

LABORATORY TEST REPORT

Name	: Mrs. V RAJESHWARI		
Sample ID	: B2623270, B2623265		
Age/Gender	: 66 Years/Female	Reg. No	: 0312505060042
Referred by	: Dr. M KAUSHIK REDDY	SPP Code	: SPL-CV-172
Referring Customer	: V CARE MEDICAL DIAGNOSTICS	Collected On	: 06-May-2025 01:46 PM
Primary Sample	: Whole Blood	Received On	: 06-May-2025 03:39 PM
Sample Tested In	: Capillary Tube, Citrated Plasm	Reported On	: 06-May-2025 08:02 PM
Client Address	: Kimtee colony ,Gokul Nagar,Tarnaka	Report Status	: Final Report

HAEMATOLOGY

Test Name	Results	Units	Biological Reference Interval
Bleeding Time & Clotting Time			
Bleeding Time (BT) <small>(Method: Capillary Method)</small>	03:00 sec	Minutes	2 - 5
Clotting Time (CT) <small>(Method: Capillary Method)</small>	05:10 sec	Minutes	3 - 7
<u>PROTHROMBIN TIME (P TIME)</u>			
PT-Patient Value <small>(Method: Photo Optical Clot Detection)</small>	14.4	Secs	10-15
PT-Mean Control Value	13.00	Seconds	
PT Ratio	1.11		
PT INR	1.20		0.9-1.2

Interpretation :

Prothrombin time measures the extrinsic coagulation pathway which consists of activated Factor VII (VIIa), Tissue factor and Proteins of the common pathway (Factors X, V, II & Fibrinogen). This assay is used to control long term oral anticoagulant therapy, evaluation of liver function & to evaluate coagulation disorders specially factors involved in the extrinsic pathway like Factors V, VII, X, Prothrombin & Fibrinogen.

Note

1. INR is the parameter of choice in monitoring adequacy of oral anticoagulant therapy. Appropriate therapeutic range varies with the disease and treatment intensity
2. Prolonged INR suggests potential bleeding disorder / bleeding complications
3. Results should be clinically correlated
4. Test conducted on Citrated plasma

*** End Of Report ***



Page 1 of 7
Swarnabala - M
DR.SWARNA BALA
MD PATHOLOGY

LABORATORY TEST REPORT

Name	: Mrs. V RAJESHWARI		
Sample ID	: B2623263		
Age/Gender	: 66 Years/Female	Reg. No	: 0312505060042
Referred by	: Dr. M KAUSHIK REDDY	SPP Code	: SPL-CV-172
Referring Customer	: V CARE MEDICAL DIAGNOSTICS	Collected On	: 06-May-2025 01:46 PM
Primary Sample	: Whole Blood	Received On	: 06-May-2025 03:39 PM
Sample Tested In	: Whole Blood EDTA	Reported On	: 06-May-2025 04:31 PM
Client Address	: Kimtee colony , Gokul Nagar, Tarnaka	Report Status	: Final Report

HAEMATOLOGY

SURGICAL PROFILE-II

Test Name	Results	Units	Biological Reference Interval
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Complete Blood Picture(CBP)

 Haemoglobin (Hb) <small>(Method: Cynmeth Method)</small>	12.2	g/dL	12-15
 Haematocrit (HCT) <small>(Method: Calculated)</small>	36.7	%	40-50
 RBC Count <small>(Method: Cell Impedance)</small>	4.12	10 ¹² /L	3.8-4.8
 MCV <small>(Method: Calculated)</small>	89	fl	81-101
 MCH <small>(Method: Calculated)</small>	29.6	pg	27-32
 MCHC <small>(Method: Calculated)</small>	33.2	g/dL	32.5-34.5
 RDW-CV <small>(Method: Calculated)</small>	13.3	%	11.6-14.0
 Platelet Count (PLT) <small>(Method: Cell Impedance)</small>	305	10 ⁹ /L	150-410
 Total WBC Count <small>(Method: Impedance)</small>	7.2	10 ⁹ /L	4.0-10.0

Differential Leucocyte Count (DC)

 Neutrophils <small>(Method: Cell Impedance)</small>	70	%	40-70
 Lymphocytes <small>(Method: Cell Impedance)</small>	20	%	20-40
 Monocytes <small>(Method: Microscopy)</small>	06	%	2-10
 Eosinophils <small>(Method: Microscopy)</small>	04	%	1-6
 Basophils <small>(Method: Microscopy)</small>	00	%	1-2
 Absolute Neutrophils Count <small>(Method: Impedance)</small>	5.04	10 ⁹ /L	2.0-7.0
 Absolute Lymphocyte Count <small>(Method: Impedance)</small>	1.44	10 ⁹ /L	1.0-3.0
 Absolute Monocyte Count <small>(Method: Calculated)</small>	0.43	10 ⁹ /L	0.2-1.0
 Absolute Eosinophils Count <small>(Method: Calculated)</small>	0.29	10 ⁹ /L	0.02-0.5
 Absolute Basophil ICount <small>(Method: Calculated)</small>	0.00	10 ⁹ /L	0.0-0.3

Morphology Normocytic normochromic blood picture.

Blood Grouping (A B O) O

Rh Typing Positive



LABORATORY TEST REPORT

Name	: Mrs. V RAJESHWARI		
Sample ID	: B2623267, B2623268		
Age/Gender	: 66 Years/Female	Reg. No	: 0312505060042
Referred by	: Dr. M KAUSHIK REDDY	SPP Code	: SPL-CV-172
Referring Customer	: V CARE MEDICAL DIAGNOSTICS	Collected On	: 06-May-2025 01:46 PM
Primary Sample	: Whole Blood	Received On	: 06-May-2025 03:39 PM
Sample Tested In	: Plasma-NaF(R), Serum	Reported On	: 06-May-2025 05:59 PM
Client Address	: Kimtee colony ,Gokul Nagar, Tarnaka	Report Status	: Final Report

CLINICAL BIOCHEMISTRY

SURGICAL PROFILE-II

Test Name	Results	Units	Biological Reference Interval
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Glucose Random (RBS) . 101 mg/dL 70-140
(Method: Hexokinase (HK))

Interpretation of Plasma Glucose based on ADA guidelines 2024

Diagnosis	Fasting Plasma Glucose(mg/dL)	2hrs Plasma Glucose(mg/dL)	HbA1c(%)	RBS(mg/dL)
Prediabetes	100-125	140-199	5.7-6.4	NA
Diabetes	> = 126	> = 200	> = 6.5	>=200(with symptoms)

Reference: Diabetes care 2024 Jan (1:47 (suppl.1):S20- S42.

- The random blood glucose if it is above 200 mg/dL and the patient has increased thirst, polyuria, and polyphagia, suggests diabetes mellitus.
- As a rule, two-hour glucose samples will reach the fasting level or it will be in the normal range.

 Creatinine <small>(Method: Sarcosine Oxidase Method)</small>	0.64	mg/dL	0.55-1.02
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Interpretation:

- This test is done to see how well your kidneys are working. Creatinine is a chemical waste product of creatine. Creatine is a chemical made by the body and is used to supply energy mainly to muscles.
- **A higher than normal level may be due to:**
- Renal diseases and insufficiency with decreased glomerular filtration, urinary tract obstruction, reduced renal blood flow including congestive heart failure, shock, and dehydration; rhabdomyolysis can cause elevated serum creatinine.
- **A lower than normal level may be due to:**
- Small stature, debilitation, decreased muscle mass; some complex cases of severe hepatic disease can cause low serum creatinine levels. In advanced liver disease, low creatinine may result from decreased hepatic production of creatinine and inadequate dietary protein as well as reduced muscle mass.

 Urea-Serum <small>(Method: Urease-GLDH, UV Method)</small>	19.7	mg/dL	17.1-49.2
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Interpretation:

- Catabolism of proteins and amino acids results in the formation of urea, which is predominantly cleared from the body by the kidneys.
- Increased urea with normal creatinine concentrations indicates a pre-renal increase in urea which may be due to a high protein diet, increased protein catabolism, reabsorption of blood proteins after GI haemorrhage, glucocorticoid treatment, dehydration or decreased perfusion of the kidneys.
- An increase in both urea and creatinine concentrations may indicate an obstructive post-renal condition such as malignancy, nephrolithiasis or prostatism.
- A low urea and increased creatinine may indicate acute tubular necrosis, low protein intake, starvation or severe liver disease.




DR. LAVANYA LAGISETTY
MD BIOCHEMISTRY

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Referring Customer	: V CARE MEDICAL DIAGNOSTICS	Collected On	: 06-May-2025 01:46 PM
Primary Sample	: Whole Blood	Received On	: 06-May-2025 03:39 PM
Sample Tested In	: Serum	Reported On	: 06-May-2025 07:36 PM
Client Address	: Kimtee colony ,Gokul Nagar,Tarnaka	Report Status	: Final Report

IMMUNOLOGY & SEROLOGY

SURGICAL PROFILE-II

Test Name	Results	Units	Biological Reference Interval
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VDRL- Syphilis Antibodies
(Method: Slide Flocculation)

Non Reactive

Non Reactive

The serological diagnosis of syphilis is classified into two groups: Nontreponemal tests (RPR/VDRL) and Treponemal tests (TPHA/CLIA). Syphilis serology is a treponemal assay for the qualitative determination of antibodies to T. pallidum in human serum or plasma as an aid in the diagnosis of syphilis. Treponemal tests may remain reactive for life, even following adequate therapy thus a positive result suggests infection with Treponema pallidum but does not distinguish between treated and untreated infections. Therefore, the results of a nontreponemal assay, such as rapid plasma reagin, are needed to provide information on a patient's disease state and history of therapy. Nontreponemal tests lack sensitivity in late stage of infection and screening with these tests alone may yield false positive reactions in various acute and chronic conditions in the absence of syphilis (biological false positive reactions).

*** End Of Report ***



Page 4 of 7


DR. RUTURAJ MANIKLAL KOLHAPURE
MD, MICROBIOLOGIST

TESTS CONDUCTED @ CENTRAL LAB, HYDERABAD

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Primary Sample	: Whole Blood	Received On	: 06-May-2025 03:39 PM
Sample Tested In	: Serum	Reported On	: 06-May-2025 07:28 PM
Client Address	: Kimtee colony , Gokul Nagar, Tarnaka	Report Status	: Final Report

IMMUNOLOGY & SEROLOGY

SURGICAL PROFILE-II

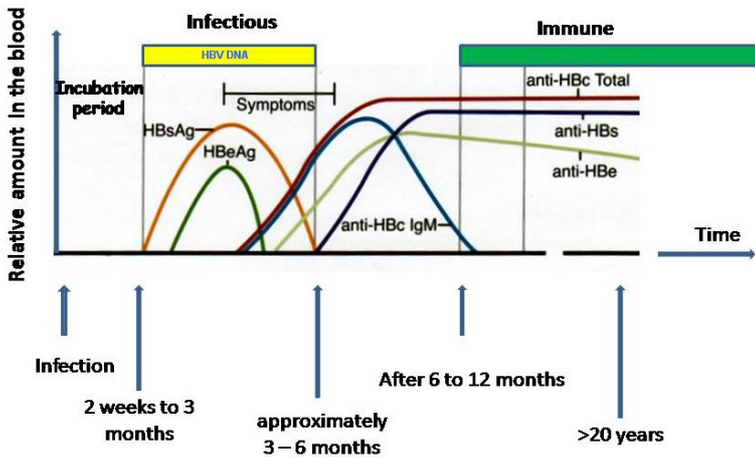
Test Name	Results	Units	Biological Reference Interval
Hepatitis B Surface Antigen (HBsAg) <small>(Method: ELISA)</small>	0.24	S/Co	<1.00 :Negative >1.00 :Positive

Interpretation:

- Negative result implies that antibodies to HBsAg have not been detected in the sample. This means the patient has either not been exposed to HBsAg 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 HBsAg.
- Positive result implies that antibodies to HBsAg have been detected in the sample.

Hepatitis B Virus (HBV) is a member of the Hepadna virus family causing infections 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 adolescents and adults develop Chronic Hepatitis. Frequency of chronic HBV infection is 5-10% in immunocompromised patients and 80% in 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 symptoms. Persistence of HBsAg for more than six months indicates development of carrier state or Chronic liver disease.

HBV antigens and antibodies in the blood



Note:

1. All Reactive results are tested additionally by Specific antibody Neutralization assay . For further confirmation Molecular assays are recommended For diagnostic purposes, results should be used in conjunction with clinical history and other hepatitis markers for Acute or Chronic infection

*** End Of Report ***



Page 5 of 7
[Signature]

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IMMUNOLOGY & SEROLOGY

SURGICAL PROFILE-II

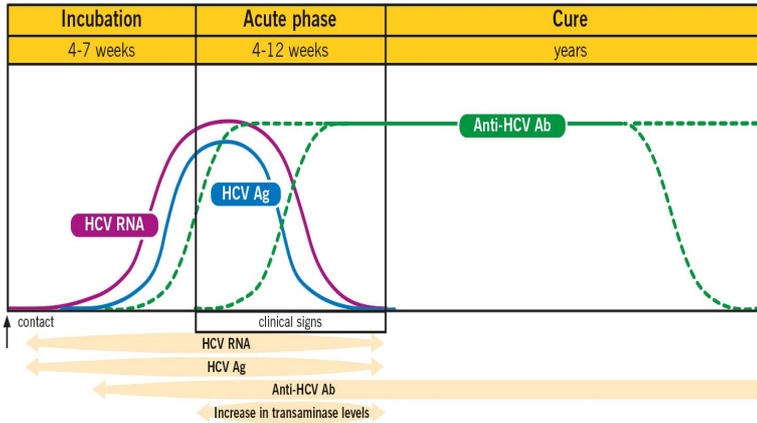
Test Name	Results	Units	Biological Reference Interval
Hepatitis C Virus Antibody <small>(Method: ELISA)</small>	0.22	S/Co	< 1.00 : Negative > 1.00 : Positive

Interpretation:

- Negative result implies that antibodies to HCV have not been detected in the sample. This means the patient has either not been exposed to HCV 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 HCV.
- Positive result implies that antibodies to HCV have been detected in the sample.

Comments :-

Hepatitis C (HCV) is an RNA virus of Flavivirus group transmitted via blood transfusions, transplantation, injection drug users, accidental needle punctures in healthcare workers, dialysis patients and rarely from mother to infant. 10% of new cases show sexual transmission. As compared to HAV & HBV, chronic infection with HCV occurs in 85% of infected individuals. In high risk populations, the predictive value of Anti HCV for HCV infection is > 99% whereas in low risk populations it is only 25%.



Note:

- False positive results are seen in Autoimmune diseases, Rheumatoid factor, Hypergammaglobulinemia, Paraproteinemia, passive antibody transfer, Anti- idiotypes & Anti superoxide dismutase
- False negative results are seen in early Acute infection, Immunosuppression & Immuno-incompetence
- HCV RNA PCR recommended in all Reactive results to differentiate between past and present infection

*** End Of Report ***




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Primary Sample	: Whole Blood	Received On	: 06-May-2025 03:39 PM
Sample Tested In	: Serum	Reported On	: 06-May-2025 06:57 PM
Client Address	: Kimtee colony ,Gokul Nagar,Tarnaka	Report Status	: Final Report

IMMUNOLOGY & SEROLOGY

SURGICAL PROFILE-II

Test Name	Results	Units	Biological Reference Interval
HIV (1& 2) Antibody <small>(Method: ELISA)</small>	0.31	S/Co	< 1.00 : Negative > 1.00 : Positive

*** End Of Report ***



Page 7 of 7


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MD, MICROBIOLOGIST

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