

Tests you can trust

Name : Bg Manjunath Swamy (57Y/M)

Date : 13 Sep 2024

Test Asked: Aarogyam Purush Profile With Utsh, Fbs

Report Status: Complete Report



9 out of 10 Doctors trust that Thyrocare reports are accurate & reliable*

















Accredited by





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CAP From 2007

Thyrocare,

5CA-711, 3rd Floor, HRBR 2nd Block, Hennur, Bengaluru-560043





TEST ASKED

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NAME : BG MANJUNATH SWAMY (57Y/M)

REF. BY : SELF

: AAROGYAM PURUSH PROFILE WITH UTSH,FBS

SAMPLE COLLECTED AT:

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

Report Availability Summary

Note: Please refer to the table below for status of your tests.

21 Ready



Ready with Cancellation



O Processing



(X) O Cancelled in Lab

21 Reduy	• Ready with Cancellation	• • • • • • • • • • • • • • • • • • •	Cancelled III Lab
TEST DETAILS			REPORT STATUS
FASTING BLOOD	SUGAR(GLUCOSE)		Ready ⊘
AAROGYAM PURI	USH PROFILE WITH UTSH		Ready ⊘
CARCINO EMBRY	ONIC ANTIGEN (CEA)		Ready ⊘
CHLORIDE			Ready ⊘
FOLATE			Ready ⊘
MAGNESIUM			Ready ⊘
PHOSPHOROUS			Ready ⊘
PROSTATE SPECI	IFIC ANTIGEN (PSA)		Ready ⊘
SODIUM			Ready ⊘
TESTOSTERONE			Ready ⊘
CARDIAC RISK M	MARKERS		Ready ⊘
COMPLETE URIN	E ANALYSIS		Ready ⊘
HBA PROFILE			Ready ⊘
HEMOGRAM - 6 I	PART (DIFF)		Ready ⊘
ANTI CCP (ACCP)		Ready ⊘
LIVER FUNCTION	N TESTS		Ready ⊘
IRON DEFICIENC	CY PROFILE		Ready ⊘
KIDPRO			Ready ⊘
LIPID PROFILE			Ready ⊘
T3-T4-USTSH			Ready ⊘

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Report Availability Summary

Note: Please refer to the table below for status of your tests.

(> 21 Ready

(v) **0** Ready with Cancellation

(Processing

(x) 0 Cancelled in Lab

TEST DETAILS REPORT STATUS

VITAMIN D TOTAL AND B12 COMBO

Ready 🕢

ANTI NUCLEAR ANTIBODIES (ANA)

Ready 🕢

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: AAROGYAM PURUSH PROFILE WITH UTSH,FBS

T C PALYA MAIN ROAD, BENGALURU

Summary Report

Tests outside reference ran	ge	
OBSERVED VALUE	UNITS	Bio. Ref. Interval.
7.53	Ratio	9:1-23:1
55.9	mg/dL	< 30
4.45	Ratio	< 3.12
8.4	%	1-6
0.74	X 10³ / μL	0.02 - 0.5
3.4	X 10³ / μL	1.0-3.0
26.4	pq	27.0-32.0
80.6	fL	83.0-101.0
37.1	fL	39-46
5.87	X 10^6/μL	4.5-5.5
Present 1+(30-100 mg/dl)	mg/dL	Absent
102.8	mg/dL	70-100
5.8	%	< 5.7
39	mg/dL	40-60
174	mg/dL	< 150
1.29	mg/dL	0.72-1.18
61	mL/min/1.73 m2	>= 90
2.2	ng/mL	> 5.38
	7.53 55.9 4.45 8.4 0.74 3.4 26.4 80.6 37.1 5.87 Present 1+(30-100 mg/dl) 102.8 5.8 39 174 1.29 61	7.53 Ratio 55.9 mg/dL 4.45 Ratio 8.4 % 0.74 X 10³ / μL 3.4 X 10³ / μL 26.4 pq 80.6 fL 37.1 fL 5.87 X 10^6/μL Present 1+(30-100 mg/dL mg/dl) 102.8 mg/dL 5.8 % 39 mg/dL 174 mg/dL 1.29 mg/dL 1.29 mg/dL 61 mL/min/1.73 m2

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REF. BY : SELF

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(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS
ANTI CCP (ACCP)	E.L.I.S.A	1.36	AU/mL
Bio. Ref. Interval. : Negative : < 15			,
Borderline: 15 - 25			

Clinical Significance:

Positive: > 25

Anti-Cyclic-Citrullinated-Peptide (Anti-CCP) Antibodies hold promise for early and more accurate detection of Rheumatoid Arthritis before the disease proceeds into an irreversible damage.

Specifications: Specificity: 94 %, Sensitivity: 76 %

Kit Validation reference: Vossenaar ER et al., Arthritis Rheum., 50, 3485, 2004

Method: INDIRECT SOLID PHASE ENZYME IMMUNOASSAY

ANTI NUCLEAR ANTIBODIES (ANA)

E.L.I.S.A

14.37

AU/mL

Bio. Ref. Interval. :

Negative : < 50 Borderline: 50 - 70 Positive : > 70

Clinical Significance:

Autoimmune diseases are characterized by abnormal functioning of Immune System where cell recognition mechanism fails to distinguish "Self" and "non-self" antigens. Presence of ANA autoantibodies associated with rhematic autoimmune diseases such as systemic Lupus Erythematosus (SLE), Sjogren Syndrome, Scleroderma and mixed connective tissue disease (MCTD).

Specifications:

Specification:- Precision: Intra assay (%CV): <=6.6, Inter assay (%CV): <=13.3, Sensitivity: 87.1%, Specificity: 80%.

Kit Validation Reference:

Antinuclear Antibody The Lancet, September 15, 1984: 611-13

Method: INDIRECT SOLID PHASE IMMUNOASSAY

Please correlate with clinical conditions.

Sample Collected on (SCT)

Sample Received on (SRT) Report Released on (RRT)

Sample Type

Labcode

Barcode

:13 Sep 2024 07:30

: 13 Sep 2024 11:30

:13 Sep 2024 18:13

:SERUM

:1309040578/A4391

:CW104197

Dr Syeda Sumaiya MD(Path)

Dr. Ashwin Mathew MD(Path)

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KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS
CARCINO EMBRYONIC ANTIGEN (CEA)	C.L.I.A	1.61	ng/mL

Bio. Ref. Interval. :-

Non-Smokers: < 2.50 ng/mL Smokers : < 5.00 ng/mL

Clinical Significance:

CEA is often used to monitor patients with cancers of the gastrointestinal tract (GI). Increased CEA levels can indicate some Non-Cancer related conditions, Such as some forms of inflammation, Cirrhosis, and Peptic Ulcer. Also, Smokers tend to have Higher CEA levels than Non-Smokers. When cancer spreads to other organs, CEA levels rise and may be present in other types of bodily fluids besides blood.

For Diagnostic Purpose, Results should always be assessed in Conjunction with the patients medical history, clinical examination and other findings.

Specifications:

Precision: Intra Assay (%CV): 3.6 %, Inter Assay (%CV): 4.1 %; Sensitivity: 0.5 ng/ml

Kit Validation References:

Statland Be, Winkel P. Neoplasia. In: Kaplan LA, Resc AJ, Editors. Clinical Chemistry, Theory, Analysis and Correlation. 2nd Ed. St. Louis: Cv Mosby, 1989.p 734-5.

Please correlate with clinical conditions.

Method:-FULLY AUTOMATED TWO STEP SANDWICH IMMUNOASSAY

Sample Collected on (SCT) : 13 Sep 2024 07:30 : 13 Sep 2024 11:30 Sample Received on (SRT) Report Released on (RRT) : 13 Sep 2024 18:13

. SERUM Sample Type

: 1309040578/A4391 Labcode

Dr Syeda Sumaiya MD(Path)

Dr. Ashwin Mathew MD(Path)

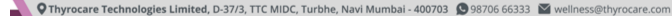
: CW104197 **Barcode** Page: 2 of 21

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TEST NAME	TECHNOLOGY	VALUE	UNITS
25-OH VITAMIN D (TOTAL)	E.C.L.I.A	39.3	ng/mL

Bio. Ref. Interval. :

Deficiency: <=20 ng/ml || Insufficiency: 21-29 ng/ml Sufficiency: >= 30 ng/ml || Toxicity: >100 ng/ml

Clinical Significance:

Vitamin D is a fat soluble vitamin that has been known to help the body absorb and retain calcium and phosphorous; both are critical for building

Decrease in vitamin D total levels indicate inadequate exposure of sunlight, dietary deficiency, nephrotic syndrome.

Increase in vitamin D total levels indicate Vitamin D intoxication.

Specifications: Precision: Intra assay (%CV):9.20%, Inter assay (%CV):8.50%

Kit Validation Reference: Holick M. Vtamin D the underappreciated D-Lightful hormone that is important for Skeletal

and cellular health Curr Opin Endocrinol Diabetes 2002:9(1)87-98.

Method: Fully Automated Electrochemiluminescence Compititive Immunoassay

VITAMIN B-12 E.C.L.I.A 470 pg/mL

Bio. Ref. Interval. : Normal: 197-771 pg/ml

Clinical significance:

Vitamin B12 or cyanocobalamin, is a complex corrinoid compound found exclusively from animal dietary sources, such as meat, eggs and milk. It is critical in normal DNA synthesis, which in turn affects erythrocyte maturation and in the formation of myelin sheath. Vitamin-B12 is used to find out neurological abnormalities and impaired DNA synthesis associated with macrocytic anemias. For diagnostic purpose, results should always be assessed in conjunction with the patients medical history, clinical examination and other findings.

Specifications: Intra assay (%CV):2.6%, Inter assay (%CV):2.3 %

Kit Validation Reference: Thomas L.Clinical laborator Diagnostics: Use and Assessment of Clinical laboratory Results 1st Edition, TH Books-Verl-Ges,1998:424-431

Method: Fully Automated Electrochemiluminescence Compititive Immunoassay

Please correlate with clinical conditions.

Sample Collected on (SCT) :13 Sep 2024 07:30

Sample Received on (SRT) : 13 Sep 2024 11:30 Report Released on (RRT) :13 Sep 2024 18:13

Sample Type :SERUM

:1309040578/A4391 Labcode

Barcode :CW104197

Dr Syeda Sumaiya MD(Path)

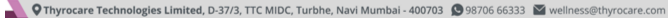
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TEST NAME	TECHNOLOGY	VALUE	UNITS
ADOLIDODDOTEIN A1 (ADO A1)	IMMUNOTUDDIDIMETOV	127	20 C / dl
APOLIPOPROTEIN - A1 (APO-A1)	IMMUNOTURBIDIMETRY	127	mg/dL
Bio. Ref. Interval. : Male : 86 - 152			
Female : 94 - 162			
Method: FULLY AUTOMATED RATE IMMUNOTURBIDIMETR	Y - BECKMAN COULTER		
APOLIPOPROTEIN - B (APO-B)	IMMUNOTURBIDIMETRY	81	mg/dL
Bio. Ref. Interval. :			
Male : 56 - 145			
Female : 53 - 138			
Method: FULLY AUTOMATED RATE IMMUNOTURBIDIMETR	Y - BECKMAN COULTER		
APO B / APO A1 RATIO (APO B/A1)	CALCULATED	0.6	Ratio
Bio. Ref. Interval. : Male : 0.40 - 1.26 Female : 0.38 - 1.14			
Method: Derived from serum Apo A1 and Apo B values			

Please correlate with clinical conditions.

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Sample Received on (SRT) : 13 Sep 2024 11:30 Report Released on (RRT) :13 Sep 2024 18:13

Sample Type :SERUM

Barcode

Labcode :1309040578/A4391

:CW104197

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KARNATAKA,560016

VALUE TEST NAME UNITS TECHNOLOGY HIGH SENSITIVITY C-REACTIVE PROTEIN (HS-CRP) **IMMUNOTURBIDIMETRY** 0.7 mg/L Bio. Ref. Interval. :-

< 1.00 - Low Risk 1.00 - 3.00 - Average Risk >3.00 - 10.00 - High Risk

> 10.00 - Possibly due to Non-Cardiac Inflammation

Disclaimer: Persistent unexplained elevation of HSCRP >10 should be evaluated for non-cardiovascular etiologies such as infection, active arthritis or concurrent illness.

Clinical significance:

High sensitivity C- reactive Protein (HSCRP) can be used as an independent risk marker for the identification of Individuals at risk for future cardiovascular Disease. A coronary artery disease risk assessment should be based on the average of two hs-CRP tests, ideally taken two weeks apart.

Kit Validation Reference:

- 1. Clinical management of laboratory date in medical practice 2003-3004, 207(2003).
- 2.Tietz: Textbook of Clinical Chemistry and Molecular diagnostics: Second edition: Chapter 47:Page no.1507-1508.

Please correlate with clinical conditions.

Method:-FULLY AUTOMATED LATEX AGGLUTINATION - BECKMAN COULTER

Sample Collected on (SCT) : 13 Sep 2024 07:30 : 13 Sep 2024 11:30 Sample Received on (SRT)

Report Released on (RRT) : 13 Sep 2024 18:13

. SERUM Sample Type

Dr Syeda Sumaiya MD(Path) Dr. Ashwin Mathew MD(Path) : 1309040578/A4391 Labcode : CW104197 **Barcode** Page: 5 of 21

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TEST NAME	TECHNOLOGY	VALUE	UNITS
IRON	PHOTOMETRY	92.3	μg/dL
Bio. Ref. Interval. : Male : 65 - 175 Female : 50 - 170			
Method: Ferrozine method without deproteinization			
TOTAL IRON BINDING CAPACITY (TIBC) Bio. Ref. Interval.: Male: 225 - 535 μg/dl Female: 215 - 535 μg/dl	PHOTOMETRY	340.22	μg/dL
Method: Spectrophotometric Assay			
% TRANSFERRIN SATURATION	CALCULATED	27.13	%
Bio. Ref. Interval. : 13 - 45			
Method: Derived from IRON and TIBC values			
UNSAT.IRON-BINDING CAPACITY(UIBC)	PHOTOMETRY	247.92	μg/dL
Bio. Ref. Interval.: 162 - 368			
Method: SPECTROPHOTOMETRIC ASSAY			
FOLATE	C.L.I.A	2.2	ng/mL
Rio Ref Interval :			

Bio. Ref. Interval. : > 5.38 ng/ml

Clinical Significance: Low folate intake, malabsorption as a result of gastrointestinal diseases, pregnancy, and drugs such as phenytoin are causes of folate deficiency.3 Folate deficiency is also associated with chronic alcoholism. Serum folate measurement provides an early index of folate

Specifications: Precision: Intra assay (%CV): 7.93, Inter assay (%CV): 7.19, Sensitivity: 0.35 ng/mL.

Kit Validation References: Steinkamp RC. Vitamin B12 and folic acid: clinical and pathophysiological considerations. In: Brewster MA, Naito HK, eds. Nutritional Elements and Clinical Biochemistry. New York: Plenum Publishing Corp.; 1980:169-240

Method: COMPETITIVE CHEMI LUMINESCENT IMMUNO ASSAY

Please correlate with clinical conditions.

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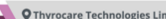
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TEST NAME VALUE UNITS TECHNOLOGY LIPOPROTEIN (A) [LP(A)] **IMMUNOTURBIDIMETRY** 55.9 mg/dL Bio. Ref. Interval. :-

Adults: < 30.0 mg/dl

Clinical Significance:

Determination of LPA may be useful to guide management of individuals with a family history of CHD or with existing disease. The levels of LPA in the blood depends on genetic factors; The range of variation in a population is relatively large and hence for diagnostic purpose, results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings.

Specifications:

Precision %CV :- Intra assay %CV- 4.55% , Inter assay %CV-0.86 %

Kit Validation Reference:

Tietz NW, Clinical Guide to Laboratory Tests Philadelphia WB. Saunders 1995: 442-444

Please correlate with clinical conditions.

Method:-LATEX ENHANCED IMMUNOTURBIDIMETRY

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. SERUM Sample Type

Dr Syeda Sumaiya MD(Path) Dr. Ashwin Mathew MD(Path) : 1309040578/A4391 Labcode : CW104197 **Barcode**

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TEST NAME	TECHNOLOGY	VALUE	UNITS
TESTOSTERONE	E.C.L.I.A	569	ng/dL
Bio. Ref. Interval. :-			

280 - 800

Clinical Significance: Clinical evaluation of serum testosterone, along with serum LH, assists in evaluation of Hypogonadal males. Major causes of lowered testosterone in males include Hypogonadotropic hypogonadism, testicular failure Hyperprolactinema, Hypopituitarism some types of liver and kidney diseases and critical illness.

Specifications: Precision: Intra assay (%CV): 11.50 %, Inter assay (%CV): 5.70%; Sensitivity: 7 ng/dL. Kit Validation Reference: Wilson JD Foster DW (Eds) Williams Textbook of Endocrinology 8th Edition WB Saunders Piladelphia Pennsylvania.

Note: The Biological Reference Range mentioned is specific to the age group and gender. Kindly correlate clinically.

Please correlate with clinical conditions.

Method:-Fully Automated Electrochemiluminescence Compititive Immunoassay

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VALUE TEST NAME UNITS TECHNOLOGY PROSTATE SPECIFIC ANTIGEN (PSA) C.L.I.A 2.83 ng/mL Bio. Ref. Interval. :-

Normal: < 4.00 ng/ml

Border line: 4.01 to 10.00 ng/ml

Clinical Significance:

Elevated levels of PSA are associated with prostate cancer, but may also be seen with prostatitis (Inflammation of the prostate) and benign prostatic hyperplasia (BPH). PSA test done along with free PSA provides additional information. Studies have suggested that the percentage of free PSA in total PSA is lower in patients with prostate cancer than those with benign prostate hyperplasia.

Specification:

Precision: Intra assay (%CV): 4.38%, Inter assay (%CV): 4.67%; Sensitivity: 0.01 ng/ml

Kit validation references:

Wang MC, Valenzuala LA, Murphy GP, and Chu TM. Purification of a human prostate-specific antigen. Invest. Urol. 1979; 17: 159

Please correlate with clinical conditions.

Method:-TWO SITE SANDWICH IMMUNOASSAY

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. SERUM Sample Type

Dr Syeda Sumaiya MD(Path) Dr. Ashwin Mathew MD(Path) : 1309040578/A4391 Labcode : CW104197 **Barcode** Page: 9 of 21

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TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
TOTAL CHOLESTEROL	PHOTOMETRY	130	mg/dL	< 200
HDL CHOLESTEROL - DIRECT	PHOTOMETRY	39	mg/dL	40-60
LDL CHOLESTEROL - DIRECT	PHOTOMETRY	74	mg/dL	< 100
TRIGLYCERIDES	PHOTOMETRY	174	mg/dL	< 150
TC/ HDL CHOLESTEROL RATIO	CALCULATED	3.3	Ratio	3 - 5
TRIG / HDL RATIO	CALCULATED	4.45	Ratio	< 3.12
LDL / HDL RATIO	CALCULATED	1.9	Ratio	1.5-3.5
HDL / LDL RATIO	CALCULATED	0.53	Ratio	> 0.40
NON-HDL CHOLESTEROL	CALCULATED	90.72	mg/dL	< 160
VLDL CHOLESTEROL	CALCULATED	34.8	mg/dL	5 - 40

Please correlate with clinical conditions.

Method:

REF. BY

CHOL - Cholesterol Oxidase, Esterase, Peroxidase

HCHO - Direct Enzymatic Colorimetric

LDL - Direct Measure

TRIG - Enzymatic, End Point

TC/H - Derived from serum Cholesterol and Hdl values

TRI/H - Derived from TRIG and HDL Values

LDL/ - Derived from serum HDL and LDL Values

HD/LD - Derived from HDL and LDL values.

NHDL - Derived from serum Cholesterol and HDL values

VLDL - Derived from serum Triglyceride values

*REFERENCE RANGES AS PER NCEP ATP III GUIDELINES:

TOTAL CHOLESTEROL	(mg/dl)	HDL	(mg/dl)	LDL	(mg/dl)	TRIGLYCERIDES	(mg/dl)
DESIRABLE	<200	LOW	<40	OPTIMAL	<100	NORMAL	<150
BORDERLINE HIGH	200-239	HIGH	>60	NEAR OPTIMAL	100-129	BORDERLINE HIGH	150-199
HIGH	>240			BORDERLINE HIGH	130-159	HIGH	200-499
				HIGH	160-189	VERY HIGH	>500
				VERY HIGH	>190		

Alert !!! 10-12 hours fasting is mandatory for lipid parameters. If not, values might fluctuate.

Sample Collected on (SCT) : 13 Sep 2024 07:30

: 13 Sep 2024 11:30 Sample Received on (SRT) Report Released on (RRT) : 13 Sep 2024 18:13

Sample Type : SERUM

Labcode : 1309040578/A4391

Barcode : CW104197 Dr Syeda Sumaiya MD(Path)

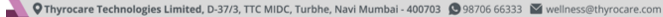
Dr. Ashwin Mathew MD(Path)

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PROCESSED AT: Thyrocare, 5CA-711, 3rd Floor, HRBR 2nd Block, Hennur, Bengaluru-560043

REF. BY





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: BG MANJUNATH SWAMY (57Y/M) NAME

: SELF

TEST ASKED : AAROGYAM PURUSH PROFILE WITH UTSH

SAMPLE COLLECTED AT:

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC, #63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
ALKALINE PHOSPHATASE	PHOTOMETRY	54.44	U/L	45-129
BILIRUBIN - TOTAL	PHOTOMETRY	0.91	mg/dL	0.3-1.2
BILIRUBIN -DIRECT	PHOTOMETRY	0.19	mg/dL	< 0.3
BILIRUBIN (INDIRECT)	CALCULATED	0.72	mg/dL	0-0.9
GAMMA GLUTAMYL TRANSFERASE (GGT)	PHOTOMETRY	23.9	U/L	< 55
ASPARTATE AMINOTRANSFERASE (SGOT)	PHOTOMETRY	22	U/L	< 35
ALANINE TRANSAMINASE (SGPT)	PHOTOMETRY	38	U/L	< 45
SGOT / SGPT RATIO	CALCULATED	0.58	Ratio	< 2
PROTEIN - TOTAL	PHOTOMETRY	7.16	gm/dL	5.7-8.2
ALBUMIN - SERUM	PHOTOMETRY	4.36	gm/dL	3.2-4.8
SERUM GLOBULIN	CALCULATED	2.8	gm/dL	2.5-3.4
SERUM ALB/GLOBULIN RATIO	CALCULATED	1.56	Ratio	0.9 - 2

Please correlate with clinical conditions.

Method:

ALKP - Modified IFCC method

BILT - Vanadate Oxidation

BILD - Vanadate Oxidation

BILI - Derived from serum Total and Direct Bilirubin values

GGT - Modified IFCC method

SGOT - IFCC* Without Pyridoxal Phosphate Activation

SGPT - IFCC* Without Pyridoxal Phosphate Activation

OT/PT - Derived from SGOT and SGPT values.

PROT - Biuret Method

SALB - Albumin Bcg1method (Colorimetric Assay Endpoint)

SEGB - DERIVED FROM SERUM ALBUMIN AND PROTEIN VALUES

A/GR - Derived from serum Albumin and Protein values

Sample Collected on (SCT)

: 13 Sep 2024 07:30

Sample Received on (SRT)

: 13 Sep 2024 11:30

Report Released on (RRT)

: 13 Sep 2024 18:13

Sample Type

: SERUM

Labcode

: 1309040578/A4391

Dr Syeda Sumaiya MD(Path)

Dr. Ashwin Mathew MD(Path)

Barcode : CW104197

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: BG MANJUNATH SWAMY (57Y/M) NAME

REF. BY

TEST ASKED : AAROGYAM PURUSH PROFILE WITH UTSH **SAMPLE COLLECTED AT:**

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS
MAGNESIUM	PHOTOMETRY	1.95	mg/dL
Bio. Ref. Interval. :-			5,

1.90 - 3.10 mg/dL

Clinical significance:

Magnesium is the fourth most abundant cation in the body and second most prevalent intracellular cation. The total body magnesium content is about 25 g or approximately 1 mol, of which 55% reside in the skeleton. About 45% of the magnesium is intracellular. In general higher the metabolic activity of cell, the greater is its magnesium content. Magnesium is a cofactor for more than 300 enzymes in the body.

Disorders of magnesium metabolism are separated into those causing hypomagnesaemia/magnesium deficiencies and hypermagnesemia. Hypomagnesaemia is common in patient in hospitals. Moderate to severe deficiency of magnesium is usually due to loss of magnesium from the gastrointestinal (gi) tract or kidneys. One of the more serious complications of magnesium deficiency is cardiac arrhythmia. Symptomatic hypermagnsemia is almost always caused by excessive intake, resulting from administration of antacids, enemas, and parenteral fluids containing magnesium. Depression of neuromuscular system is the most common manifestation of magnesium intoxication.

External quality control program participation:

College Of American Pathologists: Chemistry survey; CAP Number: 7193855-01

Please correlate with clinical conditions.

Method:-MODIFIED XYLIDYL BLUE REACTION METHOD

Sample Collected on (SCT) : 13 Sep 2024 07:30 : 13 Sep 2024 11:30 Sample Received on (SRT)

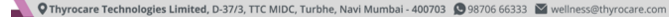
Report Released on (RRT) : 13 Sep 2024 18:13

. SERUM Sample Type

Dr Syeda Sumaiya MD(Path) Dr. Ashwin Mathew MD(Path) • 1309040578/A4391 Labcode : CW104197 **Barcode** Page: 12 of 21

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: BG MANJUNATH SWAMY (57Y/M) NAME

: SELF

TEST ASKED : AAROGYAM PURUSH PROFILE WITH UTSH

SAMPLE COLLECTED AT:

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
BLOOD UREA NITROGEN (BUN)	PHOTOMETRY	9.71	mg/dL	7.94 - 20.07
CREATININE - SERUM	PHOTOMETRY	1.29	mg/dL	0.72-1.18
BUN / SR.CREATININE RATIO	CALCULATED	7.53	Ratio	9:1-23:1
UREA (CALCULATED)	CALCULATED	20.78	mg/dL	Adult: 17-43
UREA / SR.CREATININE RATIO	CALCULATED	16.11	Ratio	< 52
CALCIUM	PHOTOMETRY	9.3	mg/dL	8.8-10.6
URIC ACID	PHOTOMETRY	6.25	mg/dL	4.2 - 7.3
PHOSPHOROUS	PHOTOMETRY	4.71	mg/dL	2.4 - 5.1

Please correlate with clinical conditions.

Method:

REF. BY

BUN - Kinetic UV Assay.

SCRE - Creatinine Enzymatic Method

B/CR - Derived from serum Bun and Creatinine values

UREAC - Derived from BUN Value.

UR/CR - Derived from UREA and Sr.Creatinine values.

CALC - Arsenazo III Method, End Point.

URIC - Uricase / Peroxidase Method

PHOS - UNREDUCED PHOSPHOMOLYBDATE METHOD

Sample Collected on (SCT) : 13 Sep 2024 07:30

: 13 Sep 2024 11:30 Sample Received on (SRT) Report Released on (RRT) : 13 Sep 2024 18:13

Sample Type : SERUM

: 1309040578/A4391 Labcode

Barcode : CW104197

Dr Syeda Sumaiya MD(Path)

Dr. Ashwin Mathew MD(Path)

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: BG MANJUNATH SWAMY (57Y/M) NAME

: SELF

TEST ASKED : AAROGYAM PURUSH PROFILE WITH UTSH **SAMPLE COLLECTED AT:**

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS
SODIUM	I.S.E	141.63	mmol/L
Bio. Ref. Interval. : ADULTS: 136-145 MMOL/L			
Method: ION SELECTIVE ELECTRODE			
CHLORIDE	I.S.E	104.36	mmol/L

Bio. Ref. Interval. : ADULTS: 98-107 MMOL/L

Clinical Significance:

An increased level of blood chloride (called hyperchloremia) usually indicates dehydration, but can also occur with other problems that cause high blood sodium, such as Cushing syndrome or kidney disease. Hyperchloremia also occurs when too much base is lost from the body (producing metabolic acidosis) or when a person hyperventilates (causing respiratory alkalosis). A decreased level of blood chloride (called hypochloremia) occurs with any disorder that causes low blood sodium. Hypochloremia also occurs with congestive heart failure, prolonged vomiting or gastric suction, Addison disease, emphysema or other chronic lung diseases (causing respiratory acidosis), and with loss of acid from the body (called metabolic alkalosis).

Method: ION SELECTIVE ELECTRODE

Please correlate with clinical conditions.

Sample Collected on (SCT) :13 Sep 2024 07:30

Sample Received on (SRT) : 13 Sep 2024 11:30 Report Released on (RRT) :13 Sep 2024 18:13

Sample Type :SERUM

Labcode :1309040578/A4391

Barcode :CW104197

Dr Syeda Sumaiya MD(Path)

Dr. Ashwin Mathew MD(Path)

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µIU/mL

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4.32

NAME : BG MANJUNATH SWAMY (57Y/M)

SAMPLE COLLECTED AT: (5600162992), SHREE KRISHNA HOMOEOPATHIC

REF. BY : SELF CLINIC, #63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU KARNATAKA,560016

TEST ASKED : AAROGYAM PURUSH PROFILE WITH UTSH

> UNITS Bio. Ref. Interval. **VALUE** 80-200 129 ng/dL 9.78 μg/dL 4.8-12.7

> > 0.54-5.30

TSH - ULTRASENSITIVE Comments:

TOTAL THYROXINE (T4)

TOTAL TRIIODOTHYRONINE (T3)

The Biological Reference Ranges is specific to the age group. Kindly correlate clinically.

Method:

TEST NAME

T3,T4 - Fully Automated Electrochemiluminescence Compititive Immunoassay USTSH - Fully Automated Electrochemiluminescence Sandwich Immunoassay

Disclaimer: Results should always be interpreted using the reference range provided by the laboratory that performed the test. Different laboratories do tests using different technologies, methods and using different reagents which may cause difference. In reference ranges and hence it is recommended to interpret result with assay specific reference ranges provided in the reports. To diagnose and monitor therapy doses, it is recommended to get tested every time at the same Laboratory.

TECHNOLOGY

E.C.L.I.A

E.C.L.I.A

E.C.L.I.A

Sample Collected on (SCT) : 13 Sep 2024 07:30 Sample Received on (SRT) : 13 Sep 2024 11:30 Report Released on (RRT) : 13 Sep 2024 18:13

: SERUM Labcode

: 1309040578/A4391

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Barcode : CW104197

Sample Type

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: BG MANJUNATH SWAMY (57Y/M) NAME

REF. BY

TEST ASKED : AAROGYAM PURUSH PROFILE WITH UTSH **SAMPLE COLLECTED AT:**

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

VALUE TEST NAME UNITS TECHNOLOGY EST. GLOMERULAR FILTRATION RATE (eGFR) CALCULATED 61 mL/min/1.73 m2 Bio. Ref. Interval. :-

> = 90 : Normal 60 - 89 : Mild Decrease

45 - 59 : Mild to Moderate Decrease 30 - 44 : Moderate to Severe Decrease

15 - 29 : Severe Decrease

Clinical Significance

The normal serum creatinine reference interval does not necessarily reflect a normal GFR for a patient. Because mild and moderate kidney injury is poorly inferred from serum creatinine alone. Thus, it is recommended for clinical laboratories to routinely estimate glomerular filtration rate (eGFR), a "gold standard" measurement for assessment of renal function, and report the value when serum creatinine is measured for patients 18 and older, when appropriate and feasible. It cannot be measured easily in clinical practice, instead, GFR is estimated from equations using serum creatinine, age, race and sex. This provides easy to interpret information for the doctor and patient on the degree of renal impairment since it approximately equates to the percentage of kidney function remaining. Application of CKD-EPI equation together with the other diagnostic tools in renal medicine will further improve the detection and management of patients with CKD.

Reference

Levey AS, Stevens LA, Schmid CH, Zhang YL, Castro AF, 3rd, Feldman HI, et al. A new equation to estimate glomerular filtration rate. Ann Intern Med. 2009;150(9):604-12.

Please correlate with clinical conditions. Method:-CKD-EPI Creatinine Equation

Sample Collected on (SCT) : 13 Sep 2024 07:30 : 13 Sep 2024 11:30 Sample Received on (SRT) Report Released on (RRT) : 13 Sep 2024 18:13

. SERUM Sample Type

: 1309040578/A4391 Labcode

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: CW104197 **Barcode** Page: 16 of 21

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NAME : BG MANJUNATH SWAMY (57Y/M)

REF. BY : SELF

TEST ASKED : COMPLETE URINE ANALYSIS **SAMPLE COLLECTED AT:**

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

TEST NAME	METHODOLOGY	VALUE	UNITS	Bio. Ref. Interval
Complete Urinogram				
Physical Examination				
VOLUME	Visual Determination	3	mL	-
COLOUR	Visual Determination	PALE YELLOW	-	Pale Yellow
APPEARANCE	Visual Determination	CLEAR	-	Clear
SPECIFIC GRAVITY	pKa change	1.01	-	1.003-1.030
PH	pH indicator	5	-	5-8
Chemical Examination				
URINARY PROTEIN	PEI	Present 1+(30-100 mg/dl)	mg/dL	Absent
URINARY GLUCOSE	GOD-POD	ABSENT	mg/dL	Absent
URINE KETONE	Nitroprusside	ABSENT	mg/dL	Absent
URINARY BILIRUBIN	Diazo coupling	ABSENT	mg/dL	Absent
UROBILINOGEN	Diazo coupling	Normal	mg/dL	<=0.2
BILE SALT	Hays sulphur	ABSENT	-	Absent
BILE PIGMENT	Ehrlich reaction	ABSENT	-	Absent
URINE BLOOD	Peroxidase reaction	ABSENT	-	Absent
NITRITE	Diazo coupling	ABSENT	-	Absent
LEUCOCYTE ESTERASE	Esterase reaction	ABSENT	-	Absent
Microscopic Examination				
MUCUS	Microscopy	ABSENT	-	Absent
RED BLOOD CELLS	Microscopy	ABSENT	cells/HPF	0-5
URINARY LEUCOCYTES (PUS CELLS)	Microscopy	ABSENT	cells/HPF	0-5
EPITHELIAL CELLS	Microscopy	1	cells/HPF	0-5
CASTS	Microscopy	ABSENT	-	Absent
CRYSTALS	Microscopy	ABSENT	-	Absent
BACTERIA	Microscopy	ABSENT	-	Absent
YEAST	Microscopy	ABSENT	-	Absent
PARASITE	Microscopy	ABSENT	-	Absent

(Reference: *PEI - Protein error of indicator, *GOD-POD - Glucose oxidase-peroxidase)

Sample Collected on (SCT) : 13 Sep 2024 07:30 Sample Received on (SRT) : 13 Sep 2024 11:01

Report Released on (RRT) : 13 Sep 2024 12:35

Sample Type : URINE

Labcode : 1309066426/A4391

Barcode : CW104196 Dr Syeda Sumaiya MD(Path)

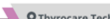
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NAME : BG MANJUNATH SWAMY (57Y/M)

REF. BY : SELF

: BLOOD SUGAR (F) **TEST ASKED**

SAMPLE COLLECTED AT:

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

TEST NAME	TECHNOLOGY	VALUE	UNITS	
FASTING BLOOD SUGAR(GLUCOSE)	PHOTOMETRY	102.8	ma/dL	

Bio. Ref. Interval. :-

As per ADA Guideline: Fasting Plasma Glucose (FPG)			
Normal 70 to 100 mg/dl			
Prediabetes	100 mg/dl to 125 mg/dl		
Diabetes	126 mg/dl or higher		

The assay could be affected mildly and may result in anomalous values if serum samples have heterophilic antibodies, hemolyzed, icteric or lipemic. The concentration of Glucose in a given specimen may vary due to differences in assay methods, calibration and reagent specificity. For diagnostic purposes results should always be assessed in conjunction with patients medical history, clinical findings and other findings.

Please correlate with clinical conditions.

Method:-**GOD-PAP METHOD**

Sample Collected on (SCT) : 13 Sep 2024 07:30

: 13 Sep 2024 11:12 Sample Received on (SRT) Report Released on (RRT) : 13 Sep 2024 11:55

. FLUORIDE Sample Type

: 1309067741/A4391 Dr Syeda Sumaiya MD(Path) Labcode

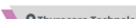
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Barcode : CW104198 Page: 18 of 21

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: BG MANJUNATH SWAMY (57Y/M) NAME

REF. BY : SELF

TEST ASKED : HBA PROFILE, HEMOGRAM **SAMPLE COLLECTED AT:**

(5600162992), SHREE KRISHNA HOMOEOPATHIC CLINIC,#63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD, BENGALURU

KARNATAKA,560016

VALUE TEST NAME TECHNOLOGY UNITS HbA1c - (HPLC) H.P.L.C 5.8 %

Bio. Ref. Interval.:

Bio. Ref. Interval.: As per ADA Guidelines

Below 5.7% : Normal 5.7% - 6.4% : Prediabetic >=6.5% : Diabetic

Guidance For Known Diabetics

Below 6.5%: Good Control 6.5% - 7% : Fair Control

7.0% - 8% : Unsatisfactory Control

>8% : Poor Control

Method: Fully Automated H.P.L.C method

AVERAGE BLOOD GLUCOSE (ABG)

CALCULATED

120

mg/dL

Bio. Ref. Interval.:

90 - 120 mg/dl : Good Control 121 - 150 mg/dl : Fair Control

151 - 180 mg/dl: Unsatisfactory Control

> 180 mg/dl : Poor Control

Method: Derived from HBA1c values

Please correlate with clinical conditions.

Sample Collected on (SCT) :13 Sep 2024 07:30

Sample Received on (SRT) : 13 Sep 2024 11:12 Report Released on (RRT) : 13 Sep 2024 12:22

Sample Type : EDTA Whole Blood Labcode :1309067683/A4391

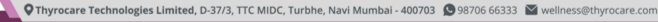
Barcode :CW104199

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: BG MANJUNATH SWAMY (57Y/M) NAME

REF. BY SELF

: HBA PROFILE, HEMOGRAM **TEST ASKED**

SAMPLE COLLECTED AT:

(5600162992), SHREE KRISHNA

HOMOEOPATHIC CLINIC, #63, AKSHAYANAGAR 1BLOCK, 12TH CROSS, T C PALYA MAIN ROAD,

BENGALURU KARNATAKA,560016

TEST NAME	METHODOLOGY	VALUE	UNITS	Bio. Ref. Interval
TOTAL LEUCOCYTES COUNT (WBC)	HF & FC	8.86	X 10 ³ / μL	4.0 - 10.0
NEUTROPHILS	Flow Cytometry	49.8	%	40-80
LYMPHOCYTE	Flow Cytometry	38.4	%	20-40
MONOCYTES	Flow Cytometry	2.9	%	2-10
EOSINOPHILS	Flow Cytometry	8.4	%	1-6
BASOPHILS	Flow Cytometry	0.2	%	0-2
IMMATURE GRANULOCYTE PERCENTAGE(IG%)	Flow Cytometry	0.3	%	0-0.5
NEUTROPHILS - ABSOLUTE COUNT	Calculated	4.41	$X~10^3$ / μL	2.0-7.0
LYMPHOCYTES - ABSOLUTE COUNT	Calculated	3.4	X 10³ / μL	1.0-3.0
MONOCYTES - ABSOLUTE COUNT	Calculated	0.26	$X~10^3$ / μL	0.2 - 1.0
BASOPHILS - ABSOLUTE COUNT	Calculated	0.02	X 10 ³ / μL	0.02 - 0.1
EOSINOPHILS - ABSOLUTE COUNT	Calculated	0.74	X 10³ / μL	0.02 - 0.5
IMMATURE GRANULOCYTES(IG)	Calculated	0.03	$X~10^3$ / μL	0-0.3
TOTAL RBC	HF & EI	5.87	X 10^6/μL	4.5-5.5
NUCLEATED RED BLOOD CELLS	Calculated	0.01	X 10 ³ / μL	0.0-0.5
NUCLEATED RED BLOOD CELLS %	Flow Cytometry	0.01	%	0.0-5.0
HEMOGLOBIN	SLS-Hemoglobin Method	15.5	g/dL	13.0-17.0
HEMATOCRIT(PCV)	CPH Detection	47.3	%	40.0-50.0
MEAN CORPUSCULAR VOLUME(MCV)	Calculated	80.6	fL	83.0-101.0
MEAN CORPUSCULAR HEMOGLOBIN(MCH)	Calculated	26.4	pq	27.0-32.0
MEAN CORP.HEMO.CONC(MCHC)	Calculated	32.8	g/dL	31.5-34.5
RED CELL DISTRIBUTION WIDTH - SD(RDW-SD) Calculated		37.1	fL	39-46
RED CELL DISTRIBUTION WIDTH (RDW-CV)	Calculated	12.8	%	11.6-14
PLATELET DISTRIBUTION WIDTH(PDW)	Calculated	10.9	fL	9.6-15.2
MEAN PLATELET VOLUME(MPV)	Calculated	10	fL	6.5-12
PLATELET COUNT	HF & EI	218	X 10 ³ / μL	150-410
PLATELET TO LARGE CELL RATIO(PLCR)	Calculated	25.6	%	19.7-42.4
PLATELETCRIT(PCT)	Calculated	0.22	%	0.19-0.39

Remarks: Alert!!! Predominantly normocytic normochromic with ovalocytes. Platelets: Appear adequate in smear.

Clinical history is asked for all the relevant abnormalities detected and in absence / failure of receiving of clinical history, results are rechecked twice and released. Advised clinical correlation.

Method: Fully automated bidirectional analyser (6 Part Differential SYSMEX XN-1000)

(Reference: *FC- flowcytometry, *HF- hydrodynamic focussing, *EI- Electric Impedence, *Hb- hemoglobin, *CPH- Cumulative pulse height)

~~ End of report ~~

Sample Collected on (SCT)

Sample Received on (SRT)

Report Released on (RRT)

Sample Type

Labcode **Barcode**



:13 Sep 2024 07:30

: 13 Sep 2024 11:12

: 13 Sep 2024 12:22

: EDTA Whole Blood

: 1309067683/A4391

: CW104199

Dr Syeda Sumaiya MD(Path)

Dr. Ashwin Mathew MD(Path)

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CONDITIONS OF REPORTING

- v The reported results are for information and interpretation of the referring doctor only.
- v It is presumed that the tests performed on the specimen belong to the patient; named or identified.
- v Results of tests may vary from laboratory to laboratory and also in some parameters from time to time for the same patient.
- v Should the results indicate an unexpected abnormality, the same should be reconfirmed.
- v Only such medical professionals who understand reporting units, reference ranges and limitations of technologies should interpret results.
- v This report is not valid for medico-legal purpose.
- v Neither Thyrocare, nor its employees/representatives assume any liability, responsibility for any loss or damage that may be incurred by any person as a result of presuming the meaning or contents of the report.
- v Thyrocare Discovery video link :- https://youtu.be/nbdYeRqYyOc
- v For clinical support please contact @8450950852,8450950853,8450950854 between 10:00 to 18:00

EXPLANATIONS

- v Majority of the specimen processed in the laboratory are collected by Pathologists and Hospitals we call them as "Clients".
- v Name The name is as declared by the client and recored by the personnel who collected the specimen.
- v Ref.Dr The name of the doctor who has recommended testing as declared by the client.
- v Labcode This is the accession number in our laboratory and it helps us in archiving and retrieving the data.
- v **Barcode** This is the specimen identity number and it states that the results are for the specimen bearing the barcode (irrespective of the name).
- v **SCP** Specimen Collection Point This is the location where the blood or specimen was collected as declared by the client.
- v SCT Specimen Collection Time The time when specimen was collected as declared by the client.
- v SRT Specimen Receiving Time This time when the specimen reached our laboratory.
- v RRT Report Releasing Time The time when our pathologist has released the values for Reporting.
- v Reference Range Means the range of values in which 95% of the normal population would fall.

SUGGESTIONS

- Values out of reference range requires reconfirmation before starting any medical treatment.
- v Retesting is needed if you suspect any quality shortcomings.
- v Testing or retesting should be done in accredited laboratories.
- v For suggestions, complaints or feedback, write to us at info@thyrocare.com or call us on 022-3090 0000 / 6712 3400
- v SMS:<Labcode No.> to **9870666333**

