



# PARKLINE DIAGNOSTICS PVT. LTD.

L.G. 3, 4 & 5, Bhuvana Towers, S.D. Road, Secunderabad - 500 003 Tel :+91 40-2784 5852, 6649 1787  
Fax : +91 40 2784 7864 Email parklinediagnostics@gmail.com www.parklinediagnostics.com



NABL Accredited  
Certificate No.MC-2566

## TEST REPORT

Name	: MR.VADDE SRI HARI [62107]	TID/SID	: UMR0638671/ 22115553
Age / Gender	: 52 Years / Male	Registered on	: 29-Dec-2021 / 08:44 AM
Ref.By	: -	Collected on	: 29-Dec-2021 / 08:46 AM
Req.No	:  BIL1618852	Reported on	: 29-Dec-2021 / 12:28 PM
		Reference	: Medi Wheel

### DEPARTMENT OF CLINICAL PATHOLOGY

### Complete Urine Examination (CUE), Urine

Investigation	Observed Value	Units	Biological Reference Interval
Colour Method:Photo detectors(instrument)	Light Yellow		Light Yellow
Appearance Method:Photo diode array sensor	Clear		Clear
Specific gravity Method:Ion concentration/colour indicator	1.005		1.003-1.030
Reaction and pH Method:Double Indicator	6.0		5.0-8.0
Protein Method:Protein Error of pH indicators	Negative		Negative
Glucose Method:Double sequential enzymatic/GOD-PAP	Negative		Negative
Urobilinogen Method:Reagent strip/Reflectance photometry	Negative	mg%	0.2-1.0
Ketones Method:Strip method/Nitroprusside method	Negative		Negative
Blood Method:Peroxidase	Negative		Negative
Bile Salt Method:Hays Method	Negative		Negative
Bile Pigment Method:Fouchets Method	Negative		Negative
<b>Microscopic Examination</b>			
Pus cells (leukocytes) Method:Microscopy Of Sediment	Nil	/hpf	0-5
RBC (erythrocytes) Method:Microscopy Of Sediment	Nil	/hpf	0-2
Epithelial cells Method:Microscopy Of Sediment	Nil	/hpf	0-8
Crystals Method:Microscopy Of Sediment	Nil	/lpf	Nil

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### Complete Urine Examination (CUE), Urine

Investigation	Observed Value	Units	Biological Reference Interval
Casts Method:Microscopy Of Sediment	Nil	/lpf	Nil
Others Method:Microscopy Of Sediment	Nil		Nil

\* Sample processed at Parkline

--- End Of Report ---

**Dr.Jyothi Kiranmai**  
Regd. No: 52272  
MD PATHOLOGY

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Call : 7995421787, 7093445852,8121147282, 9885202212



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
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BIL1618852

### DEPARTMENT OF HEMATOLOGY

### Blood Grouping ABO & Rh Typing, EDTA Whole Blood

Parameter	Results
Blood Grouping (ABO)	AB
Rh Typing (D)	POSITIVE -
Method: Agglutination	

**Method:** Hemagglutination Tube Method by Forward & Reverse Grouping

**Reference:** Tulip kit literature

**Interpretation:** The ABO grouping and Rh typing test determines blood type grouping (A,B, AB, O ) and the Rh factor (positive or negative). A person's blood type is based on the presence or absence of certain antigens on the surface of their red blood cells and certain antibodies in the plasma. ABO antigens are poorly expressed at birth, increase gradually in strength and become fully expressed around 1 year of age. In case of Rh(D) - Du(weak positive) or Weak D positive, the individual must be considered as Rh positive as donor and Rh negative as recipient.

**Note:** Records of previous blood grouping/Rh typing not available. Please verify before transfusion.

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MD PATHOLOGY

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
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### DEPARTMENT OF HEMATOLOGY

### Complete Blood Picture (CBP), EDTA Whole Blood

Investigation	Observed Value	Units	Biological Reference Interval
Hemoglobin., Method:Spectrophotometry	14.0	gm/dL	13.0-17.0 g/dL
Erythrocyte Count(RBC) Method:Electrical Impedence	<b>4.4</b>	mill /cu.mm	4.5-5.5
PCV/HCT., Method:Numeric Integration	41	%	40-50
MCV., Method:Calculated	93	fL	83-101
MCH., Method:Calculated	31.3	pg	27-32
MCHC Method:Calculated	33.6	g/dL	31.5-34.5 gm/dL
RDW (CV)., Method:Calculated	<b>14.2</b>	%	11.6-14.0
Total WBC Count Method:Impedence flowcytometry/Light scattering	5.6	cells/cumm	4-10
<b>Differential Count</b>			
Neutrophils:., Method:Flowcytometry/Microscopy	74	%	40-80
Lymphocytes:., Method:Flowcytometry/Microscopy	<b>19</b>	%	20-40
Monocytes:., Method:Flowcytometry/Microscopy	5	%	2-10
Eosinophils:., Method:Flowcytometry/Microscopy	2	%	1-6
Basophils:., Method:Flowcytometry/Microscopy	0	%	0-2
Absolute Neutrophil Count	4.14	cells/cumm	2.0-7.0
Absolute Lymphocyte Count	1.06	cells/cumm	1.0-3.0
Absolute Monocyte Count	0.28	cells/cumm	0.20-1.0
Absolute Eosinophil Count	0.11	cells/cumm	0.02-0.5

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Absolute Basophil Count	<b>0</b>	cells/cumm	0.02-0.1
Platelet Count	230	lakhs/cumm	150-410
Method:Electrical Impedence			

### Peripheral Smear

RBC	Normocytic and Normochromic
Method:Microscopy	
WBC	Within normal limits.No abnormal cells seen.
Method:Microscopy	
Platelets	Discrete and adequate.Normal in morphology
Method:Microscopy	

\* Sample processed at Parkline

--- End Of Report ---

**Dr.Jyothi Kiranmai**  
Regd. No: 52272  
MD PATHOLOGY

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### DEPARTMENT OF HEMATOLOGY

#### Erythrocyte Sedimentation Rate (ESR), Sodium Citrate Whole Blood

Investigation	Observed Value	Units	Biological Reference Intervals
ESR 1st Hour	6	mm/hour	0-10
Method:Westergren			

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
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### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

#### Blood Urea Nitrogen (BUN), Serum

Investigation	Observed Value	Units	Biological Reference Interval
Blood Urea Nitrogen.	9.3	mg/dL	7-23
Method:Calculated			

#### Creatinine, Serum

Investigation	Observed Value	Units	Biological Reference Interval
Creatinine.	0.94	mg/dL	0.60-1.30
Method:Alkaline Picrate			

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
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Ref.By	: -	Collected on	: 29-Dec-2021 / 08:46 AM
Req.No	:  BIL1618852	Reported on	: 29-Dec-2021 / 14:00 PM
		Reference	: Medi Wheel

### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

### Glucose Fasting (FBS), Sodium Fluoride Plasma

Investigation	Observed Value	Units	Biological Reference Interval
Glucose Fasting Method:GOD - PAP	<b>113</b>	mg/dL	Normal: <100 Impaired FG: 100-125 Diabetic : >=126

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
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Req.No  Reported on : 29-Dec-2021 / 14:00 PM  
BIL1618852 Reference : Medi Wheel

### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

#### Glucose Post Prandial (PPBS), Sodium Fluoride Plasma

Investigation	Observed Value	Units	Biological Reference Interval
Glucose Post Prandial Method:GOD - PAP	137	mg/dL	Normal : 90 - 140 Impaired Glucose Tolerance: 141-199 Diabetic : $\geq$ 200

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MD PATHOLOGY

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 BIL1618852 Reference : Medi Wheel

### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

### Glycosylated Hemoglobin (HbA1C), EDTA Whole Blood

Investigation	Observed Value	Units	Biological Reference Interval
Glycosylated Haemoglobin Method:High Performance Liquid Chromatography(HPLC)	5.5	%	< 5.7 : Normal 5.7 - 6.4 : Prediabetes > 6.4 Diabetes
Mean Plasma Glucose (MPG) Estimate Method:Derived from HBA1c	111	mg/dL	Excellent Control : 90 to 120 Good Control : 121 to 150 Average Control : 151 to 180 Panic Value : > 211

**Note:**Mean Plasma Glucose is calculated from HBA1c value and it indicates Average Blood Sugar level over the past three months.

#### INTERPRETATION :

- Glycated hemoglobin (glycohemoglobin / HbA1c) is a form of hemoglobin (Hb) that is chemically linked to a sugar.
- A1c is measured primarily to determine the three-month average blood sugar level and can be used as a diagnostic test for diabetes mellitus and as an assessment test for glycemic control in people with diabetes.
- In diabetes, higher amounts of glycated hemoglobin, indicating poorer control of blood glucose levels, have been associated with cardiovascular disease, nephropathy, neuropathy, and retinopathy.
- American diabetes Association (ADA) recommends an A1C goal for many non pregnant adults of < 7% (without significant hypoglycemia). On the basis of provider judgment and patient preference, achievement of lower A1C levels than the goal of 7% may be acceptable, and even beneficial, if it can be achieved safely without significant hypoglycemia or other adverse effects of treatment. Less stringent A1C goals (such as < 8%) may be appropriate for patients with severe hypoglycemia, extensive co morbid conditions etc, or where the harms of treatment are greater than the benefits.
- Glycemic goals for some older adults might reasonably be relaxed as part of individualized care, but hyperglycemia leading to symptoms or risk of acute hyperglycemia complications should be avoided in all patients.

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**Dr.Divya Panda**  
Regd. No: 84506  
MD Pathology

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
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### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

#### Lipid Profile, Serum

Investigation	Observed Value	Units	Biological Reference Interval
Total Cholesterol Method:CHOD-PAP	169	mg/dL	Desirable Level: < 200 Borderline : 200 - 239 High : > 240
HDL Cholesterol Method:Enzymatic Reaction	37	mg/dL	<40:Major risk factor for heart disease 40-59:The higher,the better >=60:Considered protective against heart disease
LDL Cholesterol Method:Calculated	<b>111</b>	mg/dL	< 100
VLDL Cholesterol Method:Calculated	21	mg/dL	10-55
Triglycerides Method:GPO-POD	109	mg/dL	Normal:<150 Borderline:150-199 High:200-499 Very High:>=500
Chol/HDL Ratio Method:Calculated	4.57		Normal : <4 Low risk : 4 - 6 High risk : >6
LDL Cholesterol/HDL Ratio	3.00		

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### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

### Liver Function Test (LFT), Serum

Investigation	Observed Value	Units	Biological Reference Interval
Total Bilirubin. Method:Diazo with sulphanilic acid	1.04	mg/dL	0.3-1.2
Direct Bilirubin. Method:Diazo with sulphanilic acid	0.14	mg/dL	0.00-0.40
Indirect Bilirubin. Method:Calculated	0.90	mg/dL	
Alanine Aminotransferase ,(ALT/SGPT) Method:IFCC without P5P	<b>42</b>	U/L	10-40
Aspartate Aminotransferase,(AST/SGOT) Method:IFCC without P5P	44	U/L	10-40
ALP (Alkaline Phosphatase). Method:AMP-IFCC	67	U/L	30-115
<b>PROTEINS</b>			
Total Protein. Method:Biuret	6.64	g/dL	6.0-8.0
Albumin. Method:Bromocresol Green (BCG)	4.58	g/dL	3.5-4.8
Globulin. Method:Calculated	<b>2.06</b>	g/dL	2.3-3.5
A/GRatio. Method:Calculated	<b>2.22</b>		0.8-2.0
Gamma GT. Method:IFCC-Enzymatic	<b>131</b>	U/L	7.0-50.0

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
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NABL Accredited  
Certificate No.MC-2566

## TEST REPORT

Name : **MR.VADDE SRI HARI [62107]** TID/SID : UMR0638671/ 22115552  
 Age / Gender : 52 Years / Male Registered on : 29-Dec-2021 / 08:44 AM  
 Ref.By : - Collected on : 29-Dec-2021 / 08:46 AM  
 Req.No  Reported on : 29-Dec-2021 / 13:08 PM  
 BIL1618852 Reference : Medi Wheel

### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

### Prostate Specific Antigen (PSA) Total, Serum

Investigation	Observed Value	Biological Reference Interval
Prostate Specific Antigen (PSA) Total	0.592 ng/mL	0-3.9
Method:Enhanced chemiluminescence		

#### Interpretation:

- 1.Prostate specific antigen (PSA) is a glycoprotein that is expressed by both normal and neoplastic prostate tissue
- 2.Elevated serum PSA concentrations are found in men with prostate cancer, benign prostatic hyperplasia (BPH) or inflammatory conditions of other adjacent genitourinary tissues. PSA can also be elevated after digital rectal examination,prostatic massage,cystoscopy,needle biopsy etc
- 3.Measurement of serum PSA by itself is not recommended as a screening procedure for the diagnosis of cancer because elevated PSA levels are also observed in patients with benign prostatic hyperplasia.
4. When employed for the management of prostate cancer patients, serial measurement of PSA is useful in detecting residual tumor and recurrent cancer after radical prostatectomy.
- 5.PSA has been demonstrated to be an accurate marker for monitoring advanced clinical stage in untreated patients and for monitoring response to therapy by radical prostatectomy, radiation therapy and anti-androgen therapy.

\* Sample processed at Parkline

--- End Of Report ---

**Dr.Jyothi Kiranmai**  
Regd. No: 52272  
MD PATHOLOGY

The Test marked with \*are not accredited by NABL.

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 BIL1618852 Reference : Medi Wheel

### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

#### Thyroid Profile (T3,T4,TSH), Serum

Investigation	Observed Value	Units	Biological Reference Interval
Triiodothyronine Total (T3) Method:Enhanced chemiluminescence	1.31	ng/mL	0.970-1.69
Thyroxine Total (T4). Method:Enhanced chemiluminescence	<b>11.3</b>	µg/dL	5.53-11.0
Thyroid Stimulating Hormone (TSH). Method:Enhanced chemiluminescence	1.92	µIU/mL	0.465-4.68 µIU/mL

Note: Change in method and reference range  
NOTE:

TSH - Reference ranges during pregnancy:\*

1st Trimester : 0.10 - 2.50

2nd Trimester : 0.20 - 3.00

3dr Trimester : 0.30 - 3.00

\*As per the Guidelines of American Thyroid Association for the diagnosis and management of thyroid disease during pregnancy and post partum.

1.Primary Hyperthyroidism is accompanied by elevated T3 & T4 values along with depressed TSH level.

2.Primary Hypothyroidism is accompanied by depressed T3 & T4 levels and elevated TSH levels.

3.Normal T4 levels accompanied by high T3 levels are seen in patients with T3 Thyrotoxicosis.

4.Slightly elevated T3 levels may be found in pregnancy and estrogen therapy, while depressed levels may be encountered in severe illness, malnutrition, renal failure and during therapy with drugs like propranolol and propylthiouracil.

5.Although elevated TSH levels are nearly always indicative of primary hypothyroidism, rarely they can result form TSH secreting pituitary tumors(secondary).

\* Sample processed at Parkline

--- End Of Report ---

**Dr.Jyothi Kiranmai**  
Regd. No: 52272  
MD PATHOLOGY

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Reference : Medi Wheel  
BIL1618852

### DEPARTMENT OF CLINICAL BIOCHEMISTRY I

#### Uric Acid, Serum

Investigation	Observed Value	Units	Biological Reference Interval
Uric Acid. Method:Uricase	4.91	mg/dL	2.5-8.0

\* Sample processed at Parkline

--- End Of Report ---

**Dr.Jyothi Kiranmai**  
Regd. No: 52272  
MD PATHOLOGY

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## TEST REPORT

Name : **MR.VADDE SRI HARI [62107]** TID/SID : UMR0638671/ 22116598  
Age / Gender : 52 Years / Male Registered on : 29-Dec-2021 / 08:44 AM  
Ref.By : - Collected on : 29-Dec-2021 / 12:24 PM  
Req.No  Reported on : 29-Dec-2021 / 15:02 PM  
Reference : Medi Wheel  
BIL1618852

### DEPARTMENT OF HEALTH CHECKUP

#### Glucose Urine Fasting

Urine Glucose Fasting Nil NIL  
Method:Reagent strip/Reflectance photometry

#### Glucose Urine Post Prandial

Urine Glucose Post Prandial Nil NIL  
Method:Reagent strip/Reflectance photometry

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--- End Of Report ---

**Dr.Divya Panda**  
Regd. No: 84506  
MD Pathology

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Ref.By	: -	Collected on	: 29-Dec-2021 / 08:44 AM
Req.No	:  BIL1618852	Reported on	: 29-Dec-2021 / 11:59 AM
		Reference	: Medi Wheel

### DEPARTMENT OF CARDIOLOGY

#### 2D Echo/Doppler Study

Mitral Valve	Normal
Aortic valve	Normal
Tricuspid valve	Normal
Pulmonary valve	Normal
Aorta	3.1 cm
Left Atrium	3.6 cm
Left Ventricle	LVDd:4.6 cm IVSd :1.0 cm EF:64% LVDs:3.0 cm LVPwd:1.1 cm FS:34%
RWMA	Nil
Right Atrium	Normal
Right Ventricle	Normal
Pulmonary Artery	Normal
IAS	Intact
IVS	Intact
Pericardium	Normal
Svc / Ivc	Normal
Intracardiac Masses	Nil
Doppler Study	Mitral flow: A > E Aortic flow : 1.0 m/sec Pulmonary flow : 1.2 m/sec
Colour Doppler	No MR / AR / TR / PR
Conclusion	No RWMA. Normal valves/ Normal chambers. No MR/ AR/ TR / PR Good LV/ RV function. No PE/ clot/ vegetation.

\* Sample processed at Parkline

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