



DI FASE SCAN OR COD

Name : Mr . CHILUKURI PRASAD TID

Age/Gender : 53 Years/Male Registered On : 09-Nov-2024 07:27 AM

Ref By : Self

Reg.No : BIL4920695 Reference : Arcofemi Health Care Ltd

- Medi Whe

Reported On

: UMR2156671

: 09-Nov-2024 02:18 PM

DEPARTMENT OF CARDIOLOGY 2D Echo/Doppler Study

MITRAL VALVE : Normal.

AORTIC VALVE : Mild sclerotic.

TRICUSPID VALVE : Normal.

PULMONARY VALVE : Normal.

RIGHT ATRIUM : Normal.

RIGHT VENTRICLE : Normal.

LEFT ATRIUM : 3.6 cms.

LEFT VENTRICLE : EDD : 3.8 cm IVS (d) : 1.1 cm LVEF : 64%

ESD: 2.6 cm PW (d): 1.1 cm FS: 32%

NO RWMA

IAS : Intact.

IVS : Intact.

AORTA : 2.5 cms.

PULMONARY ARTERY: Normal.

PERICARDIUM : Normal.

IVC/ SVC / CS : Normal.

PULMONARY VEINS : Normal.

INTRA - CARDIAC MASSES : No.





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: 09-Nov-2024 02:18 PM

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Age/Gender : 53 Years/Male Registered On : 09-Nov-2024 07:27 AM

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- Medi Whe

Reported On

DOPPLER STUDY

MITRAL FLOW : E: 0.8 m/s A: 0.7 m/s

AORTIC FLOW : AJV: 1.1 m/s

PULMONARY FLOW : PJV: 0.6 m/s

TRICUSPID FLOW : TRJV: 1.6 m/s, RVSP: 22mmHg

COLOUR FLOW MAPPING

MR : NIL
AR : NIL
TR : NIL
PR : NIL

IMPRESSION:

- * NORMAL SIZED CARDIAC CHAMBERS
- * INTACT SEPTAE
- * NO RWMA
- * GOOD LV / RV SYSTOLIC FUNCTION
- * NO MR / NO AR / NO TR
- * NO PE / NO CLOT OR VEGETATION

*** End Of Report ***

Dr.K S Reddy PGDCC(Dip. Card) Clinical Cardiologist







Name
Age / Gender
Ref By

: MR.CHILUKURI PRASAD

: 53 Years / Male

Ref.By : SELF

Req.No : BIL4920695

TID/SID : UMR2156671/ 28533464 Registered on : 09-Nov-2024 / 07:27 AM

Collected on : 09-Nov-2024 / 07:37 AM Reported on : 09-Nov-2024 / 12:12 PM

TEST REPORT Reference : Arcofemi Health Care Ltd -

DEPARTMENT OF CLINICAL PATHOLOGY

Complete Urine Examination (CUE)

Investigation	Result	Biological Reference Intervals
Physical Examination		
Colour	Pale yellow	Straw to Yellow
Method:Physical		
Appearance	Clear	Clear
Method:Physical		
Chemical Examination		
Reaction and pH	Acidic (6.0)	4.6-8.0
Method:Indicator	4.000	4.000 4.005
Specific gravity	1.009	1.000-1.035
Method:Refractometry	Namathia	Namakiya
Protein Method Bassia France folklindings	Negative	Negative
Method:Protein Error of pH indicators	Negotivo	Negativo
Glucose Method:Glucose oxidase/Peroxidase	Negative	Negative
	Negative	Negative
Blood Method:Peroxidase	regative	Negative
Ketones	Negative	Negative
Method:Sodium Nitroprusside Method	rioganio	. roga ro
Bilirubin	Negative	Negative
Method:Diazonium salt	3	S
Leucocytes	Negative	Negative
Method:Esterase reaction	-	•
Nitrites	Negative	Negative
Method:Modified Griess reaction		
Urobilinogen	Negative	Up to 1.0 mg/dl
Method:Diazonium salt		(Negative)
Microscopic Examination		
Pus cells (leukocytes)	1-2	2 - 3 /hpf
Method:Flow Digital Imaging/Microscopy		
Epithelial cells	1-2	2 - 5 /hpf
Method:Flow Digital Imaging/Microscopy		
RBC (erythrocytes)	Absent	Absent
Method:Flow Digital Imaging/Microscopy		
Casts	Absent	Occasional hyaline casts may be seen
Method:Flow Digital Imaging/Microscopy		







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TEST REPORT Reference : Arcofemi Health Care Ltd -

be seen

Crystals Absent Phosphate, oxalate, or urate crystals may

Method:Flow Digital Imaging/Microscopy

Others Nil Nil

Method:Flow Digital Imaging/Microscopy

Method: Semi Quantitative test ,For CUE

Reference: Godka**r** Clinical Diagnosis and Management by Laboratory Methods, First South Asia edition. Product kit literature.

Interpretation:

The complete urinalysis provides a number of measurements which look for abnormalities in the urine. Abnormal results from this test can be indicative of a number of conditions including kidney disease, urinary tract infecation or elevated levels of substances which the body is trying to remove through the urine. A urinalysis test can help identify potential health problems even when a person is asymptomatic. All the abnormal results are to be correlated clinically.

* Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad

--- End Of Report ---

Dr Shruti Reddy Consultant Pathologist Reg No.TSMC/FMR/22656







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TID/SID : UMR2156671/ 28533465 Registered on : 09-Nov-2024 / 07:27 AM

Collected on : 09-Nov-2024 / 07:37 AM

Reported on : 09-Nov-2024 / 13:42 PM

TEST REPORT Reference : Arcofemi Health Care Ltd -

DEPARTMENT OF HEMATOPATHOLOGY

Blood Grouping ABO And Rh Typing

Parameter Results

Blood Grouping (ABO) O

Rh Typing (D) Positive

Method:Hemagglutination Tube Method by Forward & Reverse Grouping

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 expresses 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.

* Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad

--- End Of Report ---

Dr Reenaz Shaik Consultant Pathologist







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Reported on : 09-Nov-2024 / 11:23 AM

TEST REPORT Reference : Arcofemi Health Care Ltd -

DEPARTMENT OF HEMATOPATHOLOGY

Erythrocyte Sedimentation Rate (ESR)

Investigation	Observed Value	Biological Reference Intervals
ESR 1st Hour	5	<=12 mm/hour

Method:Westergren/Vesmatic

Complete Blood Count (CBC)

Investigation	Observed Value	Biological Reference Intervals
Hemoglobin	15.7	13.0-17.0 g/dL
Method:Cyanide Free Lyse Hemoglobin		
PCV/HCT	46.6	40.0-50.0 vol%
Method:Calculated		
Total RBC Count	5.02	4.50-5.50 mill /cu.mm
Method:Electrical Impedance		
MCV	92.9	83.0-101.0 fL
Method:Calculated		
MCH	31.2	27.0-32.0 pg
Method:Calculated		
MCHC	33.6	31.5-34.5 g/dL
Method:Calculated		
RDW (CV)	13.0	11.6-14.0 %
Method:Calculated		
MPV	9.0	7.0-10.0 fL
Method:Calculated		
Total WBC Count	7500	4000-10000 cells/cumm
Method:Electrical Impedance		
Platelet Count	3.42	1.50-4.10 lakhs/cumm
Method:Electrical Impedance		
Differential count		
Neutrophils	55.2	40.0-80.0 %
Method:Microscopy	24.2	00 0 40 0 0
Lymphocytes	31.6	20.0-40.0 %
Method:Microscopy	0.5	1.0.0.0.0
Eosinophils	3.5	1.0-6.0 %
Monocytes	8.8	2.0-10.0 %
Basophils	0.9	< 1.0-2.0 %
Method:Flowcytometry/Electrical Impedance/Micro	200001/	







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TEST REPORT Reference : Arcofemi Health Care Ltd -

Absolute Neutrophil Count Method:Calculated	4140	2000-7000 cells/cumm
Absolute Lymphocyte Count (ALC)	2370	1000-3000 cells/cumm
Absolute Eosinophil Count (AEC)	263	20-500 cells/cumm
Absolute Monocyte Count Method:Calculated	660	200-1000 cells/cumm
Absolute Basophil Count Method:Calculated	68	20-100 cells/cumm
Neutrophil - Lymphocyte Ratio(NLR) Method:Calculated	1.75	0.78-3.53

Method: Automated Hematology Cell Counter, Microscopy

Reference: Dacie and Lewis Practical Hematology, 12th Edition. Wallach's interpretation of diagnostic tests, Soth Asian Edition.

Interpretation: A Complete Blood Picture (CBP) is a screening test which can aid in the diagnosis of a variety of conditions and diseases such as anemia, leukemia, bleeding disorders and infections. This test is also useful in monitoring a person's reaction to treatment when a condition which affects blood cells has been diagnosed. All the abnormal results are to be correlated clinically.

Note: These results are generated by a fully automated hematology analyzer and the differential count is computed from a total of several thousands of cells. Therefore the differential count appears in decimalised numbers and may not add upto exactly 100. It may fall between 99 and 101.

* Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad

--- End Of Report ---

Dr Reenaz Shaik Consultant Pathologist







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TEST REPORT Reference : Arcofemi Health Care Ltd -

Blood Urea Nitrogen (BUN) Investigation Observed Value Biological Reference Interval Blood Urea Nitrogen. 10.79 6-20 mg/dL Method:Calculated Urea. 23.1 12.8-42.8 mg/dL Method:Urease

Interpretation: Urea is a waste product formed in the liver when protein is metabolized. Urea is released by the liver into the blood and is carried to the kidneys, where it is filtered out of the blood and released into the urine. Since this is a continuous process, there is usually a small but stable amount of urea nitrogen in the blood. However, when the kidneys cannot filter wastes out of the blood due to disease or damage, then the level of urea in the blood will rise. The blood urea nitrogen (BUN) evaluates kidney function in a wide range of circumstances, to diagnose kidney disease, and to monitor people with acute or chronic kidney dysfunction or failure. It also may be used to evaluate a person's general health status as well.

Reference: Tietz Fundamentals of Clinical Chemistry and Molecular Diagnostics

Creatinine, Serum

Investigation	Observed Value	Biological Reference Interval	
Creatinine.	0.96	0.70-1.20 mg/dL	
Method:Alkaline Picrate			

Interpretation:

Creatinine is a nitrogenous waste product produced by muscles from creatine. Creatinine is majorly filtered from the blood by the kidneys and released into the urine, so serum creatinine levels are usually a good indicator of kidney function. Serum creatinine is more specific and more sensitive indicator of renal function as compared to BUN because it is produced from muscle at a constant rate and its level in blood is not affected by protein catabolism or other exogenous products. It is also not reabsorbed and very little is secreted by tubules making it a reliable marker. Serum creatinine levels are increased in pre renal, renal and post renal azotemia, active acromegaly and gigantism. Decreased serum creatinine levels are seen in pregnancy and increasing age.

Glucose Fasting (FBS)

	_ ·	•
Investigation	Observed Value	Biological Reference Interval
Glucose Fasting Method:Hexokinase	92	Normal: <100 mg/dL Impaired FG: 100-125 mg/dL Diabetes mellitus: >/=126 mg/dL

Interpretation: It measures the Glucose levels in the blood with a prior fasting of 9-12 hours. The test helps screen a symptomatic/ asymptomatic person who is at risk for Diabetes. It is also used for regular monitoring of glucose levels in people with Diabetes.

Reference: American Diabetes Association. Standards of Medical Care in Diabetes-2022





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Reference : Arcofemi Health Care Ltd -

Glucose Post Prandial (PPBS)

TEST REPORT

Investigation	Observed Value	Biological Reference Interval
Glucose Post Prandial Method:Hexokinase	94	Normal : <140 mg/dL Impaired PG: 140-199 mg/dL Diabetes mellitus: >/=200 mg/dL

Interpretation: This test measures the blood sugar levels 2 hours after a normal meal. Abnormally high blood sugars 2 hours after a meal reflect that the body is not producing sufficient insulin which is indicative of Diabetes.

Reference: American Diabetes Association. Standards of Medical Care in Diabetes-2022

Glycosylated Hemoglobin (HbA1C)

	<u>-</u>	
Investigation	Observed Value	Biological Reference Interval
Glycosylated Hemoglobin (HbA1c) Method:High-Performance Liquid Chromatography	5.2	Non-diabetic: <= 5.6 % Pre-diabetic: 5.7 - 6.4 % Diabetic: >= 6.5 %
Estimated Average Glucose (eAG) Method:Calculated	103	mg/dL

Interpretation:

It is an index of long-term blood glucose concentrations and a measure of the risk for developing microvascular complications in patients with diabetes. Absolute risks of retinopathy and nephropathy are directly proportional to the mean HbA1c concentration. In persons without diabetes, HbA1c is directly related to risk of cardiovascular disease.

- 1) Low glycated haemoglobin (below 4%) in a non-diabetic individual are often associated with systemic inflammatory diseases, chronic anaemia (especially severe iron deficiency & haemolytic), chronic renal failure and liver diseases. Clinical correlation suggested.
- 2) Interference of Hemoglobinopathies in HbA1c estimatiion:
- A. For HbF > 25%, an alternate platform (Fructosamine) is recommended for testing of HbA1c.
- B. Homozygous hemoglobinopathy is detected, fructosamine is recommended for monitoring diabetic status
- C. Heterozygous state detected (D10 is corrected for HbS and HbC trait).
- 3) In known diabetic patients, HbA1c can be considered as a tool for monitoring the glycemic control.

Excellent Control - 6 to 7 %,

Fair to Good Control - 7 to 8 %,

Unsatisfactory Control - 8 to 10 %

and Poor Control - More than 10 %.

Reference: American Diabetes Association. Standards of Medical Care in Diabetes-2022.

Bun/Creatinine Ratio

	Bun/Creatinin	e Halio	
Investigation	Observed Val	ue	
BUN/Creatinine Ratio Method:Calculated	10	10-20	





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TEST REPORT Reference : Arcofemi Health Care Ltd -

TID/SID

Interpretation:

The BUN/Creatinine ratio blood test is used to diagnose acute or chronic renal disease. BUN (blood urea nitrogen) and creatinine are both filtered in the kidneys and excreted in urine. The two together are used to measure overall kidney function

- 1. Increased ratio (>20) with normal creatinine occurs in the following conditions:
- a) Increased BUN (prerenal azotemia), heart failure, salt depletion, dehydration
- b) Catabolic states with tissue breakdown
- c) GI hemorrhage
- d) Impaired renal function plus excess protein intake, production, or tissue breakdown
- 2. Increased ratio (>20) with elevated creatinine occurs in the following conditions:
- a) Obstruction of urinary tract
- b) Prerenal azotemia with renal disease
- 3. Decreased ratio (<10) with decreased BUN occurs in the following conditions:
- a) Acute tubular necrosis
- b) Decreased urea synthesis as in severe liver disease or starvation
- c) Repeated dialysis
- d) SIADH
- e) Pregnancy
- 4. Decreased ratio (<10) with increased creatinine occurs in the following conditions:
- a) Phenacemide therapy (accelerates conversion of creatine to creatinine)
- b) Rhabdomyolysis (releases muscle creatinine)
- c) Muscular patients who develop renal failure

* Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad









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Note

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TEST REPORT

Reference : Arcofemi Health Care Ltd -

DEPARTMENT OF CLINICAL CHEMISTRY I Lipid Profile Observed Value Biological Reference Interval Investigation 234 Desirable: <200 mg/dL Total Cholesterol Borderline: 200-239 mg/dL Method:Cholesterol Oxidase High: >/=240 mg/dL 42 Low: <40 mg/dL **HDL Cholesterol** High: >/=60 mg/dL Method:Direct Measurement 43.20 6.0-38.0 mg/dL **VLDL Cholesterol** Method:Calculated 148.8 Optimum: <100 ma/dL LDL Cholesterol Near/above optimum: 100-129 mg/dL Method:Calculated Borderline: 130-159 mg/dL High: 160-189 mg/dL Very high: >/=190 mg/dL Normal:<150 mg/dL 216 **Triglycerides** Borderline: 150-199 mg/dL Method:Glycerol LPL/GK High: 200-499 mg/dL Very high: >/=500 mg/dL 5.57 Low Risk: 3.3-4.4 Chol/HDL Ratio Average Risk: 4.5-7.1 Method:Calculated Moderate Risk: 7.2-11.0 3.54 Desirable: 0.5-3.0 LDL Cholesterol/HDL Ratio Borderline Risk: 3.0-6.0 Method:Calculated High Risk: >6.0 192 <130 mg/dL Non HDL Cholesterol Method:Calculated

Interpretation: Lipids are fats and fat-like substances which are important constituents of cells and are rich sources of energy. A lipid profile typically includes total cholesterol, high density lipoproteins (HDL), low density lipoprotein (LDL), chylomicrons, triglycerides, very low density lipoproteins (VLDL), Cholesterol/HDL ratio .The lipid profile is used to assess the risk of developing a heart disease and to monitor its treatment. The results of the lipid profile are evaluated along with other known risk factors associated with heart disease to plan and monitor treatment. Treatment options require clinical correlation.

Kindly correlate clinically

Reference: Third Report of the National Cholesterol Education program (NCEP) Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults (Adult Treatment Panel III), JAMA 2001.



^{*} Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad





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TEST REPORT Reference : Arcofemi Health Care Ltd -

Dr Afreen Anwar Consultant Biochemist









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TEST REPORT Reference : Arcofemi Health Care Ltd -

DEPARTMENT OF CLINICAL CHEMISTRY I

Liver Function Test (LFT)

L IV	Liver i direction rest (Li 1)			
Investigation	Observed Value	Biological Reference Interval		
Total Bilirubin. Method:Diazo Method	1.58	<1.2 mg/dL		
Direct Bilirubin. Method:Diazo Method	0.60	<0.30 mg/dL		
Indirect Bilirubin. Method:Calculated	0.98	<0.9 mg/dL		
Alanine Aminotransferase ,(ALT/SGPT) Method:UV wtihout P5P	22	<45 U/L		
Aspartate Aminotransferase,(AST/SGOT) Method:UV wtihout P5P	29	<35 U/L		
ALP (Alkaline Phosphatase). Method:PNPP-AMP Buffer	102	40-129 U/L		
Gamma GT. Method:GCNA	25	10-71 U/L		
Total Protein. Method:Biuret & Bromocresol Green (BCG)	7.4	6.6-8.7 g/dL		
Albumin. Method:Bromocresol Green (BCG)	4.8	3.5-5.2 g/dL		
Globulin. Method:Calculated	2.60	1.8-3.8 g/dL		
A/GRatio. Method:Calculated	1.85	0.8-2.0		
AST/ALT Ratio Method:Calculated	1.32	<1.00		

Interpretation: Liver functions tests help to identify liver disease, its severity, and its type. Generally these tests are performed in combination, are abnormal in liver disease, and the pattern of abnormality is indicative of the nature of liver disease. An isolated abnormality of a single liver function test usually means a non-hepatic cause. If several liver function tests are simultaneously abnormal, then hepatic etiology is likely.



^{*} Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad







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DEPARTMENT OF CLINICAL CHEMISTRY I

TEST REPORT

Prostate Specific Antigen (PSA) Total

recent opening runger (recent recent		
Investigation	Observed Value	Biological Reference Interval
Prostate Specific Antigen (PSA). Total Method:ECLIA	0.67	<4.4 ng/mL Note: Biological Reference Ranges are changed due to change in method of testing.

Interpretation: PSA is a protein produced by cells in the prostate and is used to screen men for prostate cancer. PSA levels are elevated in Prostate cancer, and other conditions such as benign prostatic hyperplasia (BPH) and inflammation of the prostate. An elevated PSA may be followed by a biopsy and other tests like urinalysis and ultrasound to rule out urinary tract infections and for an accurate diagnosis. PSA levels are vital to determine the effectiveness of treatment and to detect recurrence in diagnosed cases of prostate cancer.

* Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad











TO VERIFY THE REPORT ONLINE

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DEPARTMENT OF CLINICAL CHEMISTRY I

TEST REPORT

Thyroid Profile (T3,T4,TSH)

,			
Investigation	Observed Value	Biological Reference Interval	
Triiodothyronine Total (T3) Method:ECLIA	1.27	0.80-2.00 ng/mL	
Thyroxine Total (T4) Method:ECLIA	9.9	5.1-14.1 μg/dL	
Thyroid Stimulating Hormone (TSH) Method:ECLIA	3.51	0.27-4.20 μIU/mL	

Interpretation:

A thyroid profile is used to evaluate thyroid function and/or help diagnose hypothyroidism and hyperthyroidism due to various thyroid disorders. T4 and T3 are hormones produced by the thyroid gland. They help control the rate at which the body uses energy, and are regulated by a feedback system. TSH from the pituitary gland stimulates the production and release of T4 (primarily) and T3 by the thyroid. Most of the T4 and T3 circulate in the blood bound to protein. A small percentage is free (not bound) and is the biologically active form of the hormones. Reference: Tietz textbook of Clinial Chemistry and Molecular Diagnostics, Nader Rifia, Andrea Ritas Horvath, Carl T. Wittwer.

* Sample processed at National Reference Laboratory. Tenet Diagnostics, Hyderabad

--- End Of Report ---



Dr Afreen Anwar **Consultant Biochemist**







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DEPARTMENT OF CLINICAL CHEMISTRY I Uric Acid, Serum		
Uric Acid. Method:Uricase	6.0	3.4-7.0 mg/dL

TEST REPORT

Interpretation

It is the major product of purine catabolism. Hyperuricemia can result due to increased formation or decreased excretion of uric acid which can be due to several causes like metabolic disorders, psoriasis, tissue hypoxia, preeclampsia, alcohol, lead poisoning, acute or chronic kidney disease, etc. Hypouricemia may be seen in severe hepato cellular disease and defective renal tubular reabsorption of uric acid.

* Sample processed at National Reference Laboratory, Tenet Diagnostics, Hyderabad

--- End Of Report ---



Dr Afreen Anwar Consultant Biochemist









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Reference : Arcofemi Health Care Ltd

- Medi Whe

DEPARTMENT OF ULTRASOUND Ultrasound Whole Abdomen

LIVER is normal shape, size (11.5 cms) and has uniform echopattern. No evidence of focal lesion or intrahepatic biliary ductal dilatation. Hepatic and portal vein radicals are normal.

GALL BLADDER: Partially distended.

CBD is of normal calibre.

PANCREAS has normal shape, size and uniform echopattern.

No evidence of ductal dilatation or calcification.

SPLEEN shows normal shape, size (8.1 cms) and echopattern.

KIDNEYS move well with respiration and have normal shape, size and echopattern.

Cortico- medullary differentiations are well madeout.

No evidence of calculus or hydronephrosis.

Right kidney measures 9.2 x 4.1 cms, Left kidney measures 10.1 x 4.4cms.

URINARY BLADDER shows normal shape and wall thickness.

It has clear contents. No evidence of diverticula.

PROSTATE shows normal shape, size and echopattern.

It measures 2.7 x 3.7 x 3.0 cms, Vol 16.8 cc.

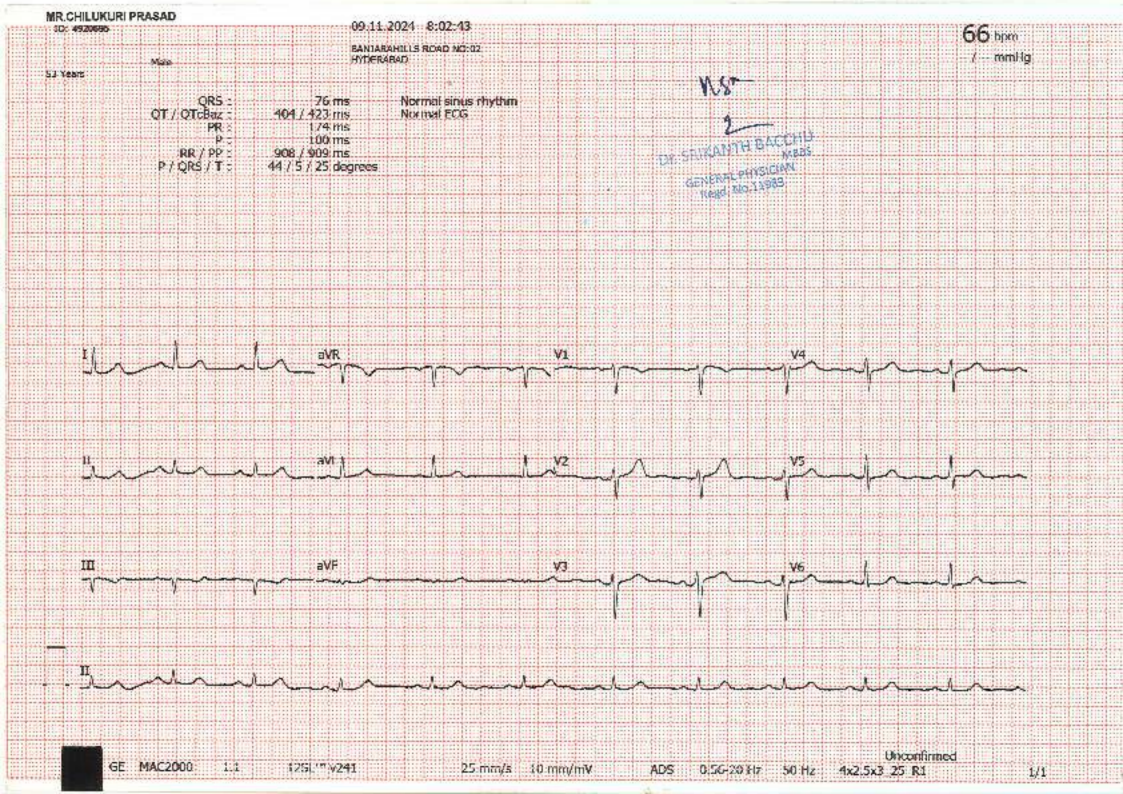
No evidence of free fluid in the abdomen and pelvis.

IMPRESSION:

- * No significant abnormality detected.
- Suggested clinical correlation and follow up

*** End Of Report ***

Dr Sheethal VConsultant Radiologist





: BIL4920695





PLEASE SCAN QR CODE

Name : Mr . CHILUKURI PRASAD TID

Age/Gender : 53 Years/Male Registered On : 09-Nov-2024 07:27 AM

Ref By : Self Reported On : 09-Nov-2024 04:45 PM

Reference : Arcofemi Health Care Ltd

- Medi Whe

: UMR2156671

DEPARTMENT OF X-RAY X-Ray Chest PA View

Radiograph was performed on GE HF ADVANTAGE 400 mA

Lung fields appear normal.

Cardiac size is within normal limits.

Aorta and pulmonary vasculature is normal.

Bilateral domes of diaphragm and costophrenic angles are normal.

Visualised bones and soft tissues appear normal.

IMPRESSION:

Reg.No

* Normal study.

Suggested clinical correlation and follow up.

Study Performed at Tenet Diagnostics Banjarahills, Hyderabad

*** End Of Report ***

Dr Sheethal VConsultant Radiologist



CHILUKURI PRASAD BIL4920695 053Y CHEST PA 09-Nov-24

TENET DIAGNOSTICS, BANJARAHILLS, HYD.