising titer (4X) indicates relapse, reinfection, or treatment failure and need for retreatment.       6.May benonreactive in early primary, late latent, and late syphillis (approx. 25% of cases).         7.Reactive and weakly reactive tests should always be confirmedwith FTA-ABS (fluorescent treponemal antibody absorptiontest).         SHORTERM FALSE POSITIVE TEST RESULTS (<6 MONTHS DURATION) MAY OCCURINE							
-	der thanage 70 years. ne anti-hypertensive drugs.						
	* * *	End Of Report **	< *				
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TEST PERFORMED AT KOS DIAGNOSTIC LAB, AMBALA CANTT.