

Tests you can trust

- Name : <u>Xxxxxxxxx xxxxxxxxx xxxx</u>
- Date : <u>19 Apr 2025</u>
- Test Asked : Master Wellness Checkup

Report Status:

Complete Report



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NABL From 2005

ISO 9001: 2015 - From 2015

CAP From 2007

PROCESSED AT Thyrocare D-37/1,TTC MIDC Navi Mumbai-400 Thyrocare 1 First Na NAME REF. BY TEST ASKED Report Avai Note: Please refer	: C,Turbhe, 0 703 Technologies Limited, D-37/3, TTC MIDC, Turbhe, M tional Diagnostic Chain to have : XXXXXXXXX XXXXX XXXX : SELF : MASTER WELLNESS CHECKUP Tability Summary r to the table below for status of your tests	Navi Mumbai - 400 703	Construction C
✓ 37 Ready	0 Ready with Cancellation	🕑 0 Processing	🚫 0 Cancelled in Lab
TEST DETAILS			REPORT STATUS
MASTER WELLI	NESS CHECKUP		Ready 😔
ERYTHROCYTE	SEDIMENTATION RATE (ESR)		Ready 😔
FASTING BLOO	DD SUGAR(GLUCOSE)		Ready ⊘
CHLORIDE			Ready ⊘
25-OH VITAMI	N D (TOTAL)		Ready 😔
CYSTATIN C			Ready ⊘
FERRITIN			Ready 😔
FRUCTOSAMIN	IE		Ready 😔
HOMOCYSTEIN	ΙE		Ready 😔
HIGH SENSITI	VITY C-REACTIVE PROTEIN (HS-CRP)		Ready 😔
INSULIN - FAS	STING		Ready ⊘
URINOGRAM			Ready ⊘
LIPASE			Ready ⊘
Lipoprotein (a)) [Lp(a)]		Ready 😔
MAGNESIUM			Ready ⊘
LP-PLA2			Ready ⊘
SERUM COPPE	R		Ready ⊘
SERUM ZINC			Ready ⊘
SODIUM			Ready ⊘
TESTOSTERON	IE		Ready ⊘
VITAMIN A			Ready ⊘

PROCESSED AT : 'hyrocare Thyrocare D-37/1,TTC MIDC,Turbhe, Tests you can trust Navi Mumbai-400 703 💡 Thyrocare Technologies Limited, D-37/3, TTC MIDC, Turbhe, Navi Mumbai - 400 703 👂 9870666333 🛯 wellness@thyrocare.com First National Diagnostic Chain to have 100% of its Labs with NABL Accreditation[#] NAME : XXXXXXXXX XXXXXXXXX XXXX **HOME COLLECTION :** REF. BY : SELF TEST ASKED : MASTER WELLNESS CHECKUP ***** ****** **Report Availability Summary** Note: Please refer to the table below for status of your tests. 37 Ready O Ready with Cancellation (Processing × 0 Cancelled in Lab **TEST DETAILS REPORT STATUS** VITAMIN E Ready ⊘ VITAMIN K Ready ⊘ **HBA PROFILE** Ready ⊘ HEMOGRAM - 6 PART (DIFF) Ready ⊘ ANTI CCP (ACCP) Ready 📿

ANTI CCF (ACCF)	Reauy 🕑
LIVER FUNCTION TESTS	Ready ⊘
ELEMENTS 22 (TOXIC AND NUTRIENTS)	Ready ⊘
IRON DEFICIENCY PROFILE	Ready ⊘
KIDPRO	Ready ⊘
LIPID PROFILE	Ready ⊘
VITAMIN B COMPLEX PROFILE	Ready ⊘
T3-T4-USTSH	Ready ⊘
URINE MICROALBUMIN	Ready ⊘
APOLIPROTEIN RATIO	Ready ⊘
AMYLASE	Ready ⊘
ANTI NUCLEAR ANTIBODIES (ANA)	Ready ⊘
BLOOD KETONE (D3HB)	Ready 🔗

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Navi Mumbai-400 703



MC-2407 Care

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NAME	: XXXXXXXXX XXXXXXXXX XXXX
REF. BY	: SELF

REF. BY: SELF**TEST ASKED**: MASTER WELLNESS CHECKUP

HOME COLLECTION :

	Summary Report			
Tests outside reference range				
TEST NAME	OBSERVED VALUE	UNITS	Bio. Ref. Interval.	
COMPLETE HEMOGRAM				
EOSINOPHILS - ABSOLUTE COUNT	0.51	X 10³ / μL	0.02 - 0.5	
HEMOGLOBIN	12.6	g/dL	13.0-17.0	
LYMPHOCYTES - ABSOLUTE COUNT	3.42	X 10³ / μL	1.0-3.0	
MEAN PLATELET VOLUME(MPV)	9	fL	7.5-8.3	
PLATELET COUNT	433	X 10³ / μL	150-410	
PLATELET DISTRIBUTION WIDTH(PDW)	8.8	fL	9.6-15.2	
PLATELET TO LARGE CELL RATIO(PLCR)	17	%	19.7-42.4	
ELECTROLYTES				
CHLORIDE	107.7	mmol/L	98 - 107	
IRON DEFICIENCY				
IRON	63.1	µg/dL	65 - 175	
LIPID				
HDL CHOLESTEROL - DIRECT	70	mg/dL	40-60	
LDL / HDL RATIO	1.3	Ratio	1.5-3.5	
TC/ HDL CHOLESTEROL RATIO	2.5	Ratio	3 - 5	
OTHER COUNTS				
ERYTHROCYTE SEDIMENTATION RATE (ESR)	25	mm / hr	0 - 15	
RENAL				
CREATININE - SERUM	0.51	mg/dL	0.72-1.18	
URIC ACID	3.07	mg/dL	4.2 - 7.3	
TOXIC ELEMENTS				
BERYLLIUM	0.08	µg/L	0.10 - 0.80	
URINE				
CREATININE - URINE	23.28	mg/dL	39 - 259	
VITAMINS				
VITAMIN B-12	1570	pg/mL	197-771	

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NAME: XXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXXX		HOME COLLECTION : xxxxxxxxx xxxxxxxxx xxxx xxxx xxxx XxxxxXX xxxxxxxx				
TEST ASKED	. PROTER WELLINESS CHECKOP	Xxxxxxxx TECHNOLOGY	XXXXXXXXXXX XXXX XXX VALUE	(XXX XXXX XXXXXXXXXXXXXXXXXXXXXXXXXXXX		
VITAMIN A		LC-MS/MS	388.37	ng/mL		
Bio. Ref. Inte	rval. :-					

 Age
 Reference range

 1 - 6 Years
 200 - 400

 7 - 12 Years
 260 - 490

 13 - 19 Years
 260 - 720

 Above 18 Years
 300 - 800

Clinical Significance:

Vitamin A or Retinol plays important role in the function retinal vision, growth, reproduction, embryonic development as well as in immune function. Vitamin A is also required for adaptive immunity and plays a role in the development of both T- helper cells and B-cells. Retinol and its metabolites and synthetic retinoids provide protective effects against the development of certain types of cancer by blocking tumor promotion, by inhibiting proliferation, by inducing apoptosis, by inducing differentiation or by performing combination of these actions.

Fat malabsorption, particularly caused by celiac disease or chronic pancreatitis and protein -energy malnutrition predispose to vitamin A deficiency. Clinical features of vitamin A deficiency include degenerative changes in eyes and skin and poor dark adaptation or night blindness. Vitamin A deficiency impairs innate immunity by impeding normal regeneration of mucosal bariers damaged by infection and by diminishing the function of neutrophils, macrophages and natural killer cells.

Toxic effects of hypervitaminosis A have occurred as a result of ingestion of excess vitamin or as a side effect of inappropriate therapy. Symptoms of acute toxicity from single massive dose present as abdominal pain, nausea,, vomiting, severe headache, dizziness, sluggishness and irritability. Chronic toxicity shows symptoms like bone, join pain, hair loss, dryness and fissures of the lips, anorexia, benign intracranial hypertension, weight loss an hepatomegaly.

Clinical reference:

Tietz Textbok of clinical chemistry and Molecular diagnostic, Carl A. Burtis, Edward R. Ashwood, David E. Bruns, Fifth edition., Elsevier.

 Please correlate with clinical conditions.

 Method: LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY



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8434.76

ng/mL

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NAME REF. BY	: XXXXXXXXX XXXXXXXXXX XXXX : SELF	HOME COLLECTION : XXXXXXXXX XXXXXXXXX XX		(xxxxxx xxxx Xxxxxxxxx	
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx xxxx xxxx xxxxx xxxx xxxxxxxx			
TEST NAME		TECHNOLOGY	VALUE	UNITS	
VITAMIN E		LC-MS/MS	8434 76	na/ml	

VITAMIN E

Bio. Ref. Interval. :-

Age

1000 - 3500
2000 - 6000
3500 - 8000
5500 - 9000
5500 - 18000

Clinical Significance: Vitamin E or Alpha-tocopherol (body's main form of vitamin) function as antioxidant which protects vitamin A, C and red blood cells from oxidative damage caused by free radicals. It has been recognized as necessary for neurologic and reproductive functions, for prevention of retinopathy in premature infants. Alpha-tocopherol also induces inhibition of cell proliferation, platelete aggregation, and monocyte adhesion, which are thought to be the results of direct interaction of alpha-tocopherol with cell components. Alpha-tocopherol reduces inflammatory mediator production.

Premature and low birth weight infants are particularly susceptible to development of vitamin E deficiency, because placental transfer is poor and infants have such limited adipose tissue where much of the vitamins is normally stored. Signs of deficiency include irritability, edema and hemolytic anemia. Although symptoms of vitamin E deficiency are rare in children and adults, deficiency can occur in some conditions.

Excess vitamin E intake usually is achieved only by dietary supplementation. A comprehensive review of tolerance and safety of vitamin E suggested that intakes upto 3000mg/d were safe and reversible side effects of gastrointestinal symptoms, increased creatinuria, and impairment of blood coagulation are seen at intakes of 1000-3000 mg/d. However as noted earlier, long term use of intakes greater than 400mg/d may cause increased mortality.

Clinical reference: Tietz Textbok of clinical chemistry and Molecular diagnostic, Carl A. Burtis, Edward R. Ashwood, David E. Bruns, Fifth edition., Elsevier.

Please correlate with clinical conditions.

Method:-LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY



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NAME : XXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXX		HOME COLLECTION : xxxxxxxxx xxxxxxxx xxxx xxxxx xxxx Xxxxx Xxxxxxx			
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx Xxxxxxxx	xxxx xxxxxx xxxx x xxxxxxxx xxx xxx xx	xxxxxxxxxx xxxx xxxx xxxxxxxxxxx	
TEST NAME		TECHNOLOGY	VALUE	UNITS	
VITAMIN K		LC-MS/MS	0.18	ng/mL	
Bio. Ref. Inte	rval. :-			5,	

- - --

0.13 - 1.19

Clinical significance:

Vitamin K assay measures the principal form of vitamin K i.e. K1 :Phylloquinone which found predominantly in green leafy vegetables, margarines and plant oils.

Vitamin K promotes clotting of the blood, is required for the conversion of several clotting factors and prothombin, and is of growing interest in bone metabolism. Vitamin K plays important role in the deposition of ionic calcium needed for proper blood coagulation and bone formation.

Although vitamin K deficiency in the adults is uncommon, the risk is increased for fat malabsorption states such as bile duct obstruction, cystic fibrosis, chronic pancreatitis and liver disease. Risk also increased by the use of drugs that interfere with vitamin K metabolism, such as warfarin, cepahlosporin. Defective blood coagulation and demonstration of abnormal noncarboxylated prothrombin are at present the only well-established signs of vitamin K deficiency.

The use of high doses of naturally occurring vitamin K (K1 and K2) appears to have no untoward effect; however menadione(K3) treatment can lead to formation of erythrocyte cytoplasmic inclusions known as Heinz bodies and hemolytic anemia. With severe hemolysis, increase bilirubin formation and undeveloped capacity for its conjugation may produce kernicterus in the newborn.

Clinical reference:

Tietz Textbok of clinical chemistry and Molecular diagnostic, Carl A. Burtis, Edward R. Ashwood, David E. Bruns, Fifth edition., Elsevier.

Please correlate with clinical conditions.

Method:- LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY





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NAME	: XXXXXXXXX XXXXXXXXX XXXX
REF. BY	: SELF
TEST ASKED	: MASTER WELLNESS CHECKUP

HOME COLLECTION :				
XXXXXXXXX	XXXXXXXXX			
XXXXXXXXXXXXXX	xxxxxxxxxxxx			
XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	XXXXXXXXXX			

XXXXXXXXX XXXXXXXXXX

xxxx xxxxxx xxxxxxxxxxxxxxxx

PATIENTID : MD22463706

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
VITAMIN B1/THIAMIN	LC-MS/MS	0.67	ng/mL	0.5-4.0
VITAMIN B2/RIBOFLAVIN	LC-MS/MS	11.02	ng/mL	1.6-68.2
VITAMIN B3/NICOTINIC ACID	LC-MS/MS	0.54	ng/mL	0.3-9.8
VITAMIN B5/PANTOTHENIC	LC-MS/MS	128.94	ng/mL	11-150
VITAMIN B6/P5P	LC-MS/MS	37.85	ng/mL	5 - 50
VITAMIN B7/BIOTIN	LC-MS/MS	1.18	ng/mL	0.2-3
VITAMIN B9/FOLIC ACID	LC-MS/MS	0.3	ng/mL	0.2-20

Please correlate with clinical conditions.

Method :

VITB1 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY VITB2 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY VITB3 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY VITB5 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY VITB6 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY VITB7 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY VITB9 - LIQUID CHROMATOGRAPHY TANDEM MASS SPECTROMETRY

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

Sample Type

Labcode

Barcode



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NAME	: XXXXXXXXX XXXXXXXXX XXXX	HOME COLLECTION :			
REF. BY : SELF TEST ASKED : MASTER WELLNESS CHECKUP	xxxxxxxx xxxxxxxx xxxx xxxx xxxx xxxx XxxxXXXXXX				
	xxxxxxxx xxxx xxxx xxxx xxxx xxxx xxxxxx				
TEST NAME		TECHNOLOGY	VALUE	UNITS	
HOMOCYSTEIN	NE	PHOTOMETRY	6.18	µmol/L	

Bio. Ref. Interval. :-

Normal Levels : <15 µmol/L Mild Hyperhomocysteinemia : 15-30 µmol/L Moderate Hyperhomocysteinemia : 30-100 µmol/L Severe Hyperhomocysteinemia : >100 µmol/L

Clinical Significance:

Homocysteine is linked to increased risk of premature coronary artery disease, stroke and thromboembolism. Moreover, alzheimers disease, osteoporosis, venous thrombosis, schizophrenia, cognitive deficiency and pregnancy complications also elevates Homocysteine levels. The results should be interpreted in conjunction with clinical history and other findings.

High Values:

Elevated homocysteine levels might be due to increasing age, genetic traits, drugs, renal dysfunction and dietary deficiency of vitamins or smoking. To lower your homocysteine, eat more green vegetables, stop smoking, alcohol. Folic acid helps lowering elevated levels.

Specifications: Kit Validation Reference: Eikelboom JW, et al Ann Intern Med 131 : 363-75 (1999) https://www.healthline.com/health/homocysteine-levels

Please correlate with clinical conditions.

Method:- SMALL MOLECULE CAPTURE TECHNOLOGY (SMT)



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NAME	: XXXXXXXXX XXXXXXXXX XXXX
REF. BY	: SELF

: MASTER WELLNESS CHECKUP

HOME COLLECTION :

VALUE

< 0.5

PATIENTID : MD22463706

TEST NAME

TEST ASKED

C.M.I.A

U/mL

UNITS

ANTI CCP (ACCP) Reference Range :-

BRI	Interpretation
Negative < 5	Absence of IgG autoantibodies to cyclic citrullinated peptides (CCP)
Positive ≥ 5	Presence of IgG autoantibodies to cyclic citrullinated peptides (CCP)

Clinical Significance :

- 1. Anti-Cyclic-Citrullinated-Peptide (Anti-CCP) titre is used for diagnosis and monitoring of Rheumatoid Arthritis (RA).
- 2. RA is one of the most common systemic autoimmune diseases characterised by chronic inflammation of the synovial joints and progressive joint degeneration eventually leading to disability of affected individuals.
- 3. The diagnosis of RA often relies on clinical manifestations and certain non-specific laboratory tests such as rheumatoid factor (RF) and C-reactive protein (CRP), which may be present in healthy elderly persons or in patients with other autoimmune and infectious diseases.
- 4. Whereas, Anti-Cyclic-Citrullinated-Peptide (Anti-CCP) Antibodies hold promise for early and more accurate detection of Rheumatoid Arthritis before the disease proceeds into irreversible damage.
- 5. Interference with pathologic levels of nonspecific IgG can not be excluded.
- 6. The anti-CCP test results can be false negative in patients with hypergammaglobulinemia.
- Results from patients suffering from this disorder should not be used for diagnostic purposes.
- 7. Heterophile antibodies may interfere with the test results.
- 8. If results are inconsistent with clinical history additional testing is suggested to confirm the results.
- 9. Some specimens may not dilute linearly because of heterogeneity of autoantibodies with respect to physicochemical properties.
- 10. HAMA (Human Anti mouse antibodies) may also interfere with the results.
- 11. For diagnostic purposes, the results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings.

References:

- Anti-CCP Reagent Kit Insert
- Feldmann M, Brennan FM, Maini RN. Rheumatoid arthritis.Cell 1996;85:307-3102.
- Landewé RB. The benefits of early treatment in rheumatoid arthritis:confounding by indication, and the issue of timing. Arthritis Rheum 2003;48(1):1-5.

Please correlate with clinical conditions.

Method:- Fully Automated ChemiLuminescent Microparticle Immunoassay (C.M.I.A)

Sample Collected on (SCT)	:1	
Sample Received on (SRT)	::	
Report Released on (RRT)	:2	
Sample Type	: 9	
Labcode	:>	
Barcode	:>	



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NAME : XXXXXXXX XXXXXXXX XXXX		HOME COLLECTION :			
REF. BY	: SELF	******	«xxxxxxx xxxx xxxxx xxxx Xxxxxxxxx		
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx xxxx xxxxx xxxx xxxx xxxx xxxxx		XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	
TEST NAME		TECHNOLOGY	VALUE	UNITS	
CYSTATIN C		IMMUNOTURBIDIMETRY	0.72	mg/L	
Bio. Ref. Inte	rval. :-			2.	

<= 60 years: <= 1.03 mg/L > 60 years : < 1.50 mg/L

Clinical significance

Cystatin c, is a small 13-kda protein and is a member of the cysteine proteinase inhibitor family, it is produced at a constant rate by all nucleated cells. Due to its small size it is freely filtered by the glomerulus and is not secreted but is fully reabsorbed and broken down by the renal tubules. This means that the primary determinate of blood Cystatin c levels is the rate at which it is filtered at the glomerulus making it an excellent gfr marker. Cystatin c is also a marker of inflammation and like many other markers of inflammation; its serum concentration may be higher in patients with decreased renal clearance. There is mounting evidence, however, that Cystatin c may be a predictor of adverse outcomes independent of renal function with its higher sensitivity to detect a reduced GFR than Creatinine determination, also in the so-called "Creatinine-blind" range. Thus, Cystatin c is suggested to be a better marker for GFR than the ubiquitous serum Creatinine.

Reference

1. Barrett aj, Davies me, Grubb a. the place of human gamma-trace (Cystatin c) among the cysteine proteinase inhibitors. Biochem biophys res common 1984; 120: 631-6.

2. Grubb a. diagnostic value of analysis of Cystatin c and protein HC in biological fluids. Clin Nephrol 1992; 38: S20-7.

Please correlate with clinical conditions. Method:- LATEX ENHANCED IMMUNOTURBIDIMETRY

Sample Collected on (SCT)	:	
Sample Received on (SRT)	:	
Report Released on (RRT)	:	
Sample Type	:	
Labcode	:	
Barcode	:	







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NAME	:	xxxxxxxx xxxxxxx xxx		HOME COLLECTIO	N :
REF. BY : SELF TEST ASKED : MASTER WELLNESS CHECKUP		XXXXXXXXX XXXXXXXX XXXXX XXXXX XXXXX		xxx xxxx xxxxxx	
			XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX		
PATIENTID	:	MD22463706			
TEST NAME			TECHNOLOGY	VALUE	UNITS
APOLIPOPROT	EIN	- A1 (APO-A1)	IMMUNOTURBIDIMETR Y	143	mg/dL
Bio. Ref. IntervaMale:86 -Female:94 -	• 152 • 162	1			
Method : FULLY	Y AU	TOMATED RATE IMMUNOTURBIDIMET	RY – BECKMAN COULTER		
APOLIPOPROT	EIN	І-В (АРО-В)	IMMUNOTURBIDIMETR Y	68	mg/dL
Bio. Ref. Interva Male : 56 -	il. : 145	i			
Female : 53 -	138				
Method : FULLY	Y AU	TOMATED RATE IMMUNOTURBIDIMET	RY – BECKMAN COULTER		
APO B / APO A	\1 R	ATIO (APO B/A1)	CALCULATED	0.5	Ratio
Bio. Ref. Interva	ıl. :				
Male : 0.40 - 1.26					
Female : 0.38 - 1.1	14				
Clinical Significanc	ce :				
Apolipoprotein B	8 is a	more potent and independent predic	tor of Coronary artery disease (CAD) than LDL Chole	sterol.
Apolipoprotein A	1 is	one of the apoproteins of HDL and is	inversely related to risk of CAD.		
The Apolipoprote	ein s	tudies help in monitoring risk of reste	enosis in patients with myocardia	al infarction, Coronary	y bypass surgery etc.
An increased rat	10 01	Apo B to A1 beyond the defined norr	nal range is indicative of CAD ris	šk.	

• All results have to be interpreted in Conjunction with clinical history and other findings.

Method : Derived from serum Apo A1 and Apo B values

Please correlate with clinical conditions.



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NAME : XXXXXXXX XXXXXXXX XXXX		HOME COLLECTION :			
REF. BY	: SELF	xxxxxxxx xxxxxxxx xxxx xxxx xxxx xxxx xxxx		xxx xxxx Xxxxxxxxx	
TEST ASKED	: MASTER WELLNESS CHECKUP			XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	
TEST NAME		TECHNOLOGY	VALUE	UNITS	
HIGH SENSITI Bio. Ref. Inte	VITY C-REACTIVE PROTEIN (HS-CRP)	IMMUNOTURBIDIMETRY	0.45	mg/L	

< 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



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NAME : XXXXXXXX XXXXXXXXXXXXXXXXXXXXXXXXXX
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HOME COLLECTION :

TEST ASKED : MASTER WELLNESS CHECKUP

: SELF

PATIENTID : MD22463706

TEST NAME	TECHNOLOGY	VALUE	UNITS
ANTI NUCLEAR ANTIBODIES (ANA)	E.L.I.S.A	24.8	AU/mL
Bio. Ref. Interval. :			

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

REF. BY

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.



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NAME : X>	****		HOME COLLECTION	1:
REF. BY : SE	LF		*****	
TEST ASKED : M/	ASTER WELLNESS CHECKUP		XXXXXXXXXXXXXXXXXXXXX XXXXXXXXXXXXXXXX	xxxxxxxx xxxx xxxxxxxxxxxxxxxxxxxxx xx
PATIENTID : M	D22463706			
TEST NAME		TECHNOLOGY	VALUE	UNITS
IRON		PHOTOMETRY	63.1	µg/dL
Bio. Ref. Interval. : Male : 65 - 175				
Female : 50 - 170				
Method : Ferrozine met	hod without deproteinization			
TOTAL IRON BINDIN	G CAPACITY (TIBC)	PHOTOMETRY	349.1	µg/dL
Bio. Ref. Interval. : Male: 225 - 535 μg/dl Fe	male: 215 - 535 µg/dl			
Method : Spectrophoto	metric Assay			
% TRANSFERRIN SA	TURATION	CALCULATED	18.08	%
Bio. Ref. Interval. : 13 - 45				
Method : Derived from	IRON and TIBC values			
FERRITIN		C.M.I.A	45.5	ng/mL
Bio. Ref. Interval. : Men: 21.81 - 274.66 ng/n	n			
Women: 4.63 - 204.00 ng	/ml			
Method : Fully Automa	ed Chemi Luminescent Microparti	cle Immunoassay		
UNSAT.IRON-BINDIN	G CAPACITY(UIBC)	PHOTOMETRY	286	µg/dL
Bio. Ref. Interval. : 162 - 368				
Method : SPECTROPHO	TOMETRIC ASSAY			

Please correlate with clinical conditions.



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NAME REF. BY TEST ASKED	: XXXXXXXXX XXXXXXXXX XXXX : SELF : MASTER WELLNESS CHECKUP	HOME COLLECTION : xxxxxxxx xxxxxxxxx xxxx xxxxx xxxx Xxxx xxxxxxxx		
TEST NAME		TECHNOLOGY	VALUE	UNITS
INSULIN - FAS	TING	C.L.I.A	7.29	uU/mL
Bio. Ref. Inte	rval. :-			F/

1.9-23 µU/mL

Clinical Significance

Type I (Insulin dependent: "Juvenile") diabetes is due to a destruction of the beta cells, with a consequence of absolute lack of insulin. In type II (Non insulin-dependent: "Maturity onset") diabetes, insulin resistance may play an important role; However after several years of evolution, beta-cells failure may occur, leading to a relative insulinopenia requiring, in some cases, insulin administration. Insulin resistance is associated with high circulation levels of the hormone.

For diagnostic purpose, results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings.

Specifications:

Precision: Intra Assay (%CV): 4.20 %, Inter Assay (%CV): 5.60%; Sensitivity: 0.03 µU/mL

External quality control program participation:

College Of American Pathologists: Insulin Survey (Ing): Cap Number: 7193855-01

Kit validation references:

Howanitz PJ, Howanitz JH, Henry JB. Carbohydrates.Clinical Diagnosis and Management by Laboratory Methods 1991 ;172-182.edited by Henry JB, Philadelphia, W.B Saunders Company.

Please correlate with clinical conditions.

Method:- One step Immunoenzymatic (Sandwich) assay.

Sample Collected on (SCT)	:	 2	
Sample Received on (SRT)	:		
Report Released on (RRT)	:		
Sample Type			
Labcode	: 3		
Barcode	: >		

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NAME REF. BY TEST ASKED	: XXXXXXXXX XXXXXXXXX XXXX : SELF : MASTER WELLNESS CHECKUP	HOME COLLECTION :			
TEST NAME		TECHNOLOGY	VALUE	UNITS	
BLOOD KETONE (D3HB)		PHOTOMETRY	0.37	ma/dL	
Bio. Ref. Inte	rval. :-		0.07		

0.21-2.81 mg/dL

Clinical Significance:

Three types of ketones can be produced in body D-3- Hydroxybutyrate, Acetoacetate and Acetone. D-3- Hydroxybutyrate accounts for approximately 75% of the ketone bodies. During periods of ketosis, D-3- Hydroxybutyrate increases more than the other two. It has been shown to be a better index of ketoacidosis. In diabetics, D-3- Hydroxybutyrate is needed for the assessment of the severity of diabetic coma and to calculate insulin requirements.

Speficcation: Precision: Intra assay (%CV): 4.53, Inter assay (%CV): 2.9, Sensitivity: 10.41 mg/dL.

Kit validation references: Mcmurray, C.H., Blanchflower, W.J., Rice, D.A., ClinChem., 1984;30:No.3.

Please correlate with clinical conditions.

Method:- ENZYMATIC (KINETIC)



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NAME	: XXXXXXXXX XXXXXXXXX XXXX	HOME COLLECTION :			
REF. BY	: SELF	xxxxxxxx xxxxxxx xxxx xxxx xxxx xxxx xxxx			XXX
TEST ASKED	: MASTER WELLNESS CHECKUP				xxxx
TEST NAME		TECHNOLOGY	VALUE	UNITS	
FRUCTOSAMINE		PHOTOMETRY	236.4	µmol/L	
Bio. Ref. Inte	rval. :-			. ,	

Normal < 286 µmol/L

Clinical Significance:

Fructosamine assay is useful in monitoring the degree of glycemia over short-to-intermediate time frames (1-3 weeks) concentration greater than the established normal range is an indication of prolonged hyperglycemia of 1-3 weeks or longer. The higher fructosamine value, poorer is the degree of glycemia control.

Specifications:

Precision %CV : Intra assay %CV- 3.2% , Inter assay %CV-4.0%, Sensitivity:- 290 umol/L

Kit Validation Reference:

Howey JEA, Browning MCK, Fraser CG. Assay of serum fructosamine that minimizes standardization and matrix problems: Use to assess components of biological va-riation. Clin Chem 1987; 33: 269-272.

Please correlate with clinical conditions. Method:-NITROBLUE TETRAZOLIUM ASSAY (NBT)



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NAME REF. BY TEST ASKED	: XXXXXXXXX XXXXXXXXXX XXXX : SELF : MASTER WELLNESS CHECKUP	HOME COLLECTION : xxxxxxxx xxxxxxxxx xxxx xxxx Xxxx Xxxx			
TEST NAME		TECHNOLOGY	VALUE	UNITS	
Lipoprotein (a) [Lp(a)] Bio Ref Interval :-		IMMUNOTURBIDIMETRY	7.3	mg/dL	

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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REF. BY	: SELF	*******	xxxxxxxx xxxxxxxx xxxx xxxx xxxx Xxxx XxxxXXX				
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx xxxx xxxxx xxxxx xxxx xxxx xxxx					
TEST NAME		TECHNOLOGY	VALUE	UNITS			
LP-PLA2		PHOTOMETRY	112	nmol/min/mL			
Bio. Ref. Inte	rval. :-						

Low Risk : < 225 nmol/min/mL High Risk : >= 225 nmol/min/mL

Clinical Significance:

Lp-PLA2, is an enzyme produced by inflammatory cells. It is predominantly associated with low-density lipoprotein (LDL) and high-density lipoprotein (HDL). Lp-PLA2 is a specific marker of vascular inflammation and found to be upregulated in atherosclerotic lesions especially in complex plaque prone to rupture. A meta-analysis found that Lp-PLA2 levels are positively correlated with increased risk of developing coronary heart disease and stroke. Lp-PLA2 is not an acute phase reactant and thus is unaffected by systemic inflammatory processes. Lp-PLA2 activity should be interpreted in conjunction with clinical evaluation and other risk factor assessment.

Specification:

Precision %CV: Intra assay %CV- 1.50%, Inter assay %CV-3.80%

Kit Validation Reference:

Alexander Thompson etal., The Lp-PLA2 Studies Collaboration (2010). "Lipoprotein-associated phospholipase A2 and risk of coronary disease, stroke, and mortality: collaborative analysis of 32 prospective studies". The Lancet 375 (9725): 1536-1544

 Please correlate with clinical conditions.

 Method: ENZYMATIC ASSAY



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TEST NAME		TECHNOLOGY		
SERUM COPPER Bio Ref Interval :-		PHOTOMETRY	126.26	µg/dL

Male : 63.5 - 150 Female : 80 - 155

Clinical significance:

Copper is an important trace element and a component of numerous enzymes and proteins involved in energy production, connective tissue formation, melanin synthesis, iron metabolism, development of central nervous system, angiogenesis as well as an antioxidant. Deficiency can cause- Malnourishment, cardiovascular disease, anemia & neuropathy, toxicity may be manifested as acute renal failure, gastroenteritis & chronic liver disease.

Specifications:

Precision: Intra assay (%CV): 1.17, Inter assay (%CV): 2.32.

Kit validation references: Thomas L. Clinical Laboratory Diagnostics. 1st ed. Frankfurt: TH-Books Verlagsgesellschaft; 1998. p. 337-8

Please correlate with clinical conditions. Method:- 3,5-DIBR-PAESA



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		HOME COLLECTION :				
REF.BY : SELF						
TEST ASKED	: MASTER WELLNESS CHECKUP	XXXXXXXXX XXXX XXXXX XXXX XXXX XXXXX XXXX				
TEST NAME		TECHNOLOGY	VALUE	UNITS		
SERUM ZINC		PHOTOMETRY	118.01	µg/dL		
Bio. Ref. Inte	rval. :-					

52 - 286

Clinical Significance:

Zinc is one of the essential trace elements in the body. Its metalloenzymes play a key role in protein and nucleic acid synthesis, gene expression, wound healing, as an antioxidant, etc. Deficiency can cause- Poor wound healing, gastroenteritis, impaired spermatogenesis, Alzheimer's disease, etc. Toxicity may be manifested as pancreatitis, gastric ulcer, anemia, pulmonary fibrosis.

Specifications: Precision: Intra assay (%CV): 2.02, Inter assay (%CV): 2.22.

Kit Validation References: Thomas L. Clinical Laboratory Diagnostics. 1st ed. Frankfurt: TH-Books Verlagsgesellschaft; 1998. p. 347-9

Please correlate with clinical conditions. Method:- NITRO - PAPS



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NAME	: XXXXXXXXX XXXXXXXXX XXXX	HOME COLLECTION :				
REF. BY	: SELF	XXXXXXXXX	xxxxxxxxx xxxx xxx	(XXXXX XXXX XXXXX XXXX XXXXXXXXXXXXXXX		
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx xxxx xxxx xxxx xxxx xxxx xxxxxx				
TEST NAME		TECHNOLOGY	VALUE	UNITS		
TESTOSTERONE		E.C.L.I.A	92.1	ng/dL		
Bio. Ref. Inte	rval. :-			5,		

4 - 836.7

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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REF. BY	: SELF	xxxxxxx x	xxxx xxxx Xxxxxxxxx			
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx xxxx xxxx xxxxx xxxx xxxx xxxxx				
TEST NAME		TECHNOLOGY	VALUE	UNITS		
AMYLASE		PHOTOMETRY	83.1	U/L		
Bio. Ref. Inte	rval. :-			•		

Adults : 28-100 U/L

Interpretation:

Lipemic Sera (Hypertriglyceridemia) may contain inhibitors, Which falsely depress results. About 20% of patients with Acute Pancreatitis have abnormal lipids. Normal serum amylase may occur in Pancreatitis, Especially relapsing and chronic pancreatitis. Moderate increases may be reported in normal pregnancy.

Clinical Significance:

Causes of high Serum Amylase include Acute Pancreatitis, Pancreatic Pseudocyst, Pancreatic Ascites, Pancreatic Abscess, Neoplasm in or adjacent to Pancreas, Trauma to Pancreas, and common Duct Stones. Nonpancreatic Causes include inflammatory salivary lesions (Eg, Mumps), Perforated Peptic Ulcer, Intestinal Obstruction, Biliary Tract Disease, Peritonitis, Acute Appendicitis, Diabetic Ketoacidosis, and Extrapancreatic Carcinomas. Amylase levels more than 25-fold the upper limit of normal are often found when metastatic tumors produce Ectopic Amylase.

Specifications:

Precision: Intra assay (%CV): 2.82, Inter assay (%CV): 2.49, Sensitivity: 10.9 U/L.

Kit Validation References:

Rauscher, E., et coll., Fresenius Z. Analyt. Chem. 324 (1986) 304-305.

Please correlate with clinical conditions.

Method:- ENZYMATIC COLORIMETRIC TEST



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REF. BY	: SELF	XXXXXXXXXX >	xxxxxxxxx xxxx xxxxx xxxx Xxxxxxxx			
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx xxxx xxxx xxxx xxxx xxxx xxxx xxxx				
TEST NAME		TECHNOLOGY	VALUE	UNITS		
LIPASE		PHOTOMETRY	25.3	U/L		
Bio. Ref. Inte	rval. :-					

Adults : 5.6 - 51.3 U/L

Interpretation:

For diagnostic purposes, the results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings like serum amylase. Serum Lipase is usually normal in patients with elevated serum amylase, having peptic ulcer, salivary adenitis, inflammatory bowel disease, intestinal obstruction, and macroamylasemia. Lipemic sera may interfere with results.

Clinical Significance:

High serum Lipase is a specific marker for pancreatitis; after acute pancreatitis the Lipase activity increases within 4-8 hours, reaches a peak after 24 hours and decreases after 8 to 14 days. However, there is no correlation between the Lipase activity determined in serum and the extent of damage to the pancreas.

Specifications:

Precision: Intra assay (%CV): 3.35, Inter assay (%CV): 2.46, Sensitivity: 3.5 U/L.

Kit Validation References:

Tietz Nw Et Al. Lipase In Serum - The Elusive Enzyme: An Overview. Clin Chem 1993; 39:746-756.

Please correlate with clinical conditions.

Method:- ENZYMATIC COLORIMETRIC ASSAY









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NAME	: XXXXXXXXX XXXXXXXXXX XXXX
REF. BY	: SELF
TEST ASKED	: MASTER WELLNESS CHECKUP

HOME COLLECTION :

xxxx xxxxxx

PATIENTID : MD22463706

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
TOTAL CHOLESTEROL	PHOTOMETRY	177	mg/dL	< 200
HDL CHOLESTEROL - DIRECT	PHOTOMETRY	70	mg/dL	40-60
HDL / LDL RATIO	CALCULATED	0.78	Ratio	> 0.40
LDL CHOLESTEROL - DIRECT	PHOTOMETRY	89	mg/dL	< 100
TRIG / HDL RATIO	CALCULATED	0.82	Ratio	< 3.12
TRIGLYCERIDES	PHOTOMETRY	57	mg/dL	< 150
TC/ HDL CHOLESTEROL RATIO	CALCULATED	2.5	Ratio	3 - 5
LDL / HDL RATIO	CALCULATED	1.3	Ratio	1.5-3.5
NON-HDL CHOLESTEROL	CALCULATED	106.9	mg/dL	< 160
VLDL CHOLESTEROL	CALCULATED	11.42	mg/dL	5 - 40

Please correlate with clinical conditions.

Method :

CHOL - Cholesterol Oxidase, Esterase, Peroxidase HCHO - Direct Enzymatic Colorimetric HD/LD - Derived from HDL and LDL values. LDL - Direct Measure TRI/H - Derived from TRIG and HDL Values TRIG - Enzymatic, End Point TC/H - Derived from serum Cholesterol and Hdl values LDL/ - Derived from serum 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.









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REF. BY	: SELF
TEST ASKED	: MASTER WELLNESS CHECKUP

HOME COLLECTION : XXXXXXXXX

Xxxxxxxx xxxxxxxx

XXXXXXXXX XXXX ***** *****

PATIENTID : MD22463706

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
ALKALINE PHOSPHATASE	PHOTOMETRY	159.6	U/L	127-403
BILIRUBIN - TOTAL	PHOTOMETRY	0.65	mg/dL	0.3-1.2
BILIRUBIN -DIRECT	PHOTOMETRY	0.12	mg/dL	< 0.3
BILIRUBIN (INDIRECT)	CALCULATED	0.53	mg/dL	0-0.9
GAMMA GLUTAMYL TRANSFERASE (GGT)	PHOTOMETRY	12.5	U/L	< 55
SGOT / SGPT RATIO	CALCULATED	1.15	Ratio	< 2
ASPARTATE AMINOTRANSFERASE (SGOT)	PHOTOMETRY	26.4	U/L	< 35
ALANINE TRANSAMINASE (SGPT)	PHOTOMETRY	23	U/L	< 45
PROTEIN - TOTAL	PHOTOMETRY	7.04	gm/dL	5.7-8.2
ALBUMIN - SERUM	PHOTOMETRY	4.51	gm/dL	3.2-4.8
SERUM GLOBULIN	CALCULATED	2.53	gm/dL	2.5-3.4
SERUM ALB/GLOBULIN RATIO	CALCULATED	1.78	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

OT/PT - Derived from SGOT and SGPT values.

SGOT - IFCC* Without Pyridoxal Phosphate Activation

SGPT - IFCC* Without Pyridoxal Phosphate Activation

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



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TEST NAME		TECHNOLOGY	VALUE	UNITS	
MAGNESIUM	rval	PHOTOMETRY	2.41	mg/dL	

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









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NAME	: XXXXXXXXX XXXXXXXXX XXXX
REF. BY	: SELF

: MASTER WELLNESS CHECKUP

HOME COLLECTION : xxxxxxxxx xxxxxxx

XXXXXXXXX XXXXXXXXXX

xxxx xxxxxx xxxxxxxxxxxxxx

PATIENTID : MD22463706

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
BLOOD UREA NITROGEN (BUN)	PHOTOMETRY	8.49	mg/dL	7.94 - 20.07
UREA (CALCULATED)	CALCULATED	18.17	mg/dL	Adult : 17-43
CREATININE - SERUM	PHOTOMETRY	0.51	mg/dL	0.72-1.18
UREA / SR.CREATININE RATIO	CALCULATED	35.63	Ratio	< 52
BUN / SR.CREATININE RATIO	CALCULATED	16.65	Ratio	9:1-23:1
CALCIUM	PHOTOMETRY	9.75	mg/dL	8.8-10.6
URIC ACID	PHOTOMETRY	3.07	mg/dL	4.2 - 7.3
SODIUM	I.S.E - INDIRECT	141.3	mmol/L	136 - 145
CHLORIDE	I.S.E - INDIRECT	107.7	mmol/L	98 - 107

Please correlate with clinical conditions.

Method :

TEST ASKED

BUN - Kinetic UV Assay. UREAC - Derived from BUN Value. SCRE - Creatinine Enzymatic Method UR/CR - Derived from UREA and Sr.Creatinine values. B/CR - Derived from serum Bun and Creatinine values CALC - Arsenazo III Method, End Point. URIC - Uricase / Peroxidase Method SOD - ION SELECTIVE ELECTRODE - INDIRECT CHL - ION SELECTIVE ELECTRODE - INDIRECT

Sample Collected on (SCT)	:		
Sample Received on (SRT)	:		
Report Released on (RRT)	:		
Sample Type	:		
Labcode	:		
Barcode	:		

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NAME	:	XXXXXXXXX XXXXXXXXX XXXX
REF. BY	:	SELF
TEST ASKED	:	MASTER WELLNESS CHECKUP

HOME COLLECTION : XXXXXXXXX XXXXXXXXX XXXX XXXXXX ***** ***** ***** XXXXXXXXX XXXXXXXXX

PATIENTID : MD22463706				
TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
TOTAL TRIIODOTHYRONINE (T3)	E.C.L.I.A	149	ng/dL	80-210
TOTAL THYROXINE (T4)	E.C.L.I.A	9.57	µg/dL	4.7-12.4
TSH - ULTRASENSITIVE	E.C.L.I.A	1.42	µIU/mL	0.72-5.77

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

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

References :

1. Elmlinger MW, Kuhnel W, Lambretch HG, et al. Reference intervals from birth to adulthood for serum thyroxine, T3, free T3, Free T4, TBG and TSH. Clin Chem lab med. 2001; 39:973

2. Edward CC, Carlo B. Paediatric Reference Intervals. 8th edition. 2021

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.







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NAME	: XXXXXXXXX XXXXXXXXX XXXX	HOME COLLECTION :
REE BY	• SELE	XXXXXXXXX XXXXXXXXX
TEST ASKED : MASTER WELLNESS CHECKUP	XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	
	: MASTER WELLNESS CHECKUP	XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
		Χχχχχχχχ χχχχχχχχ

PATIENTID : MD22463706 **TEST NAME** TECHNOLOGY VALUE UNITS Bio. Ref. Interval. ARSENIC ICP-MS 2.58 µg/L < 5 CADMIUM ICP-MS 1.09 µg/L < 1.5 MERCURY ICP-MS 1 µg/L < 5 ICP-MS LEAD 37.47 µg/L < 150 CHROMIUM ICP-MS 1.77 µg/L < 30 BARIUM ICP-MS 0.59 µg/L < 30 ICP-MS µg/L COBALT 0.39 0.10 - 1.50 CAESIUM ICP-MS 2.4 < 5 µg/L THALLIUM ICP-MS 0.01 < 1 µg/L URANIUM ICP-MS 0.04 µg/L < 1 STRONTIUM ICP-MS 20.14 µg/L 8 - 38 ANTIMONY ICP-MS 4.01 µg/L 0.10 - 18 TIN ICP-MS 1.2 µg/L < 2 MOLYBDENUM ICP-MS 1.41 µg/L 0.70 - 4.0 SILVER ICP-MS 0.68 < 4 µg/L VANADIUM ICP-MS 0.16 µg/L < 0.8 BERYLLIUM ICP-MS 0.08 µg/L 0.10 - 0.80 BISMUTH ICP-MS 0.1 µg/L 0.10 - 0.80 SELENIUM ICP-MS 221.21 µg/L 60 - 340 ALUMINIUM ICP-MS 6.43 µg/L < 30 NICKEL ICP-MS 3.87 µg/L < 15 MANGANESE ICP-MS 9.24 7.10 - 20 µg/L

Please correlate with clinical conditions.

Method :

ICP - MASS SPECTROMETRY Note:Reference range has been obtained after considering 95% population as cutoff.



X







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NAME REF. BY TEST ASKED	: XXXXXXXXX XXXXXXXXXX XXXX : SELF : MASTER WELLNESS CHECKUP			HOME COLLECTION	l: xx xxxx xxxxxx xxxxxxxx xxxx xxxxxx xx
TEST NAME	: MD22463706	TECHNOL	002		
	`	TECHNOL	UGT	VALUE	UNITS
HDAIC - (HPLC)	H.P.L.C		4.9	%
Bio. Ref. Inte Bio. Ref. Inte	erval. : erval.: As per ADA Guidelines		Guidan	ice For Known Dia	betics
Below 5.7% 5.7% - 6.4% >=6.5%	: Normal : Prediabetic : Diabetic		Below 6 6.5% - 7.0% - >8%	5.5% : Good Control 7% : Fair Control 8% : Unsatisfactor : Poor Control	y Control
Method : Fully	Automated H.P.L.C method		FD		/ II
AVERAGE BLO Bio. Ref. Inte	OD GLUCOSE (ABG) erval.:	CALCOLAI	20	94	mg/dL
90 - 120 mg/ 121 - 150 mg 151 - 180 mg > 180 mg/dl	dl : Good Control /dl : Fair Control J/dl : Unsatisfactory Control : Poor Control				
method : Deriv					

Please correlate with clinical conditions.

Sample Collected on (SCT)

:

:

:

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

Barcode

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ERYTHROCYT Bio. Ref. Inte	E SEDIMENTATION RATE (ESR) rval.:-	MODIFIED WESTERGREN	25	mm / hr
TEST NAME		TECHNOLOGY	VALUE	UNITS
TEST ASKED	: MASTER WELLNESS CHECKUP	Xxxxxxx xxxx XXXX	XXXXX XXXX XX	xxxx xxxx xxxxxxxxx
REF. BY	: SELF		xxxx xxxx Xxxxxxxxx	
NAME	: XXXXXXXXX XXXXXXXXX XXXX	HOME COLLECT	ION :	

Male : 0-15 Female : 0-20

Clinical Significance:

- An erythrocyte sedimentation rate (ESR) is a blood test that can rise if you have inflammation in your body. Its also used as a marker to monitor prognosis of an existing inflammatory/infective condition.
- Inflammation is your immune systems response to injury, infection, and many types of conditions, including immune system disorders, certain cancers and blood disorders.
- A high ESR test result may be from a condition that causes inflammation, such as: Arteritis, Arthritis, Systemic vasculitis, Polymyalgia rheumatica, Inflammatory bowel disease, Kidney disease, Infections like Tuberculosis etc, Rheumatoid arthritis and other autoimmune diseases, Heart disease, Certain cancers and many other Conditions.
- A low ESR test result may be caused by conditions such as: A blood disorder, such as: Polycythemia, Sickle cell disease (SCD), Leukocytosis, Heart failure, Certain kidney and liver problems etc.
- Certain physiological conditions also affect ESR results, these include : Pregnancy, menstrual cycle, ageing, obesity, drinking alcohol regularly, and exercise, Certain medicines and supplements also can affect ESR results.
- Hence Its always suggested to interpret ESR results in conjunction with Clinical History and other findings.

References :

https://medlineplus.gov/lab-tests/erythrocyte-sedimentation-rate-esr/

Please correlate with clinical conditions.

Method:- MODIFIED WESTERGREN



NAME





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:	XXXXXXXXXX XXXXXXXXXX XXXX

REF. BY	:	SELF
TEST ASKED	:	MASTER WELLNESS CHECKUP

HOME COLLECTION :

PATIENTID : MD22463706

TEST NAME	METHODOLOGY			
		VALUE	UNITS	Bio. Ref. Interval.
HEMOGLOBIN	SLS-Hemoglobin Method	12.6	g/dL	13.0-17.0
Hematocrit (PCV)	CPH Detection	40	%	40.0-50.0
Total RBC	HF & EI	4.62	X 10^6/µL	4.5-5.5
Mean Corpuscular Volume (MCV)	Calculated	86.6	fL	83.0-101.0
Mean Corpuscular Hemoglobin (MCH)	Calculated	27.3	pq	27.0-32.0
Mean Corp.Hemo. Conc (MCHC)	Calculated	31.5	g/dL	31.5-34.5
Red Cell Distribution Width - SD (RDW-SD)	Calculated	40.1	fL	39-46
Red Cell Distribution Width (RDW - CV)	Calculated	12.9	%	11.6-14.0
RED CELL DISTRIBUTION WIDTH INDEX (RDWI)	Calculated	241.8	-	*Refer Note below
MENTZER INDEX	Calculated	18.7	-	*Refer Note below
TOTAL LEUCOCYTE COUNT (WBC)	HF & FC	9.09	X 10 ³ / μL	4.0 - 10.0
DIFFERENTIAL LEUCOCYTE COUNT				
Neutrophils Percentage	Flow Cytometry	52.6	%	40-80
Lymphocytes Percentage	Flow Cytometry	37.6	%	20-40
Monocytes Percentage	Flow Cytometry	2.9	%	2-10
Eosinophils Percentage	Flow Cytometry	5.6	%	1-6
Basophils Percentage	Flow Cytometry	1.1	%	0-2
Immature Granulocyte Percentage (IG%)	Flow Cytometry	0.2	%	0.0-0.5
Nucleated Red Blood Cells %	Flow Cytometry	0.01	%	0.0-5.0
ABSOLUTE LEUCOCYTE COUNT				
Neutrophils - Absolute Count	Calculated	4.78	X 10³ / μL	2.0-7.0
Lymphocytes - Absolute Count	Calculated	3.42	Χ 10³ / μL	1.0-3.0
Monocytes - Absolute Count	Calculated	0.26	X 10 ³ / µL	0.2 - 1.0
Basophils - Absolute Count	Calculated	0.1	X 10 ³ / μL	0.02 - 0.1
Eosinophils - Absolute Count	Calculated	0.51	Χ 10³ / μL	0.02 - 0.5
Immature Granulocytes (IG)	Calculated	0.02	X 10 ³ / μL	0.0-0.03
Nucleated Red Blood Cells	Calculated	0.01	X 10 ³ / μL	0.0-0.5
PLATELET COUNT	HF & EI	433	Χ 10³ / μL	150-410
Mean Platelet Volume (MPV)	Calculated	9	fL	7.5-8.3
Platelet Distribution Width (PDW)	Calculated	8.8	fL	9.6-15.2
Platelet to Large Cell Ratio (PLCR)	Calculated	17	%	19.7-42.4
Plateletcrit (PCT)	Calculated	0.39	%	0.19-0.39

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

*Note - Mentzer index (MI), RDW-CV and RDWI are hematological indices to differentiate between Iron Deficiency Anemia (IDA) and Beta Thalassemia Trait (BTT). MI >13, RDWI >220 and RDW-CV >14 more likely to be IDA. MI <13, RDWI <220, and RDW-CV <14 more likely to be BTT. Suggested Clinical correlation. BTT to be confirmed with HB electrophoresis if clinically indicated.

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)





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NAME	: XXXXXXXXX XXXXXXXXX XXXX
REF. BY	: SELF
TEST ASKED	: MASTER WELLNESS CHECKUP

TEST NAME	TECHNOLOGY	VALUE	UNITS
DIABETES SCREEN (URINE)			
URINARY MICROALBUMIN	PHOTOMETRY	< 5.5	µg/mL
Bio. Ref. Interval. : Adults: Less than 25 μg/ml			
Method : Fully Automated Immuno Turbidometry			
CREATININE - URINE	PHOTOMETRY	23.28	mg/dL
Bio. Ref. Interval. : Male: 39 - 259 mg/dl Female: 28 - 217 mg/dl			
Method : Creatinine Jaffe Method, Rate-Blanked and Con	npensated		
URI. ALBUMIN/CREATININE RATIO (UA/C)	CALCULATED	23.6	µg/mg of Creatinine
Bio. Ref. Interval. : Adults : Less than 30 μg/mg of Creatinine			
Method : Derived from Albumin and Creatinine values			

Please correlate with clinical conditions.

Sample Collected on (SCT) Sample Received on (SRT) Report Released on (RRT) Sample Type

Labcode

Barcode



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NAME	: XXXXXXXXX XXXXXXXXX XXXX
REF. BY	: SELF
TEST ASKED	: MASTER WELLNESS CHECKUP

HOME COLLECTION : ***** ***** xxxxxxxxxxxx Xxxxxx Xxxxxxx xxxxxxx

TEST NAME	METHODOLOGY	VALUE	UNITS	Bio. Ref. Interval.
Complete Urinogram				
Physical Examination				
SPECIFIC GRAVITY	pKa change	1.015	-	1.003-1.030
PH	pH indicator	6	-	5-8
Chemical Examination				
URINARY GLUCOSE	GOD-POD	ABSENT	mg/dL	Absent
URINARY BILIRUBIN	Diazo coupling	ABSENT	mg/dL	Absent
URINE KETONE	Nitroprusside	ABSENT	mg/dL	Absent
URINE BLOOD	Peroxidase reaction	ABSENT	-	Absent
URINARY PROTEIN	PEI	ABSENT	mg/dL	Absent
UROBILINOGEN	Diazo coupling	Normal	mg/dL	<=0.2
NITRITE	Diazo coupling	ABSENT	-	Absent
Microscopic Examination				
URINARY LEUCOCYTES (PUS CELLS)	Microscopy	1	cells/HPF	0-5

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

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NAME	: XXXXXXXXX XXXXXXXXXX XXXX	HOME COL	LECTION :	
REF. BY	: SELF	xxxxxxx x	xxxxxxxx xxxx xxx	xxxx xxxx Xxxxxxxxx
TEST ASKED	: MASTER WELLNESS CHECKUP	xxxxxxxx x Xxxxxxxx x	xxx xxxxxx xxxx x xxxxxxxx xxxx xxx	xxxxxxxxxx xxxx xxxx xxxxxxxxxx
TEST NAME		TECHNOLOGY	VALUE	UNITS
FASTING BLOC	DD SUGAR(GLUCOSE)	PHOTOMETRY	94.9	mg/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		

Note :

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

~~ End of report ~~



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: (a) 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, (b) any claims of any nature whatsoever arising from or relating to the performance of the requested tests as well as any claim for indirect, incidental or consequential damages. The total liability, in any case, of Thyrocare shall not exceed the total amount of invoice for the services provided and paid for.
- v Thyrocare Discovery video link :- <u>https://youtu.be/nbdYeRgYyQc</u>

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

- v 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, clinical support or feedback, write to us at **customersupport@thyrocare.com** or call us on **022-3090 0000**



+T&C Apply, #As on 5th December 2024, *As per a survey on doctors' perception of laboratory diagnostics (IJARIIT, 2023)