DIAGNOSTICS REPORT

Patient Name	: Mr. IVL SRIDHAR	Order Date	: 30/03/2023 08:42
Age/Sex	: 55 Year(s)/Male	Report Date	: 30/03/2023 12:41
UHID	: SHHM.61755	IP No	:
Ref. Doctor	: Self	Facility	: SEVENHILLS HOSPITAL, MUMBAI

2D ECHOCARDIOGRAPHY WITH COLOUR DOPPLER STUDY

Fair LV and normal RVsystolic function.
Estimated LVEF = 50%
Dyssynchronous septal motion consistent with LBBB.
All valves are structurally and functionally normal.
Normal sized cardiac chambers.
Grade I LV Diastolic dysfunction .
No pulmonary arterial hypertension.
No regurgitation across any other valves.
Normal forward flow velocities across all the cardiac valves.
Aorta and pulmonary artery dimensions: normal.
IAS / IVS: Intact.
No evidence of clot, vegetation, calcification, pericardial effusion.
COLOUR DOPPLER: NO MR/AR.



Dr.Jayashree Dash,

(Junior Consultant NIC) RegNo: 3393/09/2003

Patient Name: Mr. IVL SRIDHARUHID: SHHM.61755Episode: OPRef. Doctor: Self

Age/Sex	: 55 Year(s) / Male
Order Date	: 30/03/2023 08:42
Mobile No	: 9035281468
DOB	: 12/06/1967
Facility	: SEVENHILLS HOSPITAL, MUMBAI

Blood Bank

 Test Name
 Result

 Sample No :
 00264832A
 Collection Date :
 30/03/23
 08:44
 Ack Date :
 30/03/2023
 11:57
 Report Date :
 30/03/23
 13:24

BLOOD GROUPING/ CROSS-MATCHING BY SEMI AUTOMATION

BLOOD GROUP (ABO)

UP (ABO) ' A ' POSITIVE

Method - Column Agglutination

REMARK: THE REPORTED RESULTS PERTAIN TO THE SAMPLE RECEIVED AT THE BLOOD CENTRE.

Interpretation:

Rh Type

Blood typing is used to determine an individual's blood group, to establish whether a person is blood group A, B, AB, or O and whether he or she is Rh positive or Rh negative. Blood typing has the following significance,

• Ensure compatibility between the blood type of a person who requires a transfusion of blood or blood components and the ABO and Rh type of the unit of blood that will be transfused.

• Determine compatibility between a pregnant woman and her developing baby (fetus). Rh typing is especially important during pregnancy because a mother and her fetus could be incompatible.

• Determine the blood group of potential blood donors at a collection facility.

• Determine the blood group of potential donors and recipients of organs, tissues, or bone marrow, as part of a workup for a transplant procedure.

End of Report

Dr.Ritesh Kharche MD, PGD HOD, Laboratory Medicine Dept.

RegNo: 2006/03/1680

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Patient Name	: Mr. IVL SRIDHAR	Age/Sex	: 55 Year(s) / Male
UHID	: SHHM.61755	Order Date	: 30/03/2023 08:42
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Ref. Doctor	: Self	Mobile No	: 9035281468
		DOB	: 12/06/1967
		Facility	: SEVENHILLS HOSPITAL, MUMBAI

Biochemistry									
est Name			Result			Unit	Ref.	Range	
Sample No :	O0264832A	Collection Date :	30/03/23 08:44	Ack Date :	30/03/2023 10:45	Repo	rt Date :	30/03/23	12:20
GLYCOSLY	ATED HAEMOGI	LOBIN (HBA1C)						
HbA1c			6.78	•		%		4 to 6% Non-diab 6.07	
Method - Calco NOTES :- 1. HbA1c is us 2. HbA1c n evaluates diat 3. Inapprop hypertrigiycer interference w 4. HbA1c may 5. Inapprop hyperbilirubine 6. Trends in H 7. Any sam below 4% sho 8. HbA1c targ 9. HbA1c targ	ed for monitoring diaba nay be falsely low petes over 15 days. priately low HbA1 ridemia, chronic l with estimation of HbA1. the increased in patien riately higher value emia and large doses o lbA1c are a better indic pole with >15% Hi puld prompt additional s et in pregnancy is to at et in paediatric age gro	etic control. It reflect in diabetics with ac values may liver disease.Drug c, causing falsely low ts with polycythemia of aspirin. ator of diabetic contr bA1c should be studies to determine tain level <6 % .	s the mean plasma gluce hemolytic disease. I be reported due s like dapsone, values. or post-splenectomy. v be caused due ol than a solitary test. suspected of having the possible presence of < 7.5 %.	n these ind to hen ribavirin, to iron d a hemoglo, variant hemo	lividuals a plasma fi nolysis, recent blou antiretroviral drugs, eficiency, vitamin B1 bin variant, especiall)	od transfus , trimetho 12 deficienc	sion, ac prim, n y, alcoho	rute blood nay also ol intake,	t loss, cause uremia,
		,, ,	r hemolyzed whole blood Medical Care in Diabete						
Sample No :	O0264832B	Collection Date :	30/03/23 08:44	Ack Date :	30/03/2023 10:08	Repo	rt Date :	30/03/23	12:20

GLUCOSE-PLASMA-FASTING

Glucose,Fasting

127.88 🔺

mg/dl 70 - 110

Patient Name : Mr. IVL SRIDHAR

UHID : SHHM.61755 Episode : OP Ref. Doctor : Self

: 55 Year(s) / Male
: 30/03/2023 08:42
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: SEVENHILLS HOSPITAL, MUMBAI

American Diabetes Association Reference Range :

Normal : < 100 mg/dl Impaired fasting glucose(Prediabetes) : 100 - 126 mg/dl Diabetes : >= 126 mg/dl

References:

Pack Insert of Bio system
 Tietz Textbook Of Clinical Chemistry And Molecular Diagnostics, 6th Ed, Editors: Rifai et al. 2018

Interpretation :-

Conditions that can result in an elevated blood glucose level include: Acromegaly, Acute stress (response to trauma, heart attack, and stroke for instance), Chronic kidney disease, Cushing syndrome, Excessive consumption of food, Hyperthyroidism, Pancreatitis.

A low level of glucose may indicate hypoglycemia, a condition characterized by a drop in blood glucose to a level where first it causes nervous system symptoms (sweating, palpitations, hunger, trembling, and anxiety), then begins to affect the brain (causing confusion, hallucinations, blurred vision, and sometimes even coma and death). A low blood glucose level (hypoglycemia) may be

seen with:Adrenal insufficiency, Drinking excessive alcohol, Severe liver disease, Hypopituitarism, Hypothyroidism, Severe infections, Severe heart failure, Chronic kidney (renal) failure, Insulin overdose, Tumors that produce insulin (insulinomas),Starvation.

<u>Lipid Profile</u>			
Total Cholesterol	185.72	mg/dl	Reference Values : Up to 200 m
Triglycerides	135.23	mg/dl	Reference Values: Up to 150 mg
Method - Enzymatic			
HDL Cholesterol	35.28	mg/dl	0 - 60
Method - Enzymatic immuno inhibition			
LDL Cholesterol	123.39	mg/dl	0 - 130
Method - Calculated			
VLDL Cholesterol	27.05	mg/dl	0 - 40
Method - Calculated			
Total Cholesterol / HDL Cholesterol Ratio -	5.26 ▲	RATIO	0 - 5
Calculated			
Method - Calculated			
LDL / HDL Cholesterol Ratio - Calculated	3.50	RATIO	0 - 4.3
Method - Calculated			

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Interpretation

1. Triglycerides: When triglycerides are very high greater than 1000 mg/dL, there is a risk of developing pancreatitis in children and adults. Triglycerides change dramatically in response to meals, increasing as much as 5 to 10 times higher than fasting levels just a few hours after eating. Even fasting levels vary considerably day to day. Therefore, modest changes in fasting triglycerides measured on different days are not considered to be abnormal.

2. HDL-Cholesterol: HDL- C is considered to be beneficial, the so-called "good" cholesterol, because it removes excess cholesterol from tissues and carries it to the liver for disposal. If HDL-C is less than 40 mg/dL for men and less than 50 mg/dL for women, there is an increased risk of heart disease that is independent of other risk factors, including the LDL-C level. The NCEP guidelines suggest that an HDL cholesterol value greater than 60 mg/dL is protective and should be treated as a negative

risk factor.

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3. LDL-Cholesterol: Desired goals for LDL-C levels change based on individual risk factors. For young adults, less than 120 mg/dL is acceptable. Values between 120-159 mg/dL are considered Borderline high. Values greater than 160 mg/dL are considered high. Low levels of LDL cholesterol may be seen in people with an inherited lipoprotein deficiency and in people with hyperthyroidism, infection, inflammation, or cirrhosis.

Uric Acid (Serum)				
Uric Acid	5.6	mg/dl	3.5 - 7.2	
Method - Uricase				
References:				
1)Pack Insert of Bio system				

2) TIETZ Textbook of Clinical chemistry and Molecular DiagnosticsEdited by: Carl A.burtis, Edward R. Ashwood, David e. Bruns

Interpretation:-

Uric acid is produced by the breakdown of purines. Purines are nitrogen-containing compounds found in the cells of the body, including our DNA. Increased concentrations of uric acid can cause crystals to form in the joints, which can lead to the joint inflammation and pain characteristic of gout. Low values can be associated with some kinds of liver or kidney diseases, Fanconi syndrome, exposure to toxic compounds, and rarely as the result of an inherited metabolic defect (Wilson disease).

Liver Function Test (LFT)

SGOT (Aspartate Transaminase) - SERUM	18.97	U/L	0 - 35
Method - IFCC SGPT (Alanine Transaminase) - SERUM	31.11	U/L	0 - 45
<i>Method - IFCC</i> Total Bilirubin - SERUM	0.87	mg/dl	0 - 2
<i>Method - Diazo</i> Direct Bilirubin SERUM	0.34	mg/dl	0 - 0.4
Method - Diazotization			
Indirect Bilirubin - Calculated Method - Calculated	0.53	mg/dl	0.1 - 0.8
Alkaline Phosphatase - SERUM Method - IFCC AMP Buffer	97.99	U/L	0 - 115
Total Protein - SERUM	7.81 🔺	gm/dl	6 - 7.8
<i>Method - Biuret</i> Albumin - SERUM	5.01	gm/dl	3.5 - 5.2
Method - Bromo Cresol Green(BCG)			

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			Facility	: SEVENHILLS HOSPITAL, MUMBAI		
Globulin - Ca	lculated	2.80		gm/dl	2 - 4	
Method - Calcula	ated					
A:G Ratio		1.79		:1	1 - 3	
<i>Method - Calculated</i> Gamma Glutamyl Transferase (GGT) - Gglutamyl						
		19.2		U/L	0 - 55	
carboxy nitro	oanilide - SERUM					
Method - G glut	amyl carboxy nitroanilide					

References:

1)Pack Insert of Bio system

2) Tietz Textbook Of Clinical Chemistry And Molecular Diagnostics, 6th Ed, Editors: Rifai et al. 2018

Interperatation :-

Bilirubin is a yellowish pigment found in bile and is a breakdown product of normal heme catabolism. Elevated levels results from increased bilirubin production (eg hemolysis and ineffective erythropoiesis); decreased bilirubin excretion (eg; obstruction and hepatitis); and abnormal bilirubin metabolism (eg; hereditary and neonatal jaundice).conjugated (direct) bilirubin is also elevated more than unconjugated (indirect) bilirubin when there is some kind of blockage of the bile ducts like in Gallstonesgetting into the bile ducts tumors & Scarring of the bile ducts. Increased unconjugated (indirect) bilirubin may be a result of hemolytic or pernicious anemia, transfusion reaction & a common metabolic condition termed Gilbert syndrome.

AST levels increase in viral hepatitis, blockage of the bile duct ,cirrhosis of the liver, liver cancer, kidney failure, hemolytic anemia, pancreatitis, hemochromatosis.Ast levels may also increase after a heart attck or strenuous activity. ALT is commonly measured as a part of a diagnostic evaluation of hepatocellular injury, to determine liver health. Elevated ALP levels are seen in Biliary Obstruction, Osteoblastic Bone Tumors, Osteomalacia, Hepatitis, Hyperparathyriodism, Leukemia,Lymphoma, paget's disease, Rickets, Sarcoidosis etc.

Elevated serum GGT activity can be found in diseases of the liver, Biliary system and pancreas. Conditions that increase serum GGT are obstructive liver disease, high alcohol consumption and use of enzyme-including drugs etc.

Serum total protein, also known as total protein, is a biochemical test for measuring the total amount of protein in serum..Protein in the plasma is made up of albumin and globulin. Higher-than-normal levels may be due to: Chronic inflammation or infection, including HIV and hepatitis B or C, Multiple myeloma,Waldenstrom's disease. Lower-than-normal levels may be due to: Agammaglobulinemia, Bleeding (hemorrhage), Burns, Glomerulonephritis, Liver disease, Malabsorption, Malnutrition, Nephrotic - Human serum albumin is the most abundant protein in human blood plasma. It is produced in the liver.Albumin constitutes about half of the blood serum protein. Low blood albumin levels (hypoalbuminemia) can be caused by: Liver disease like cirrhosis of the liver, nephrotic syndrome, protein-losing enteropathy, Burns, hemodilution, increased vascular permeability or decreased lymphatic clearance, malnutrition and wasting etc.

Renal Function Test (RFT)

Urea - SERUM	13.27 🔻	mg/dl	15 - 39
Method - Urease			
BUN - SERUM	6.20	mg/dl	4 - 18
Method - Urease-GLDH			
Creatinine - SERUM	0.77	mg/dl	0.5 - 1.3
Method - Jaffes Kinetic			
References:			
1)Pack Insert of Bio system			

2) Tietz Textbook Of Clinical Chemistry And Molecular Diagnostics, 6th Ed, Editors: Rifai et al. 2018

Interpretation:-

The blood urea nitrogen or BUN test is primarily used, along with the creatinine test, to evaluate kidney function in a wide range of circumstances, to help 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.

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GLUCOSE-PLASMA POST PRANDIAL

Glucose, Post Prandial

335.59 🛦

mg/dl

70.00 - 140.00

American Diabetes Association Reference Range :

Post-Prandial Blood Glucose:

Non- Diabetic: Up to 140mg/dL Pre-Diabetic: 140-199 mg/dL Diabetic :>200 mg/dL

References:

1)Pack Insert of Bio system 2) Tietz Textbook Of Clinical Chemistry And Molecular Diagnostics, 6th Ed, Editors: Rifai et al. 2018

Interpretation :-

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seen with:Adrenal insufficiency, Drinking excessive alcohol, Severe liver disease, Hypopituitarism, Hypothyroidism, Severe infections, Severe heart failure, Chronic kidney (renal) failure, Insulin overdose, Tumors that produce insulin (insulinomas),Starvation.

End of Report

Dr.Ritesh Kharche MD, PGD HOD, Laboratory Medicine Dept.

RegNo: 2006/03/1680

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HAEMATOLOGY

Test Name	Result		Unit	Ref. Range
Sample No : 00264832A Collection Date	30/03/23 08:44	Ack Date : 30/03/2023 10:4	5 Report Date	e: 30/03/23 12:20
COMPLETE BLOOD COUNT (CBC) - EDT	A WHOLE BLOOD			
Total WBC Count	6.73		x10^3/ul	4.00 - 10.00
Neutrophils	58		%	40.00 - 80.00
Lymphocytes	34.8		%	20.00 - 40.00
Eosinophils	1.9		%	1.00 - 6.00
Monocytes	5.3		%	2.00 - 10.00
Basophils	0.0	,	%	1.00 - 2.00
Absolute Neutrophils Count	3.91		x10^3/ul	2.00 - 7.00
Absolute Lymphocytes Count	2.34		x10^3/ul	0.80 - 4.00
Absolute Eosinophils Count	0.12		x10^3/ul	0.02 - 0.50
Absolute Monocytes Count	0.36		x10^3/ul	0.12 - 1.20
Absolute Basophils Count	0.00		x10^3/ul	0.00 - 0.10
RBCs	5.28		x10^6/ul	4.50 - 5.50
Hemoglobin	16.3		gm/dl	13.00 - 17.00
Hematocrit	49.0		%	40.00 - 50.00
MCV	92.6		fl	83.00 - 101.00
МСН	30.8		pg	27.00 - 32.00
МСНС	33.2		gm/dl	31.50 - 34.50
RED CELL DISTRIBUTION WIDTH-CV (RDW	/-CV) 12.7		%	11.00 - 16.00
RED CELL DISTRIBUTION WIDTH-SD (RDW	/-SD) 43.1		fl	35.00 - 56.00
Platelet	187		x10^3/ul	150.00 - 410.00
MPV	8.7		fl	6.78 - 13.46
PLATELET DISTRIBUTION WIDTH (PDW)	16.0		%	9.00 - 17.00
PLATELETCRIT (PCT)	0.163	3	%	0.11 - 0.28

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Method:-

HB Colorimetric Method. RBC/PLT Electrical Impedance Method. WBC Flow Cytometry by Laser Method. MCV, MCH, MCHC, RDW - Calculated. Differential Count - Manual.

NOTE: Wallach's Interpretation of Diagnostic Tests. 11th Ed, Editors: Rao LV. 2021

NOTE :-

The International Council for Standardization in Haematology (ICSH) recommends reporting of absolute counts of various WBC subsets for clinical decision making. This test has been performed on a fully automated 5 part differential cell counter which counts over 10,000 WBCs to derive differential counts. A complete blood count is a blood panel that gives information about the cells in a patient's blood, such as the cell count for each cell type and the concentrations of Hemoglobin and platelets. The cells that circulate in the bloodstream are generally divided into three types: white blood cells (leukocytes), red blood cells (erythrocytes), and platelets (thrombocytes). Abnormally high or low counts may be physiological or may indicate disease conditions, and hence need to be interpreted clinically.

ERYTHROCYTE SEDIMENTATION RATE (ESR)

ESR 15 mm/hr 0 - 20

Method: Westergren Method

INTERPRETATION :-

ESR is a non-specific phenomenon, its measurement is clinically useful in disorders associated with an increased production of acute-phase proteins. It provides an index of progress of the disease in rheumatoid arthritis or tuberculosis, and it is of considerable value in diagnosis of temporal arteritis and polymyalgia rheumatica. It is often used if multiple myeloma is suspected, but when the myeloma is non-secretory or light chain, a normal ESR does not exclude this diagnosis.

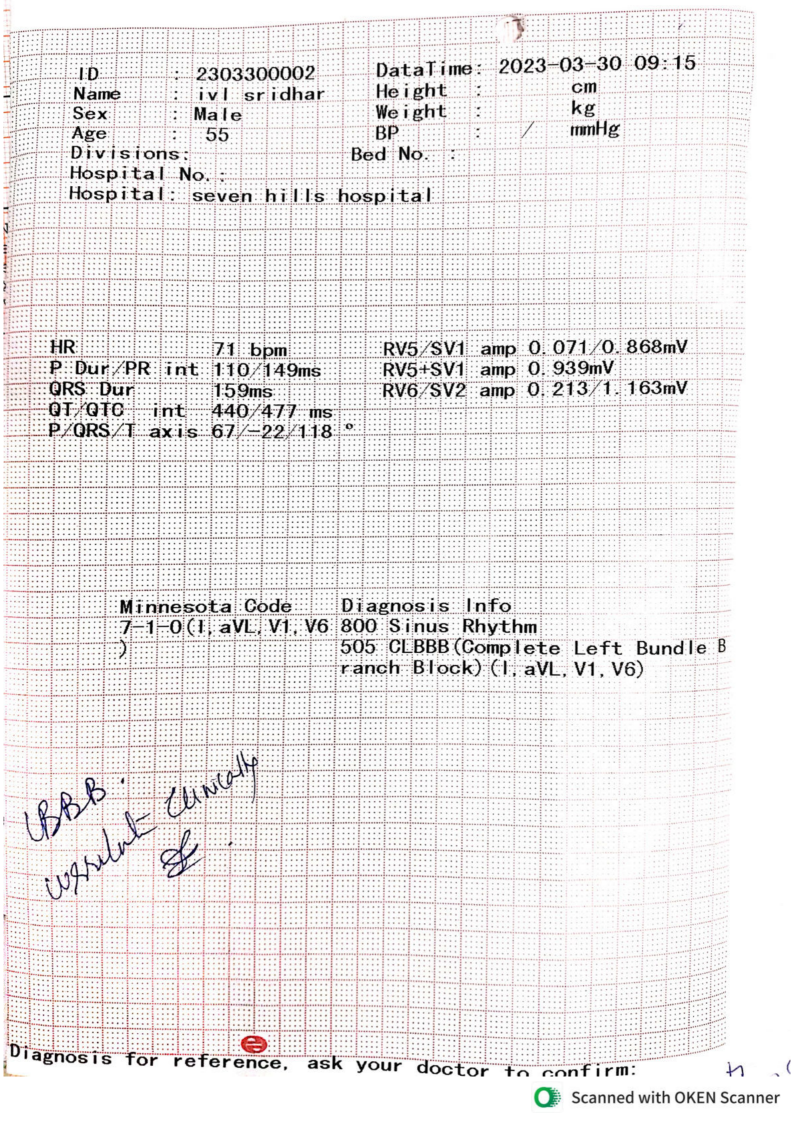
An elevated ESR may occur as an early feature in myocardial infarction. Although a normal ESR cannot be taken to exclude the presence of organic disease, the vast majority of acute or chronic infections and most neoplastic and degenerative diseases are associated with changes in the plasma proteins that increased ESR values.

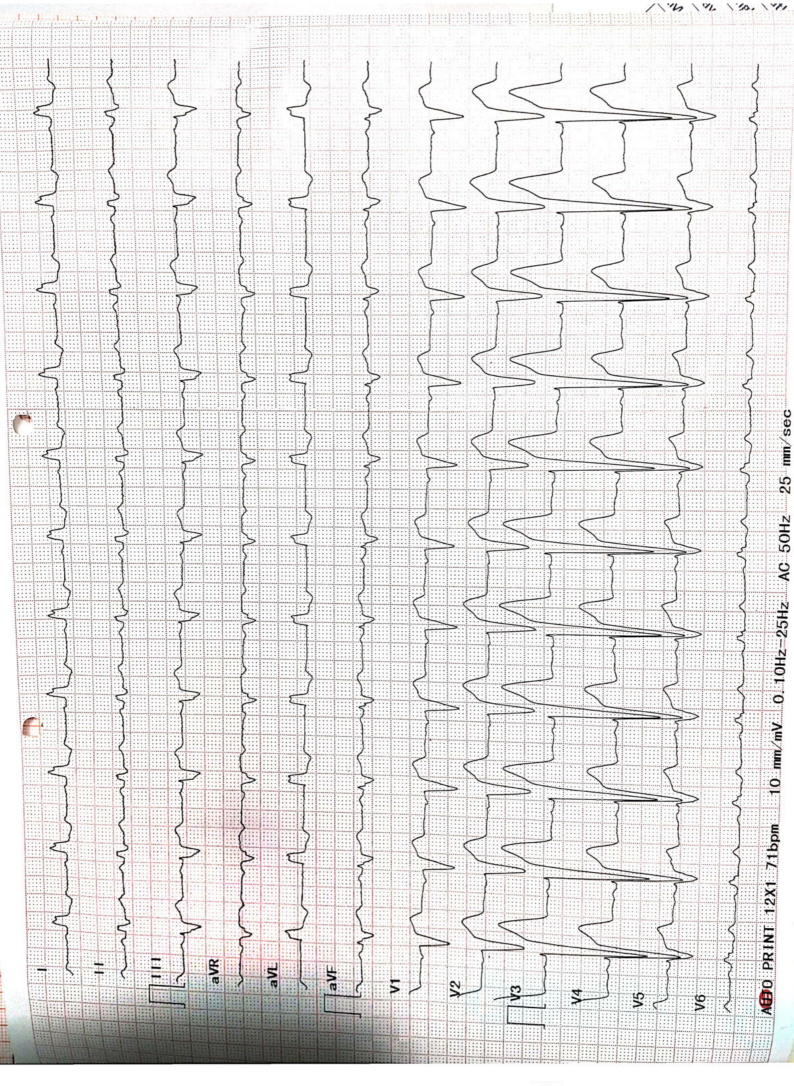
The ESR is influenced by age, stage of the menstrual cycle and medications taken (corticosteroids, contraceptive pills). It is especially low (0–1 mm) in polycythaemia, hypofibrinogenaemia and congestive cardiac failure and when there are abnormalities of the red cells such as poikilocytosis, spherocytosis, or sickle cells. In cases of performance enhancing drug intake by athletes the ESR values are generally lower than the usual value for the individual and as a result of the increase in haemoglobin (i.e. the effect of secondary polycythaemia).

End of Report

Dr.Ritesh Kharche MD, PGD HOD, Laboratory Medicine Dept. RegNo: 2006/03/1680

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				2		MUMBAI		+				
						MUMBAI						
	IVL SHRIDHA	R				TREADMILL	TEST REPORT					
	TD : 4	7125					: Bruce					
	DATE : 3	0-03-202	23			PROTOCOL HISTORY	: Bruce					
	AGE/SEX : 1 HT/WT : 1	165 / 73	3			INDICATION						
	REF.BY : Se	elf				MEDICATION	: NIL					
	PHASE	TOTAL	STAGE	SPEED	GRADE	H.R.		RPP	ST	LEVEL (MM)		MET
		TIME	TIME	Km/Hr	8	bpm	mmHg	x100	II	V1	V5	-
									<u> </u>		1 1 3	
UPINE						81	118 / 80	95	0.3	0.4	0.1	
tage 1		2:55	2:55	2.7	10	113	118 / 80	133	-0.8	1.3	-0.4	4.6
tage 2			2:55	4	12	127	125 / 85	158	-1	1	-1.9	7.0
K-EXERCI: ECOVERY	SE	7:39 10:49	2.55	5.4	14	142 102	125 / 85 125 / 85	177 127	-1.2	1.9	-1.9	8.7
LCOTENI		10.95	2.55			102	125 / 85	127	-0.0	1.2	-0.5	
	RESULTS											
	EXERCISE DURA	ATION	: 7:3				MAX WORK L	OAD	: 8.70	METS		
				hom 86	% of tar	got boart r				and the second s		
	MAX HEART RAT		: 142	Dpm 00		get neart 1	ate 165 bpm					
	MAX HEART RAT MAX BLOOD PRE	SSURE	: 125	/ 85 mm	Hg	get neart I	ate 165 bpm					
	MAX HEART RAT MAX BLOOD PRE REASON OF TEF	SSURE	: 125	/ 85 mm	Hg	get heart h	ate 165 bpm					
	MAX HEART RAT MAX BLOOD PRI REASON OF TEF BP RESPONSE	SSURE	: 125	/ 85 mm	Hg		ate 165 bpm					
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Technician : VIKESH JADHAV





Patient Name: Mr. IVL SRIDHARUHID: SHHM.61755Episode: OPRef. Doctor: Self

Age/Sex	: 55 Year(s) / Male
Order Date	: 30/03/2023 08:42
Mobile No	: 9035281468
DOB	: 12/06/1967
Facility	: SEVENHILLS HOSPITAL, MUMBAI

IMMUNOLOGY

Test Name			Result			Unit R	ef. Range
Sample No :	O0264832C	Collection Date :	30/03/23 08:44	Ack Date :	30/03/2023 10:09	Report Date	: 30/03/23 12:21
	AL-SERUM ate Specific Antige	n - SERUM	1.29			ng/ml	0.00 - 4.00
-	erence Interval :- for all ages: <=4) - 4.5						

Note : Change in method and Reference range

INTERPRETATION :

Prostate-specific antigen (PSA) is a glycoprotein that is produced by the prostate gland, the lining of the urethra, and the bulbourethral gland. PSA exists in serum mainly in two forms, complexed to alpha-1-anti-chymotrypsin (PSA-ACT complex) and unbound (free PSA). Increases in prostatic glandular size and tissue damage caused by benign prostatic hypertrophy, prostatitis, or prostate cancer may increase circulating PSA levels. Transient increase in PSA can also be seen following per rectal digital or sonological examinations.

NOTE:

Patients on Biotin supplement may have interference in some immunoassays. With individuals taking high dose Biotin (more than 5 mg per day) supplements, at least 8-hour wait time before blood draw is recommended. Ref: Arch Pathol Lab Med—Vol 141, November 2017

Nell Alen Tatlor Lab Field Vol 111, November 2017			
T3 - SERUM	104.7	ng/dl	47.00 - 200.00
Method - CLIA			
T4 - SERUM	9.05	ug/dL	4.60 - 10.50
Method - CLIA			
TSH - SERUM	3.76	uIU/ml	0.40 - 5.50
Method - CLIA			

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Reference Ranges (T3) Pregnancy: First Trimester 81 - 190 Second Trimester & Third Trimester 100 - 260

Reference Ranges (TSH) Pregnancy: 1st Trimester : 0.1 – 2.5 2nd Trimester : 0.2 – 3.0 3rd Trimester : 0.3 – 3.0

Reference:

1. Clinical Chemistry and Molecular Diagnostics, Tietz Fundamentals, 7th Edition & Endocronology Guideliens

Interpretation :-

It is recommended that the following potential sources of variation should be considered while interpreting thyroid hormone results:

1. Thyroid hormones undergo rhythmic variation within the body this is called circadian variation in TSH secretion: Peak levels are seen between 2-4 am. Minimum levels seen between 6-10 am. This variation may be as much as 50% thus, influence of sampling time needs to be considered for clinical interpretation.

2. Circulating forms of T3 and T4 are mostly reversibly bound with Thyroxine binding globulins (TBG), and to a lesser extent with albumin and Thyroid binding PreAlbumin. Thus the conditions in which TBG and protein levels alter such as chronic liver disorders, pregnancy, excess of estrogens, androgens, anabolic steroids and glucocorticoids may cause misleading total T3, total T4 and TSH interpretations.

3. Total T3 and T4 levels are seen to have physiological rise during pregnancy and in patients on steroid treatment.

4. T4 may be normal the presence of hyperthyroidism under the following conditions : T3 thyrotoxicosis, Hypoproteinemia related reduced binding, during intake of certain drugs (eg Phenytoin, Salicylates etc)

5. Neonates and infants have higher levels of T4 due to increased concentration of TBG

6. TSH levels may be normal in central hypothyroidism, recent rapid correction of hypothyroidism or hyperthyroidism, pregnancy, phenytoin therapy etc.

7. TSH values of <0.03 uIU/mL must be clinically correlated to evaluate the presence of a rare TSH variant in certain individuals which is undetectable by conventional methods.

8. Presence of Autoimmune disorders may lead to spurious results of thyroid hormones

9. Various drugs can lead to interference in test results.

10. It is recommended that evaluation of unbound fractions, that is free T3 (fT3) and free T4 (fT4) for clinic-pathologic correlation, as these are the metabolically active forms.

End of Report

Dr.Ritesh Kharche MD, PGD HOD, Laboratory Medicine Dept. RegNo: 2006/03/1680

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Patient Name : Mr. IVL SRIDHAR UHID : SHHM.61755 : OP Episode Ref. Doctor : Self

Age/Sex : 55 Year(s) / Male **Order Date** : 30/03/2023 08:42 Mobile No :9035281468 DOB : 12/06/1967 : SEVENHILLS HOSPITAL, MUMBAI Facility

Stool Examination

Fest Name			Result							
Sample No :	O0264832D	Collection Date :	30/03/23 08:44	Ack Date :	30/03/2023 1	0:29	Report Date :	30/03/23 12:44		
Gross and	Chemical Exa	mination								
Consistency	/		Sem	i-Solid						
COLOUR ST	TOOL		Brov	vn						
Visible Bloo	Blood Absent									
Mucus			Abse	ent						
Occult Bloo	d		NEG	ATIVE						
Microscop	ic Examinatio	<u>n</u>								
Puscells			occa	isional						
RBC			Abse	Absent						
Epithelial Ce	ells		Abse	Absent						
Parasites			Not	Not Seen						
Bacteria			Abse	ent						
				End of Rep	ort					



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Facility	: SEVENHILLS HOSPITAL, MUMBAI

		U	rinalysis				
est Name		Resul	t		Unit	Ref	. Range
Sample No : 00264834E Col	llection Date :	30/03/23 08:51	Ack Date :	30/03/2023 10:29		Report Date :	30/03/23 12:39
Physical Examination							
QUANTITY		40			ml		
Colour		Pale	e Yellow				
Appearance		Clea	ar				
DEPOSIT		Abs	ent				Absent
рН		Acio	dic				
Specific Gravity		1.0	15				
Chemical Examination							
Protein		Abs	ent				Absent
Sugar		Abs	ent				Absent
ketones		Abs	ent				Absent
Occult Blood		NEG	GATIVE				Absent
Bile Salt		Abs	ent				Absent
Bile Pigments		Abs	ent				Absent
Urobilinogen		Nor	mal				Absent
NITRATE		Abs	ent				
LEUKOCYTES		Abs	ent				
Microscopic Examination							
Puscells		1-2			/HP	۶F	
Epithelial Cells		occ	asional		, /Hb		
RBC		Abs	ent		, /Hb	۶F	Absent
Cast		Abs	ent		, /LP		Absent
Crystal		Abs	ent		, /HP		Absent
Amorphous Materials		Abs	ent				Absent
Yeast		Abs	ent				Absent
Bacteria		Absent					Absent
URINE SUGAR AND KETONE	(FASTING)						
Sugar	<u></u>	Abs	ent				
ketones		Abs	ent				
	llection Date :	30/03/23 11:14	Ack Date :	30/03/2023 11:25		Report Date :	30/03/23 12:50

URINE SUGAR AND KETONE (PP)

Sugar

Trace

Patient Name: Mr. IVL SRIDHARUHID: SHHM.61755Episode: OP

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ketones

Absent

End of Report

Dr.Ritesh Kharche MD, PGD HOD, Laboratory Medicine Dept. RegNo: 2006/03/1680

URINE SUGAR AND KETONE (PP)- Report has been amended at Mar 30 2023 12:46PM by Ritesh kharche.

DIAGNOSTICS REPORT

Patient Name	: Mr. IVL SRIDHAR	Order Date	: 30/03/2023 08:42
Age/Sex	: 55 Year(s)/Male	Report Date	: 30/03/2023 13:20
UHID Ref. Doctor	: SHHM.61755 : Self	IP No Facility	: : : SEVENHILLS HOSPITAL, MUMBAI

USG ABDOMEN

Liver is normal in size (13.6 cm) and shows bright echotexture. No focal liver parenchymal lesion is seen. Intrahepatic portal and biliary radicles are normal.

Gall-bladder is physiologically distended. No evidence of intraluminal calculus is seen. Wall thickness appears normal. No evidence of peri-cholecystic fluid is seen.

Portal vein and CBD are normal in course and calibre.

Visualised part of pancreas appears normal in size and echotexture. No evidence of duct dilatation or parenchymal calcification seen.

Spleen is normal in size (10.2 cm) and echotexture. No focal lesion is seen in the spleen.

Right kidney measures 9.3 x 4.0 cm. Left kidney measures 10.8 x 4.8 cm.

Both the kidneys are normal in size, shape and echotexture. Cortico-medullary differentiation is maintained. No evidence of calculus or hydronephrosis on either side.

There is no free fluid in abdomen and pelvis. **IMPRESSION:**

•Grade I fatty liver.

Dr.Bhavesh Rajesh Dubey, MBBS, MD

RegNo: 2017/03/0656

DIAGNOSTICS REPORT

Patient Name	: Mr. IVL SRIDHAR	Order Date	: 30/03/2023 08:42	
Age/Sex	: 55 Year(s)/Male	Report Date	: 30/03/2023 16:10	
UHID	: SHHM.61755	IP No	:	
Ref. Doctor	: Self	Facility	: SEVENHILLS HOSPITAL, MUMBAI	

X-RAY CHEST PA VIEW

Both lungs are clear.

The frontal cardiac dimensions are normal.

The pleural spaces are clear.

Both hilar shadows are normal in position and density.

No diaphragmatic abnormality is seen.

The soft tissues and bony thorax are normal.

IMPRESSION: No pleuroparenchymal lesion is seen.

Dr.Bhavesh Rajesh Dubey, MBBS, MD

RegNo: 2017/03/0656