





Lab No. 012411200230

Mr. JAGDISH

Ref. Dr. **MEDIWEEL**

NAME

Rpt. Centre

BarcodeNo

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Age/ Gender 54 YRS/MALE

01200230

Coll. ON

20/Nov/2024 09:16AM

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20/Nov/2024

Approved ON 20/Nov/2024 02:49PM

20/Nov/2024 06:52PM

Test Name	Value	Unit	Biological Reference Interval
Complete Haemogram, EDTA wh	ole blood		
Haemoglobin (Hb) Method : Colorimetry	14.70	gm/dl	13.0 - 17.0
RBC count Method: Electrical impedence	4.45	Millons/cmm	4.5 - 5.5
PCV / Haematocrit Method : Calculated	41.80	%	40.0 - 50.0
MCV Method : Calculated	93.80	fl	83.0 - 101.0
MCH Method: Calculated	33.00	picogram	27.0 - 32.0
MCHC Method : Calculated	35.10	%	31.5 - 34.5
RDW - CV Method : Calculated	13.80	%	11.6 - 14.0
Mentzer I ndex Method : Calculated	21.08		>= 13.0

The Mentzer index (MCV/RBC count) is a useful tool for initial screening of patients with a microcytic hypochromic blood picture to rule out a thalassemia trait. If the index is less than 13, thalassemia is said to be more likely. If the result is greater than 13, then iron-deficiency anemia is said to be more likely. All patients with a low normal to low hemoglobin and a Mentzer index below 13 should be screened for thalassemia trait by HPLC.

TLC (Total Leucocyte Count) Method: Flowcytometry	7,050	/cmm	4000 - 10000
DLC (Flow cytometry)			
Neutrophils	45.80	%	35.0 - 75.0
Lymphocytes	43.40	%	25.0 - 45.0
Eosinophils	4.30	%	1.0 - 5.0
Monocytes	6.00	%	1.0 - 6.0
Basophils	0.50	%	0 - 1
Absolute Leucocyte Count (Calculated)			
Absolute Neutrophil Count	3,228.90	/cmm	2000 - 7000
Absolute Lymphocyte Count	3,059.70	/cmm	1000 - 3000
Absolute Eosinophil count	303.15	/cmm	20 - 500
Absolute Monocyte count	423.00	/cmm	200 - 1000
Absolute Basophil count	35.25	/cmm	0 - 100
Platelet count Method: Electrical impedence	1.67	Lakh/cmm	1.5 - 4.1
ESR (Erythrocyte Sedimentation Rate) Method: Westergren method	20	mm/1st hr	0 - 22

RBCs are normocytic and normochromic.

Leucocytic series is numerically and morphologically within normal limits.

Platelets are adequate in number and are normal in morphology.

No atypical cells or haemoparasites are seen.

Impression: Normal peripheral smear.

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Biological Reference Test Name Value Unit Interval

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Blood Group (ABO + RH)

Blood Group , EDTA blood Method : Slide agglutination (Forward & Reverse grouping)

Rh type , EDTA blood Method : Slide agglutination Positive



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Test Name	Value	Unit	Biological Reference Interval
Glucose Fasting, plasma Method: GOD POD	90.10	mg/dL	60 - 100

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Interpretation (In accordance with the American diabetes association guidelines):

- A fasting plasma glucose level below 100 mg/dl is considered normal.
- A fasting plasma glucose level between 100-126 mg/dl is considered as glucose intolerant or pre diabetic. A fasting and post-prandial blood sugar test (after consumption of 75 gm of glucose) is recommended for all such patients.
- A fasting plasma glucose level of above 126 mg/dl is highly suggestive of a diabetic state. A repeat fasting test is strongly recommended for all such patients. A fasting plasma glucose level in excess of 126 mg/dl on both the occasions is confirmatory of a diabetic state.

Glucose PP, plasma 90 - 140 Method : GOD POD

Interpretation (In accordance with the American diabetes association guidelines):

- A post-prandial plasma glucose level below 140 mg/dl is considered normal.
- A post-prandial plasma glucose level between 140-199 mg/dl is considered as glucose intolerant or pre diabetic. A fasting and post-prandial blood sugar test (after consumption of 75 gm of glucose) is recommended for all such patients.
- · A post-prandial plasma glucose level of above 200 mg/dl is highly suggestive of a diabetic state. A repeat post-prandial test is strongly recommended for all such patients. A post-prandial plasma glucose level in excess of 200 mg/dl on both the occasions is confirmatory of a diabetic state.

Blood Urea Nitrogen (BUN), serum Method : Calculated Serum Creatinine Method : Jaffe kinetic

7.92 0.82

7.8 - 20.2

mg/dl

0.7 - 1.2

Serum Uric Acid Method: Uricase-Peroxidase

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6.16

mg/dl

3.6 - 8.2

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Value	Unit	Biological Reference Interval
	Value	Value Unit

HbA1c (Glycosylated haemoglobin), EDTA whole blood 5.80

< 5.7

Method : HPLC

Estimated average plasma Glucose Method : Calculated

119.76

mg/dL

65 - 136

The test is approved by NGSP for patient sample testing.

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Metabolically normal patients	%	< 5.7
Pre-diabetic	%	5.7 - 6.4
Diabetic	%	> 6.4

Glycosylated hemoglobin or HbA1C is a reliable indicator of mean plasma glucose levels for a period of 8-12 weeks preceeding the date on which the test is performed and is a more reliable indicator of overall blood sugar control in known diabetic patients than blood sugar levels. A value of less than 5.7 % is usually seen in metabolically normal patients, however diabetics with very good control can also yield similar values. The HbA1c test, thus can not be used to differentiate between diabetic patients with very good control over the plasma glucose levels from metabolically normal, non-diabetic subjects as both groups may reveal very similar values in the assay.



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Test Name	Value	Unit	Biological Reference Interval
LFT (Liver Function Test)			
Serum Bilirubin Total Method : Diazotized Sulfanilic Acid (DSA)	0.71	mg/dl	0.1 - 1.2
Serum Bilirubin Direct Method: Diazotized Sulfanilic Acid (DSA)	0.21	mg/dl	0.0 - 0.3
Serum Bilirubin Indirect Method : Calculated	0.50	mg/dl	0.1 - 1.1
Serum SGOT/AST Method : IFCC without P5P	104.10	U/I	<= 35.0
Serum SGPT/ALT Method : IFCC without P5P	168.80	U/I	<= 45.0
Serum Alkaline Phosphatase Method : PNP. AMP Buffer	148.30	U/I	30.0 - 120.0
Serum GGT (Gamma Glutamyl Transpeptidase) Method : UV-assay according to Szasz	27.80	U/I	11.0 - 61.0
Serum total Protein Method : Biuret	8.21	g/dl	6.6 - 8.3
Serum Albumin Method: Bromo Cresol Green	4.80	g/dl	3.5 - 5.2
Serum Globulin Method : Calculated	3.41	g/dl	2.0 - 3.5
Albumin / Globulin ratio Method : Calculated	1.41		1.5 - 2.5

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Test Name		Value	Unit	Biological Reference Interval
Lipid Profile basic (direct H	DL,calculated L	DL)		
Total Cholesterol, , serum Method : CHOD-POD		216.50	mg/dl	< 200.0
Triglycerides , serum Method : GPO-POD		127.50	mg/dl	< 150
HDL Cholesterol , serum Method : Direct measure PEG (CHE-CH	1 O)	44.20	mg/dl	> 40
/LDL Cholesterol , serum Method : Calculated	,	25.50	mg/dl	< 30
.D.L Cholesterol , serum Method : Calculated		146.80	mg/dl	< 100
Cholesterol, Non HDL , serum Method : Calculated		172.30	mg/dl	< 130
Fotal Cholesterol / HDL Cholester Method : Calculated	ol Ratio <i>, serum</i>	4.90		< 5.0
LDL / HDL Cholesterol ratio , seru Method : Calculated	m	3.32		< 3.5
Interpretation:				
National Lipid Association Recommend	ation (NLA-2014)		E-20	
Total Cholesterol Desirable: <200 mg/dL Borderline high: 200-239 mg/dL High: > or =240 mg/dL	Triglycerides Normal: <150 m Borderline high High: 200-499 m Very high: > or	: 150-199 mg/dL mg/dL		
Non HDL Cholesterol Desirable: <130 mg/dL Borderline high: 130-159 mg/dL High: 160-189 mg/dL Very high: > or =190 mg/dL	LDL Choleste Optimal: <100: Near Optimal: Borderline high High: 160-189: Very high: > or	mg/dL 100-129 mg/dL : 130-159 mg/dL mg/dL		
HDL Cholesterol Low (Men) <40 mg/dL Low (Women) <50 mg/dL		2		

Method : Phosphomolybdate Method

Eighty-eight percent of the phosphorus contained in the body is localized in bone in the form of hydroxyapatite. The remainder is involved in intermediary carbohydrate metabolism and in physiologically important substances such as phospholipids, nucleic acids, and adenosine triphosphate (ATP). Phosphorus occurs in blood in the form of inorganic phosphate and organically bound phosphoric acid. The small amount of extracellular organic phosphorus is found exclusively in the form of phospholipids. Serum phosphate concentrations are dependent on meals and variation in the secretion of hormones such as parathyroid hormone (PTH) and may vary widely.

Hypophosphatemia may have 4 general causes: shift of phosphate from extracellular to intracellular, renal phosphate wasting, loss from the gastrointestinal tract, and loss

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from intracellular stores.

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Hyperphosphatemia is usually secondary to an inability of the kidneys to excrete phosphate. Other factors may relate to increased intake or a shift of phosphate from the tissues into the extracellular fluid.

Phosphate levels may be used in the diagnosis and management of a variety of disorders including bone, parathyroid and renal disease.

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Hypophosphatemia is relatively common in hospitalized patients. Levels less than 1.5 mg/dL may result in muscle weakness, hemolysis of red cells, coma, and bone deformity and impaired bone growth.

The most acute problem associated with rapid elevations of serum phosphate levels is hypocalcemia with tetany, seizures, and hypotension. Soft tissue calcification is also an important long-term effect of high phosphorus levels.

Phosphorus levels less than 1.0 mg/dL are potentially life-threatening and are considered a critical value.

Note: Phosphorus has a very strong biphasic circadian rhythm. Values are lowest in the morning, peak first in the late afternoon and peak again in the late evening. The second peak is quite elevated and results may be outside the reference range.



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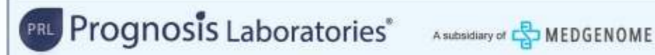
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Test Name Value Unit Biological Reference Interval
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01200230

Vitamin B 12, serum Method: CLIA Microparticles

183.0 - 822.0 pg/ml

Please note change in biological reference interval.

Vitamin B12 (cobalamin) is necessary for hematopoiesis and normal neuronal function. In humans, it is obtained only from animal proteins and requires intrinsic factor (IF) for absorption. The body uses its vitamin B12 stores very economically, reabsorbing vitamin B12 from the ileum and returning it to the liver; very little is excreted.

Vitamin B12 deficiency may be due to lack of IF secretion by gastric mucosa (eg, gastrectomy, gastric atrophy) or intestinal malabsorption (eg, ileal resection, small intestinal diseases).

Vitamin B12 deficiency frequently causes macrocytic anemia, glossitis, peripheral neuropathy, weakness, hyperreflexia, ataxia, loss of proprioception, poor coordination, and affective behavioral changes. These manifestations may occur in any combination; many patients have the neurologic defects without macrocytic anemia. Serum methylmalonic acid and homocysteine levels are also elevated in vitamin B12 deficiency states.

Follow-up testing for antibodies to intrinsic factor (IF) is recommended to identify this potential cause of vitamin B12 malabsorption.

A normal serum concentration of vitamin B12 does not rule out tissue deficiency of vitamin B12. The most sensitive test for vitamin B12 deficiency at the cellular level is the assay for MMA. If clinical symptoms suggest deficiency, measurement of MMA and homocysteine should be considered, even if serum vitamin B12 concentrations are

The commonest cause of increased level of vitamin B12 is therapeutic intake of vitamin B12 in the form of multivitamin tablets or as intramuscular injections.

Many other conditions are known to cause an increase or decrease in the serum vitamin B12 concentration including:

Increased Serum B12	Decreased Serum B12	
Ingestion of vitamin C	Pregnancy	
Ingestion of estrogens	Aspirin	
Ingestion of vitamin A	Anticonvulsants	
Hepatocellular injury	Colchicine	
Myeloproliferative disorder	Ethanol ingestion	
Uremia	Contraceptive hormones	
	Smoking	
	Hemodialysis	
	Multiple myeloma	

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Test Name	Value	Unit	Biological Reference Interval	
Vitamin D (25 Hydroxy), serum Method: CLIA Microparticles	21.72	ng/ml	30.0 - 100.0	

Interpretation:

Deficiency	ng/ml	< 20
Insufficiency	ng/ml	21 - 29
Sufficiency	ng/ml	30 - 100
Intoxication	ng/ml	> 150

Vitamin D compounds are derived from dietary ergocalciferol (from plants, VitD2) or cholecalciferol (from animals, VitD3), or by conversion of 7-dihydrocholesterol to VitD3 in the skin upon ultraviolet exposure. VitD2 and VitD3 are subsequently 25-hydroxylated in the liver to 25-OH-VitD. 25-OH-VitD represents the main body reservoir and transport form of vitamin D, being stored in adipose tissue and tightly bound by a transport protein while in circulation. A fraction of circulating 25-OH-VitD is converted to its active metabolites 1,25-dihydroxy vitamin D2 and D3 (1,25-OH-VitD), mainly by the kidneys. This process is regulated by parathyroid hormone (PTH). VitD plays a primary role in the maintenance of calcium homeostasis. It promotes intestinal calcium absorption and, in concert with PTH, skeletal calcium deposition, or less commonly, calcium mobilization. Renal calcium and phosphate reabsorption are also promoted. In addition to its effects on calcium and bone metabolism, 1,25-OH-VitD regulates the expression of a multitude of genes in many other tissues including immune cells, muscle, vasculature, and reproductive organs.

The exact 25-OH-VitD level reflecting optimal body stores remains unknown. Mild-to-modest deficiency can be associated with osteoporosis or secondary hyperparathyroidism. Severe deficiency may lead to failure to mineralize newly formed osteoid in bone, resulting in rickets in children and osteomalacia in adults. The consequences of vitamin D deficiency on organs other than bone are not fully known, but may include increased susceptibility to infections, muscular discomfort, and an increased risk of colon, breast, and prostate cancer.

Reasons for suboptimal 25-OH-VitD levels include lack of sunshine exposure, a particular problem in India; inadequate intake; malabsorption (eg. due to Celiac disease); depressed hepatic vitamin D 25-hydroxylase activity, secondary to advanced liver disease; and enzyme-inducing drugs, in particular many antiepileptic drugs, including phenytoin, phenobarbital, and carbamazepine, that increase 25-OH-VitD metabolism.

Hypervitaminosis D is rare, and is only seen after prolonged exposure to extremely high doses of vitamin D. When it occurs, it can result in severe hypercalcemia and

Caution: Replacement therapy in deficient individuals must be monitored by periodic assessment of Vitamin D levels in order to prevent hypervitaminosis D.

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Test Name	Value	Unit	Biological Reference Interval	
DCA Total	0.20	ng/ml	0 2 1	

Interpretation:

Method : ECLIA

Prostate-specific antigen (PSA) is a glycoprotein that is produced by the prostate gland, the lining of the urethra, and the bulbourethral gland. Normally, very little PSA is secreted in the blood. Increases in glandular size and tissue damage caused by benign prostatic hypertrophy, prostatitis, or prostate cancer may increase circulating PSA

In patients with previously diagnosed prostate cancer, PSA testing is advocated as an early indicator of tumor recurrence and as an indicator of response to therapy.

The test is also useful for initial screening for prostate cancer:

Total PSA levels < 2 ng/ml almost rule out the possibility of prostatic malignancy.

Total PSA levels between 2 and 10 ng/ml lie in the grey zone. Such values may be obtained in prostatitis, benign hyperplasia and malignancy. Further testing including a free PSA/PSA ratio and prostate biopsy is recommended for these patients for confirmation of the diagnosis.

Total PSA values >10 ng/ml are highly suspicious for prostate cancer but further testing, such as prostate biopsy, is needed to diagnose the exact pathology.



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Test Name	Value	Unit	Biological Reference Interval
Thyroid Profile Total (T3, T4, TSH)			
T3, (Triiodothyronine) , serum Method : ECLIA	1.07	ng/mL	0.80 - 2.0
T4, (Thyroxine) , serum Method : ECLIA	6.55	ug/dL	5.1 - 14.1
TSH (Thyroid Stimulating Hormone) , serum Method : ECLIA	1.29	uIU/mI	0.27 - 4.2

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Interpretation:

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· Primary hyperthyroidism is accompanied by elevated serum T3 and T4 values alongwith depressed TSH levels

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- Primary hypothyroidism is accompanied by depressed serum T3 and T4 values and elevated serum TSH levels.
- High T3 levels coupled with normal T4 and suppressed TSH may be seen in T3 toxicosis.

Note: Total T3 and total T4 are highly bound to plasma proteins and are amenable to fluctuations with plasma protein content as well as due to binding defects in the thyroid hormone binding proteins.

The following ranges are recommended for pregnant females:

Gestation period	TSH (uIU/ml)
First trimester	0.1 - 2.5
Second trimester	0.2 - 3.0
Third trimester	0.3 - 3.0

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Negative

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Test Name	Value	Unit	Biological Reference Interval

Urine Routine & Microscopic Examination

Physical examination

Volume	40 m	nL
Colour	Pale Yellow	Pale yellow
Transparency	Clear	Clear
Specific gravity Method: pKa change	1.015	1.003 - 1.035

<u>Onemical examination</u>		
Protein Method: error-of-indicator	Nil	Nil
Glucose Method: GOD-POD	Nil	Nil
pH Method : Double indicator	5.0	

Negative Bilirubin Negative Method: Azo-coupling reaction Urobilinogen Normal Normal

Method: Azo- coupling reaction Negative Negative Ketone Method : Legals test

Erythrocytes Absent Absent Method: Peroxidase

Negative

Method: Griess reaction Leu/uL Absent Negative Method: Esterase activity of granulocytes

Microscopic examination

Method : Light microscopy

MBC	0 - 1	/ HPF	0 - 2
RBC	Nil	/ HPF	0 - 2
Casts	Nil	/ HPF	Nil
Crystals	Nil	/ HPF	Nil
Epithelial cells	0 - 1	/ HPF	0 - 15
Bacteria	Absent		Absent
Others	Nil		

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Nitrite





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Test Name	Value	Unit	Biological Reference Interval	
Urine Sugar fasting Method : Hexokinase	Nil		Nil	
Urine Sugar PP Method : Hexokinase	NIL		NIL	

01200230



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Mobile:9313817732

Dr. Deepak Sadwani MD Pathology

Dr. Mayank Gupta MD, DNB Pathology Consultant Pathologist Dr. Moushmi Mukherjee MD Pathology

Mousheei Mukkaezee

Consultant Pathologist

Page 13 of 18

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Lab No. 012411200230

Mr. JAGDISH

Ref. Dr. **MEDIWEEL** Rpt. Centre Courier

NAME

Mobile:9313817732

Age/ Gender 54 YRS/MALE

01200230

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20/Nov/2024

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20/Nov/2024 06:52PM

ECG Electro-cardiography Normal ECG.



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Dr. Smita Sadwani

MBBS. MD Director DMC Regd. No. 48732

Dr. Mukesh Sharma MD(Microbiology) Consultant Microbiologist Lab Director

Dr. Deepak Sadwani Dr. Ashish Gautam MD(Pathology)

MD, PGDCC

Dr. Moushmi Mukherjee MBBS,MD (Pathology) Consultant Cardiologist Consultant Pathologist

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Lab No. 012411200230

Mr. JAGDISH

Courier

Ref. Dr. **MEDIWEEL**

NAME

Rpt. Centre

Age/ Gender 54 YRS/MALE

01200230

BarcodeNo

Coll. ON

20/Nov/2024 09:16AM

Reg. ON

20/Nov/2024

Approved ON 20/Nov/2024 01:38PM

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TMT (Treadmill Test)

Negative For RMI.



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Dr. Anil Sahoo MD. PGDCO Reg. No.33201

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Lab No. 012411200230

Courier

Age/ Gender 54 YRS/MALE Coll. ON 20/Nov/2024 09:16AM

Reg. ON

20/Nov/2024

NAME Mr. JAGDISH Ref. Dr.

Rpt. Centre

MEDIWEEL BarcodeNo 01200230

Approved ON 20/Nov/2024 01:35PM

Printed ON

20/Nov/2024 06:52PM

Eye Vision									
	Right Eye	Left Eye							
NEAR VISION	NI/6 (Mith Class)	N/6 (With							
INDAK VISION	N/6 (With Glass)	Glass)							
DISTANCE	6/6 (With Glass)	6/6 (With Glace)							
VISION	0/0 (With Glass)	6/6 (With Glass)							
COLOR VISION	Normal	Normal							

MER

Mobile:9313817732

General	Fair, no pallor, no icterus, no anemia
Condition	observed
Height (cm)	178
Weight (kg)	80
Pulse (bpm)	73
BP (mm/hg)	147/88

Please note: Kindly review with clinician in view of abnormal reports (if any).

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Tr. Dr. Smita Sadwani MBBS. MD

Director

Dr. Mukesh Sharma MD(Microbiology) Consultant Microbiologist Lab Director

Dr. Deepak Sadwani Dr. Ashish Gautam MD(Pathology)

MD, PGDCC

Dr. Moushmi Mukherjee MBBS,MD (Pathology) Consultant Cardiologist Consultant Pathologist

DMC Regd. No. 48732

Page 16 of 18





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Lab No. 012411200230

NAME

Ref. Dr.

Rpt. Centre

Mr. JAGDISH

MEDIWEEL Courier

Age/ Gender 54 YRS/MALE

01200230

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20/Nov/2024

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20/Nov/2024 06:52PM

X-Ray Chest PA view

Trachea and mediastinum are central.

Bilateral lung fields are clear.

Bilateral hilar shadows are normal.

Bilateral costophrenic angles are clear.

Cardiac shadow is normal.

Soft tissue shadows and bony rib cage is normal.

Impression: No significant abnormality seen.

Please correlate clinically

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DR AMIT JAISWAL MBBS,DMRD.DNB (RADIO DIAGNOSIS) Page 17 of 18 DMC No. 55709





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Lab No. 012411200230

Mr. JAGDISH

Courier

Ref. Dr. **MEDIWEEL**

NAME

Rpt. Centre

Age/ Gender 54 YRS/MALE

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20/Nov/2024

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SONOGRAPHY OF ABDOMEN AND PELVIS

The liver is normal in size (13.6 cm) and shows mild diffuse increased parenchymal echogenicity. There is no evidence of any focal hepatic lesion. The hepatic and portal veins are normal. There is no intrahepatic biliary dilatation.

The gall bladder is adequately distended. There is no evidence of any calculi. There is no evidence of any wall thickening seen. The CBD is not dilated.

The pancreas is well visualized and shows a normal parenchymal echotexture. There is no evidence of any focal mass, calcification or ductal dilatation seen. There is no peripancreatic fluid collection seen.

The spleen is normal in size (9.3 cm) and shows a normal parenchymal echotexture. There is no focal lesion seen.

The right kidney measures 11.2 x 4.4 cm and the left kidney measures 11.3 x 5.1 cm. Both kidneys are normal in size and shape. The kidneys show normal echotexture with a well-maintained cortical thickness. There is no evidence of hydronephrosis, cortical scarring or calculus disease in right kidney.

Left kidney shows few simple cortical cysts largest of them measuring 18 x 14 mm at interpolar region.

There is no ascites or bowel wall thickening.

The urinary bladder shows normal contours.

The prostate is normal in size.

IMPRESSION

• Grade I fatty liver.

Kindly correlate clinically.

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*** Partial Report ***



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A SUBSIDIARY OF MEDGENOME 515-516 DWARKA SEC19 NEW DELHI 110075

Mr. JAGDISH

I.D. : 361 AGE/SEX: 54 Yr/M HT/WT : /

DATE : 20-11-2024 10:43:43 AM

REF.BY: Dr.MEDIWEEL

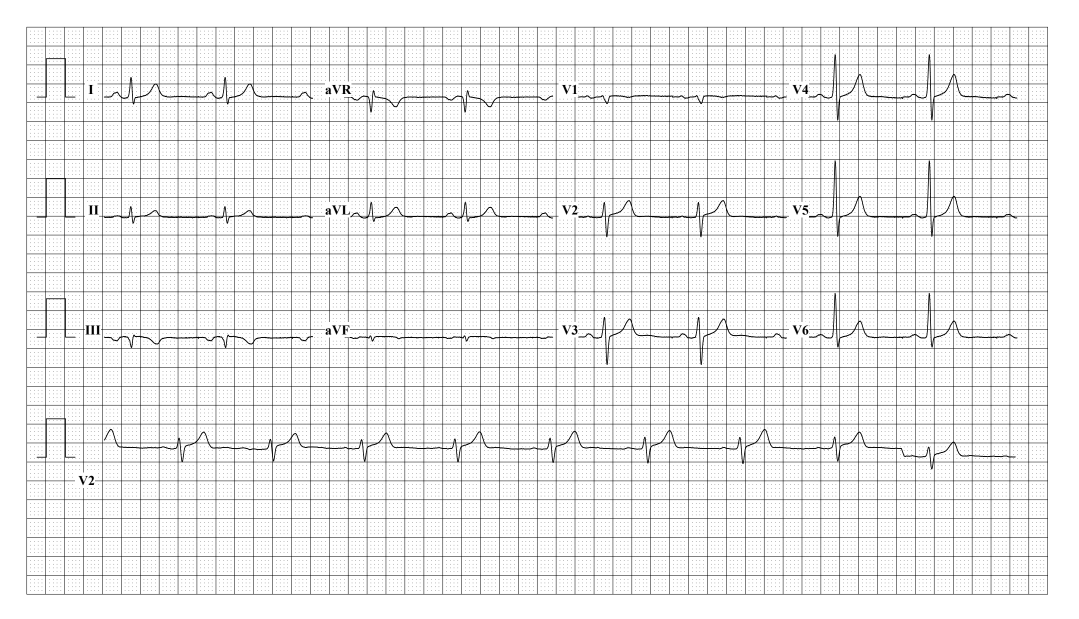
MACHINE INTERPRETATION: Normal ECG.

RATE : 61 bpm P Duration : 125 ms PR Duration : 168 ms BP : N/A

P Axis : -5 deg. QRS Duration: 85 ms QRS Axis : -8 deg. QT Interval : 350 ms

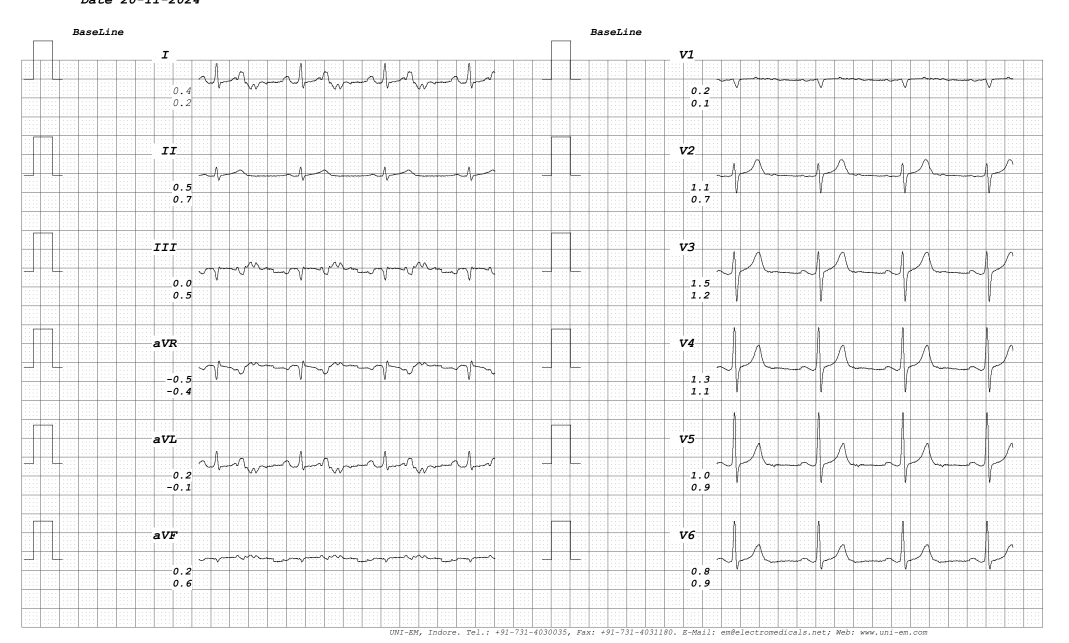
T Axis : -1 deg. QTc Interval : 354 ms **Linked Median**

Speed: 25 mm/s Sensitivity: 10 mm/mV



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

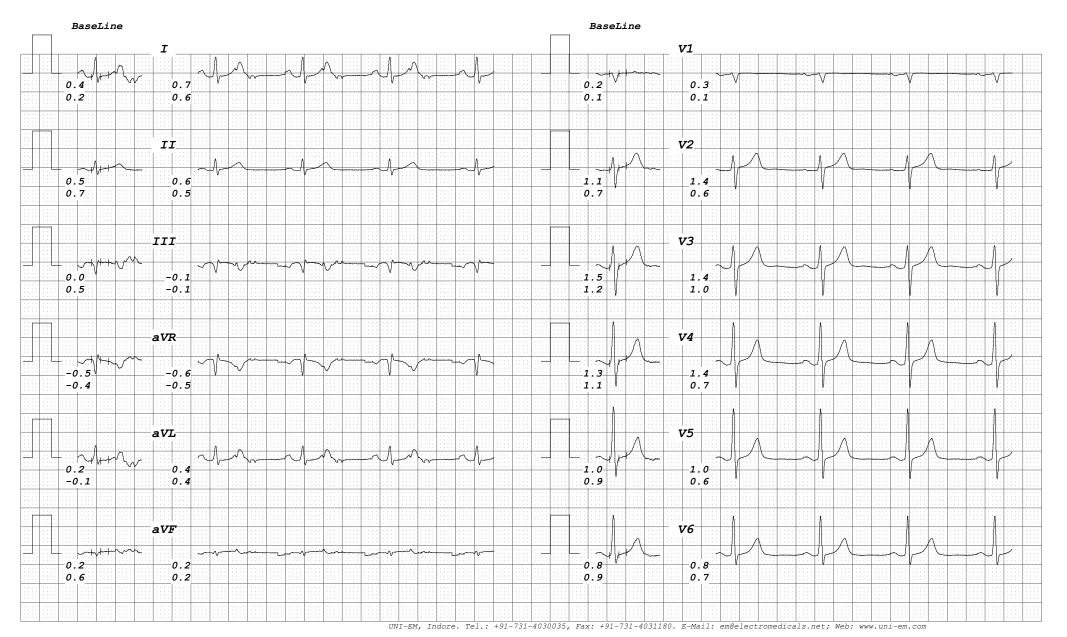
RATE 68bpm B.P. 140/88 PRETEST SUPINE ST @ 10mm/mV 80ms PostJ



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

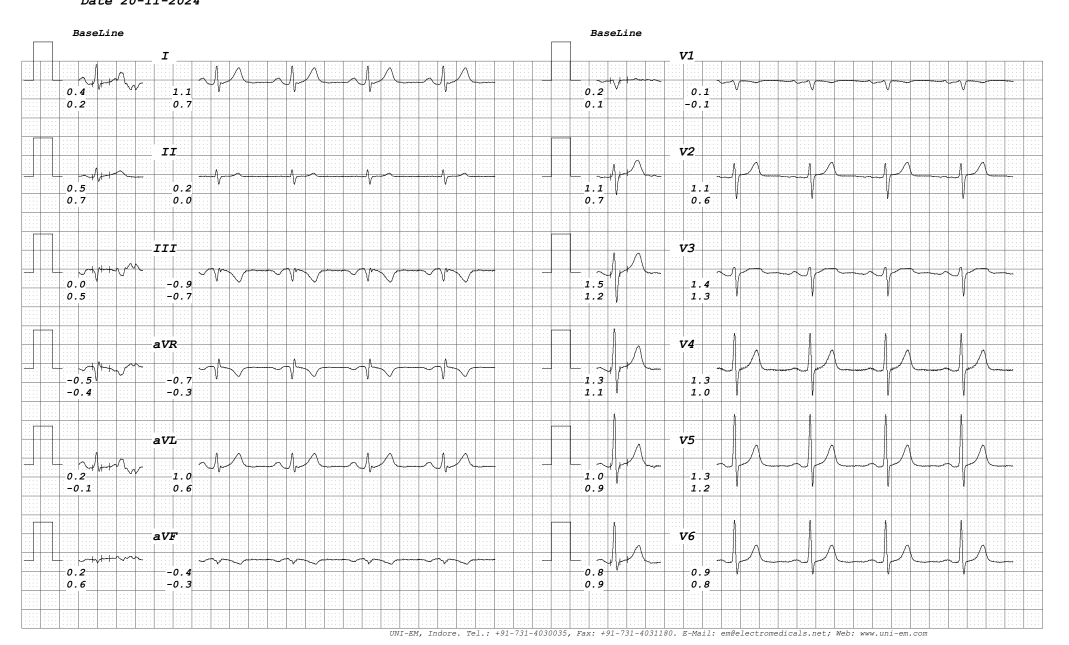
RATE 66bpm B.P. 140/88 PRETEST HYPERVENT ST @ 10mm/mV 80ms PostJ

PHASE TIME 0:16



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

RATE 76bpm B.P. 140/88 PRETEST STANDING ST @ 10mm/mV 80ms PostJ

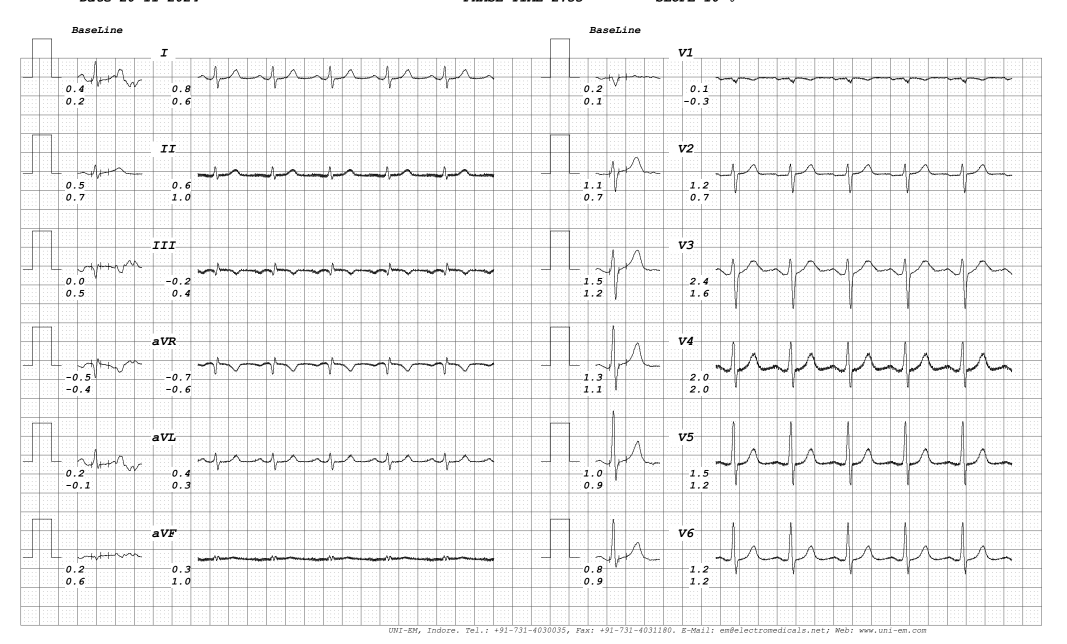


JAGDISH I.D. 583 Age 54/M Date 20-11-2024

RATE 100bpm B.P. 142/90 Bruce Stage 1 TOTAL TIME 2:55 PHASE TIME 2:55

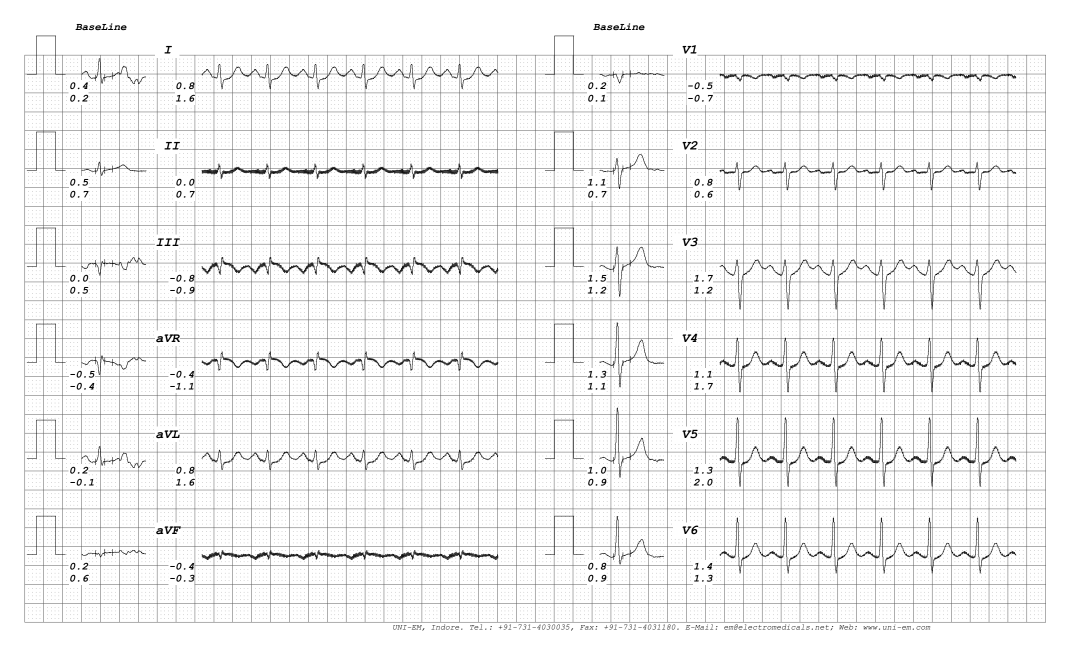
80ms PostJ Speed 2.7 km/hr SLOPE 10 %

ST @ 10mm/mV



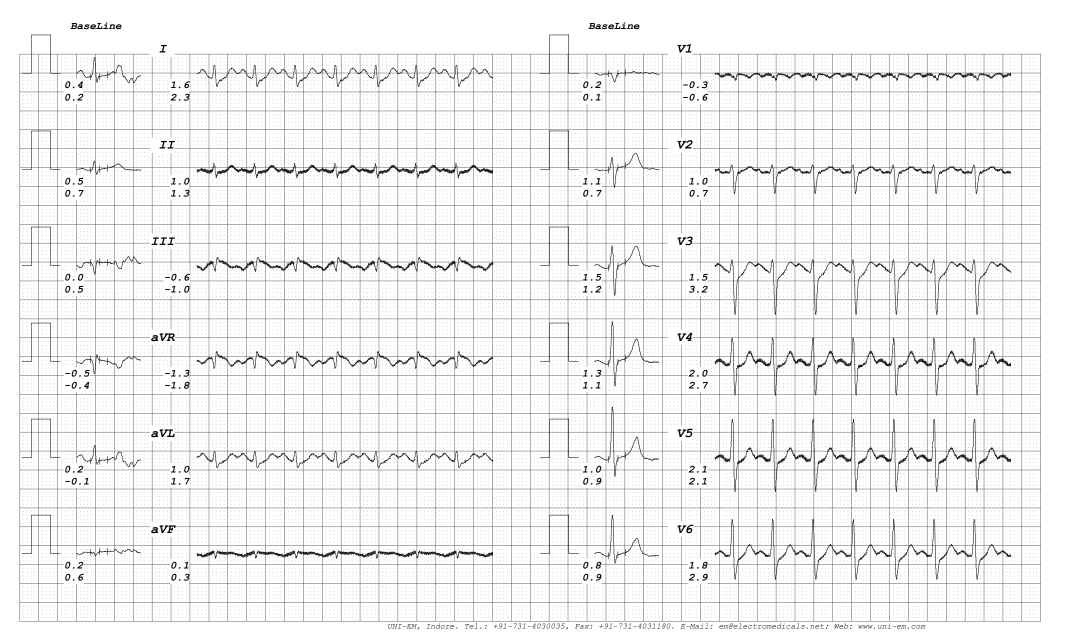
JAGDISH I.D. 583 Age 54/M Date 20-11-2024

RATE 120bpm B.P. 144/92 Bruce Stage 2 TOTAL TIME 5:55 PHASE TIME 2:55 ST @ 10mm/mV 80ms PostJ Speed 4 km/hr SLOPE 12 %



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

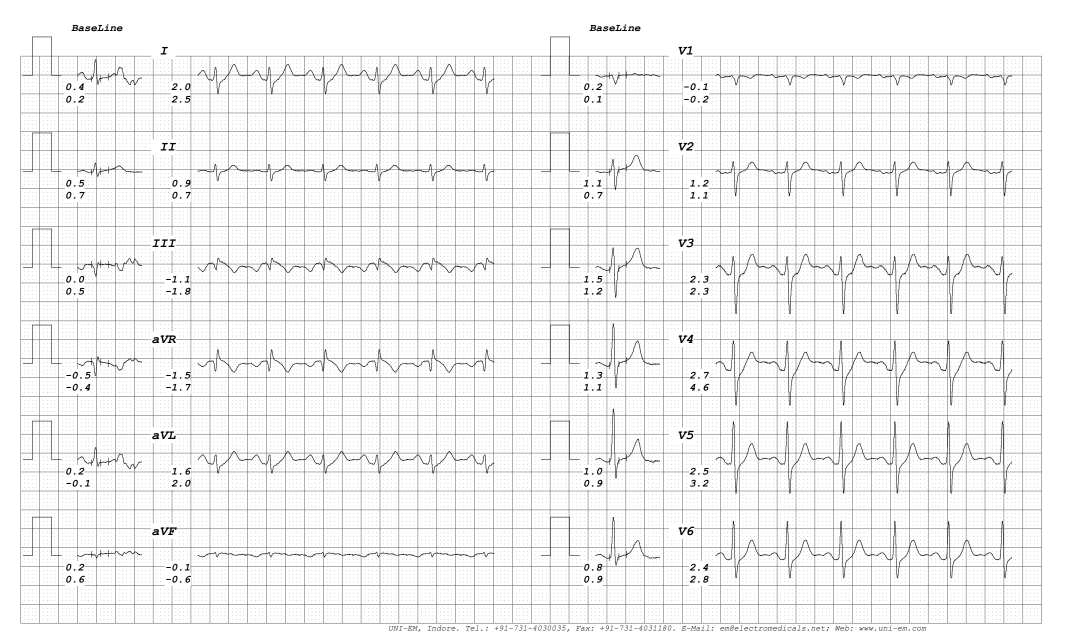
RATE 142bpm B.P. 146/94 Bruce PK-EXERCISE TOTAL TIME 7:47 PHASE TIME 1:47 ST @ 10mm/mV 80ms PostJ Speed 5.4 km/hr SLOPE 14 %



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

RATE 107bpm B.P. 144/92 Bruce RECOVERY TOTAL TIME 8:57 ST @ 10mm/mV 80ms PostJ

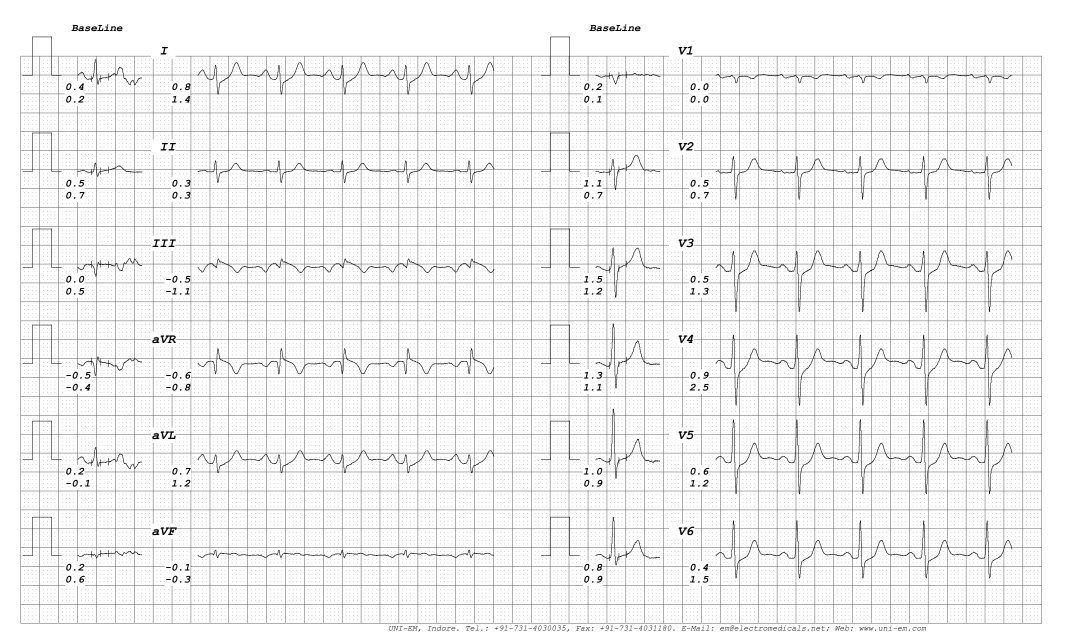
PHASE TIME 0:55



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

RATE 90bpm B.P. 140/88 Bruce RECOVERY TOTAL TIME 9:57 ST @ 10mm/mV 80ms PostJ

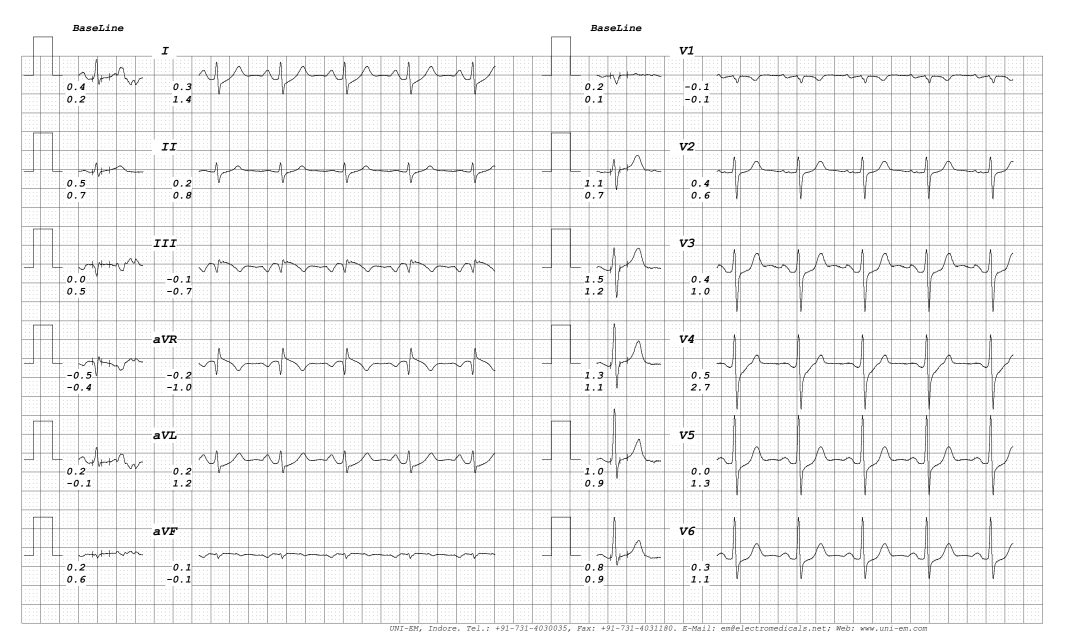
TOTAL TIME 9:57
PHASE TIME 1:55



JAGDISH I.D. 583 Age 54/M Date 20-11-2024

RATE 90bpm B.P. 140/88 Bruce RECOVERY TOTAL TIME 10:57 ST @ 10mm/mV 80ms PostJ

PHASE TIME 2:55



A SUBSIDIARY OF MEDGENOME 515-516 DWARKA SEC 19 NEW DELHI 110075

JAGDISH TREADMILL TEST REPORT

ID : 583

DATE : 20-11-2024 PROTOCOL : Bruce

AGE/SEX: 54 /M HISTORY: HT/WT: 0 / 0 INDICATION: REF.BY: MEDIWEEL MEDICATION:

PHASE	TOTAL TIME	STAGE TIME	SPEED Km/Hr	GRADE %	H.R.	B.P.		ST LEVEL(MM)		METS	
	TIME	TIME	MIII/ HI	6	bpm	mmHg	XIOO	II	V1	V5	
SUPINE HYPERVENT STANDING Stage 1 Stage 2 PK-EXERCISE RECOVERY RECOVERY RECOVERY	2:55 5:55 7:47 8:57 9:57 10:57	0:16 2:55 2:55 1:47 0:55 1:55 2:55	2.7 4 5.4	10 12 14	68 66 76 100 120 142 107 90	140 / 88 140 / 88 140 / 88 142 / 90 144 / 92 146 / 94 144 / 92 140 / 88	92 106 142 172 207 154 126	0.5 0.6 0.2 0.6 0 1 0.9 0.3	0.2 0.3 0.1 0.1 -0.5 -0.3 -0.1	1 1 1.3 1.5 1.3 2.1 2.5 0.6	4.67 7.04 8.82

RESULTS

EXERCISE DURATION : 7:47 MAX WORK LOAD : 8.82 METS

MAX HEART RATE : 142 bpm 85 % of target heart rate 166 bpm

MAX BLOOD PRESSURE : 146 / 94 mm Hg REASON OF TERMINATION : Achieved THR,

BP RESPONSE
ARRYTHMIA
H.R. RESPONSE
IMPRESSIONS



भारतीय विशिष्ट पहचान प्राधिकरण भारत सरकार



Linique Kleofilicanon Authority of Inche Government of India

E-ABOURS LEDS

अपमानन सम्पन्धEnrolatert No. 1452/18694/90122

Jagdish Prasad Yadav (अगरील प्रमाद बादन)

RZF-907/1 S/F, M.G MARG Raj Nagar-2, Bagdola. South West Delhi.

Deitri - 110077

STITE SEETS WHEN Your Andhear No.:

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आधार-आम आदमी का अधिकार

Hot da nat

- अपार के लिए अलगर एक है। इस नामांश्रेस इस कर बाँग की आवश्याकता है.
- 🔊 प्रमुक्त अञ्चल नुसीलगण मीबाइल नेबर तथा है सल प्रण सुर्ज कराएँ, इसमें जापको ब्रिमिध सुविधाएँ धाम कर में महक्तियन

机制型

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- 🚁 पर राजका प्रमाणा जानलाहम अभिन्दिर्गान द्वारा प्राप्त संग्र
- महाभक्त इलेक्ट्रोनिक प्रक्रिका द्वारा ध्रमा १७० ५% है।

INFORMATION

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भारत धरकार



अगदील प्रसाद गावक Jegdish Prasad Yadav men (vita) DOB: 02/01/1970 HER / MALE



भूगतीय विशिष्ट प्रदेशीन गामिकरण CHIES THE EPPARTURA MODERNIA OF THE STATE OF

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