

Age

Jeevan Jyoti HLM

Pathkind Diagnostics Pvt. Ltd.

162, Lowther Road, Bai Ka Bagh, Prayagraj

PATHKIND REFERENCE LAB PATHKIND DIAGNOSTICS PVT. LTD.

Plot No. 55-56, Udyog Vihar, Phase IV, Sector-18, Gurugram-122015 E-Mail: care@pathkindlabs.com | Website: www.pathkindlabs.com Customer Care: 75000 75111

Processed By

Pathkind Diagnostics Pvt. Ltd.

162, Lowther Road, Bai Ka Bagh, Prayagraj

Uttar Pradesh-211003

: Mrs. PRATIMA VIASHYA REG-315077 ECHS Name

: 33 Yrs

Sex : Female : P1212100008608 P. ID No.

Accession No : 121222028678

Referring Doctor: MR SELF

Referred By

Billing Date

16/01/202311:24:26

Sample Collected on

16/01/2023 14:41:00

Sample Received on

16/01/2023 15:02:21

Report Released on

16/01/2023 15:23:53

gm/dL

thou/µL

million/µL

fL

pg

g/dL

%

%

Barcode No.

994858101

Ref no.

12.0 - 15.0

4.0 - 10.0

3.8 - 4.8

36.0 - 46.0

83.0 - 101.0

27.0 - 32.0

31.5 - 34.5

11.9 - 15.5

Report Status - Preliminary Report

Test Name Result **Biological Ref. Interval** Unit

11.9 L

9.6

4.1

36.6

89.6

29.2

32.6

12.6

64

27

HAEMATOLOGY

Complete Blood Count (CBC)

Ha	en	nog	gle	obi	n	(Hb
_						

Sample: Whole Blood EDTA Method: Photometric measurement

Total WBC Count / TLC

Sample: Whole Blood EDTA Method: Impedance

RBC Count Sample: Whole Blood EDTA

Method: Impedance

PCV / Hematocrit

Sample: Whole Blood EDTA Method: Impedance

Sample: Whole Blood EDTA

Method: Calculated

Sample: Whole Blood EDTA Method: Calculated

MCHC

Sample: Whole Blood EDTA

RDW (Red Cell Distribution Width)

Sample: Whole Blood FDTA Method: Calculated

DLC (Differential Leucocyte Count)

Method: Flowcytometry/Microscopy

Neutrophils Sample: Whole Blood EDTA

Method: VCS Technology & Microscopy

Lymphocytes

Sample: Whole Blood EDTA Method: VCS Technology & Microscopy

40 - 80

20 - 40

%

121222028678 Mrs. PRATIMA VIASHYA REG-3150









from the promoters of





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st Name	Result	Biological Ref. Interval	Unit
Eosinophils Sample: Whole Blood EDTA Method: VCS Technology & Microscopy	05	01 - 06	%
Monocytes Sample: Whole Blood EDTA Method: VCS Technology & Microscopy	04	02 - 10	%
Basophils Sample: Whole Blood EDTA Method: VCS Technology & Microscopy	00	00 - 02	%
Absolute Neutrophil Count Sample: Whole Blood EDTA	6144	2000 - 7000	/μL
Absolute Lymphocyte Count Sample: Whole Blood EDTA	2592	1000 - 3000	/μL
Absolute Eosinophil Count Sample: Whole Blood EDTA	480	20 - 500	/μL
Absolute Monocyte Count Sample: Whole Blood EDTA	384	200 - 1000	/μL
Absolute Basophil Count Sample: Whole Blood EDTA	0 L	20 - 100	/μL
DLC Performed By Sample: Whole Blood EDTA	EDTA Smear		
Platelet Count Sample: Whole Blood EDTA Method: Impedance	273	150 - 410	thou/μL
MPV (Mean Platelet Volume) Sample: Whole Blood EDTA Method: Calculated	10.1	6.8 - 10.9	fL
Sample: Whole Blood EDTA ythrocyte Sedimentation Rate (ESR)	26 H	<12	mm 1st Hour

Sample: Whole Blood EDTA

Method: Modified Westergren Method









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Test Name	Result	Biological Ref. Interval	Unit
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Blood Group

"O" **Blood Grouping**

Sample: Whole Blood EDTA

POSITIVE Rh (D) Typing

Sample: Whole Blood EDTA

BIOCHEMISTRY

Fasting Plasma Glucose Sample: Fluoride Plasma - F	100	74 - 106	mg/dl
Glucose Post-Prandial Sample: Fluoride Plasma - PP Method: Hexokinase	108	70 - 140	mg/dl
Thyroid Profile Total Total T3 (Triiodothyronine) Sample: Serum Method: ECLIA	1.32	0.80 - 2.00	ng/mL
Total T4 (Thyroxine)	7.94	5.10 - 14.10	μg/dL

Total T4 (Thyroxine) Sample: Serum

TSH 3rd Generation 4.680 H 0.270 - 4.200 μIU/mL

Sample: Serum Method: ECLIA

Method: ECLIA









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Test Name	Result	Biological Ref. Interval	Unit
<u>Liver Function Test (LFT)</u>			
Bilirubin Total Sample: Serum Method: Spectrophotometery	0.4	<1.1	mg/dL
Bilirubin Direct Sample: Serum Method: Spectrophotometery	0.1	<0.2	mg/dL
Serum Bilirubin (Indirect) Sample: Serum Method: Calculated	0.3	<0.90	mg/dL
SGOT / AST Sample: Serum Method: Spectrophotometery	20	<31	U/L
SGPT / ALT Sample: Serum Method: Spectrophotometery	22	<33	U/L
AST / ALT Ratio Sample: Serum Method: Calculated	0.91		
Alkaline Phosphatase (ALP) Sample: Serum Method: Spectrophotometery	133 H	<98	U/L
Total Protein Sample: Serum Method: Spectrophotometry	8.0	6.4 - 8.3	g/dL
Albumin Sample: Serum Method: Spectrophotometery	4.9	4.0 - 4.9	g/dL
Globulin Sample: Serum Method: Calculated	3.1	1.9 - 3.7	g/dL
Albumin/Globulin (A/G) Ratio Sample: Serum	1.6	1.0 - 2.1	g/dL

121222028678 Mrs. PRATIMA VIASHYA REG-3150







Method: Calculated









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Preliminary Report

Test Name	Result	Biological Ref. Interval	Unit
ipid Profile			
Total Cholesterol Sample: Serum Method: Spectrophotometery	210 H	No risk : < 200 Moderate risk : 200–239 High risk : =240	mg/dL
Triglycerides Sample: Serum Method: Spectrophotometry	198	Desirable : < 150 Borderline High : 150 - 199 High : 200 - 499 Very High : >/= 500	mg/dL
LDL Cholesterol (Calculated) Sample: Serum Method: Calculated	123 H	Optimal : <100 Near Optimal : 100 - 129 Borderline High : 130 - 160 High : 161 - 189 Very High : >/=190	mg/dL
HDL Cholesterol Sample: Serum Method: Spectrophometry	47	Low : < 40 Optimal : 40 - 60 High : > 60	mg/dl
Non HDL Cholesterol Sample: Serum	163 H	< 130	mg/dL
VLDL Cholesterol Sample: Serum Method: Calculated	39.6 Н	Desirable 10 - 35	mg/dL
Total Cholesterol / HDL Ratio Sample: Serum Method: Calculated	4.47 H	Low Risk : 3.3 - 4.4 Average Risk : 4.5 - 7.0 Moderate Risk : 7.1 - 11.0 High Risk : > 11.0	
LDL / HDL Ratio Sample: Serum Method: Calculated	2.6	0.5 - 3.0	
		Low Risk : 0.5 - 3.0 Moderate Risk : 3.1 - 6.0 High Risk : > 6.0	

Kidney Profile (KFT)

Blood Urea









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: 33 Yrs : Female

Sex : Female
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Defender Destant MAD CELE

Referring Doctor: MR SELF

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Report Status - Preliminary Report

Test Name	Result Biological Ref. Interval		Unit
Blood Urea Nitrogen (BUN) Sample: Serum Method: Spectrophotometry-Urease / GLDH	9.50	7.00 - 18.69	mg/dL
Urea Sample: Serum Method: Spectrophotometery	20.33	17.00 - 43.00	mg/dL
Creatinine Sample: Serum Method: Spectrophotometry	0.53	0.50 - 1.10	mg/dL
BUN Creatinine Ratio Sample: Serum Method: Calculated	18	10 - 20	
Calcium Sample: Serum Method: Spectrophotometery	9.9	8.6 - 10.0	mg/dL
Uric Acid Sample: Serum Method: Spectrophotometery	4.9	2.4 - 5.7	mg/dL
Total Protein Sample: Serum Method: Spectrophotometry	8.0	6.4 - 8.3	g/dL
Albumin Sample: Serum Method: Spectrophotometery	4.9	4.0 - 4.9	g/dL
Globulin Sample: Serum Method: Calculated	3.1	1.9 - 3.7	g/dL
Albumin/Globulin (A/G) Ratio Sample: Serum	1.6	1.0 - 2.1	g/dL

Sample: Serum Method: Calculated









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Test Name Result **Biological Ref. Interval** Unit

CLINICAL PATHOLOGY

Urine Routine & Microscopic Examination

Method: Reflectance Photometry

Physical Examination

Colour

Sample: Urine Method: Physical Examination

Appearance

Sample: Urine Method: Physical Examination

Specific Gravity

Sample: Urine

Method: pKa change of pretreated polyelectrolytes

pН

yellow

6.0

Slightly Hazy Clear

1.015 1.003 - 1.035

Sample: Urine

Method: Double indicator principle

Chemical Examination

Glucose Sample: Urine

Method: Glucose oxidase/peroxidase

Protein

Sample: Urine

Method: Protein-error-of-indicators principle

Ketones

Sample: Urine

Method: Sodium nitroprusside reaction

Sample: Urine Method: Peroxidase

Bilirubin

Sample: Urine Method: Diazo reaction

Not Detected

Not Detected

4.7 - 7.5

Pale Yellow

Not Detected

Not Detected

Not Detected

Detected

Not Detected

Trace

Not Detected

Not Detected



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Test Name	Result	Biological Ref. Interval	Unit
Urobilinogen Sample: Urine Method: Ehrlich's reaction	Normal	Normal	
Nitrite Sample: Urine Method: Nitrite Test	Not Detected	Not Detected	
Microscopic Examination Method: Microscopy			
Pus Cells Sample: Urine	2 - 3	0 - 5	/hpf
RBC Sample: Urine	1 - 2	Not Detected	/hpf
Epithelial Cells Sample: Urine	3 - 5	0 - 5	/hpf
Casts Sample: Urine	Not Detected	Not Detected	/hpf
Crystals Sample: Urine	Not Detected	Not Detected	/hpf
Bacteria Sample: Urine	Not Detected	Not Detected	/hpf
Remarks Sample: Urine			

Remarks: Microscopic Examination is performed on urine sediment

BIOCHEMISTRY

Electrolytes (Na/K/CI)

Sodium 139 136 - 145 mmol/L

Sample: Serum Method: ISE













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Test Name	Result	Biological Ref. Interval	Unit
Potassium Sample: Serum Method: ISE	4.4	3.5 - 5.1	mmol/L
Chloride Sample: Serum Method: ISE	107	97 - 107	mmol/L

Complete Blood Count (CBC)

Clinical Significance:

CBC comprises of estimation of the cellular components of blood including RBCs, WBCs and Platelets. Mean corpuscular volume (MCV) is a measure of the size of the average RBC, MCH is a measure of the hemoglobin cointent of the average RBC and MCHC is the hemoglobin concentration per RBC. The red cell distribution width (RDW) is a measure of the degree of variation in RBC size (anisocytosis) and is helpful in distinguishing between some anemias. CBC examination is used as a screening tool to confirm a hematologic disorder, to establish or rule out a diagnosis, to detect an unsuspected hematologic disorder, or to monitor effects of radiation or chemotherapy. Abnormal results may be due to a primary disorder of the cell-producing organs or an underlying disease. Results should be interpreted in conjunction with the patient's clinical picture and appropriate additional testing performed.

Erythrocyte Sedimentation Rate (ESR)

Clinical Significance

The erythrocyte sedimentation rate (ESR) is a simple but non-specific test that helps to detect inflammation associated with conditions such as infections, cancers, and autoimmune diseases.

Total T3 (Triiodothyronine)

Clinical Significance:

Thyroid hormones, T3 and T4, which are secreted by the thyroid gland, regulate a number of developmental, metabolic, and neural activities throughout the body. The thyroid gland synthesizes 2 hormones - T3 and T4. T3 production in the thyroid gland constitutes approximately 20% of the total circulating T3, 80% being produced by peripheral conversion from T4. T3 is more potent biologically. Total T3 comprises of Free T3 and bound T3. Bound T3 remains bound to carrier proteins like thyroid-binding globulin, prealbumin, and albumin). Only the free forms are metabolically active. In hyperthyroidism, both T4 and T3 levels are usually elevated, but in some rare cases, only T3 elevation is also seen. In hypothyroidism T4 and T3 levels are frequently low in sick or hospitalized euthyroid patients.

Total T4 (Thyroxine)









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			· · · · · ·

Clinical Significance:

Referred By

Total T4 is synthesized in the thyroid gland. About 0.05% of circulating T4 is in the free or biologically active form. The remainder is bound to thyroxine-binding globulin (TBG), prealbumin, and albumin. High levels of T4 (and FT4) causes hyperthroidism and low levels lead to hypothyroidism.

TSH 3rd Generation

Clinical Significance:

TSH levels are elevated in primary hyporthyroidism and low in primary hyperthyroidism. Evaluation of TSH is useful in the differential diagnosis of primary from secondary and tertiary hypothyroidism. In primary hypothyroidism, TSH levels are elevated, while in secondary and tertiary hypothyroidism, TSH levels are low or normal. High TSH level in the presence of normal FT4 is called subclinical hypothyroidism and low TSH with normal FT4 is called subclinical hyperthyroidism. Sick, hospitalized patients may have falsely low or transiently elevated TSH. Significant diurnal variation is also seen in TSH levels.

Guidelines for TSH levels in pregnancy, as per American Thyroid Association, are as follows:

PREGNANCY TRIMESTER	BIOLOGICAL REFERENCE INTERVAL	UNIT
FIRST TRIMESTER	0.100 - 2.500	μIU/mL
SECOND TRIMESTER	0.200 - 3.000	μIU/mL
THIRD TRIMESTER	0.300 - 3.000	uIU/mL

Bilirubin Total

Clinical Significance:

"Total Bilirubin is one of the most commonly used tests to assess liver function. A number of inherited and acquired diseases affect bilirubin production, metabolism, storage and excretion and causes hyperbilirubinemia resulting in jaundice. Hyperbilirubinemia may be due to increased bilirubin production (eg, hemolysis and ineffective erythropoiesis), decreased bilirubin excretion (eg, obstruction and hepatitis), and abnormal bilirubin metabolism (eg, hereditary and neonatal jaundice). Unconjugated hyperbilirubinemia is seen in newborn andd known as physiological jaundice. Elevated unconjugated bilirubin in the neonatal period may result in brain damage (kernicterus). Crigler-Najjar syndromes type I and type II are also associated with elevated levels of indirect bilirubin. Both conjugated and unconjugated bilirubin are increased in hepatitis and space-occupying lesions of the liver; and obstructive lesions such as carcinoma of the head of the pancreas, common bile duct, or ampulla of Vater."

Bilirubin Direct











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Clinical Significance:

"Direct bilirubin is a measurement of conjugated bilirubin. Jaundice can occur as a result of increased bilirubin production (eg, hemolysis and ineffective erythropoiesis), decreased bilirubin excretion (eg, obstruction and hepatitis), and abnormal bilirubin metabolism (eg, hereditary and neonatal jaundice). Inherited disorders in which direct bilirubin levels are increased are seen in Dubin-Johnson syndrome and Rotor syndrome, idiopathic neonatal hepatitis and biliary atresia. The most commonly occurring form of jaundice of the newborn called physiological jaundiceis due to increase in levels of indirect bilirubin.Both conjugated and unconjugated bilirubin are increased in hepatocellular diseases such as hepatitis and space-occupying lesions of the liver, bstructive lesions such as carcinoma of the head of the pancreas, common bile duct, or ampulla of Vater."

SGOT / AST

Clinical Significance:

"Elevated aspartate aminotransferase (AST) values are seen most commonly in parenchymal liver diseases. Values can be elevated from 10 to 100 times the normal range, though commonly 20 to 50 times elevations are seen. AST levels are raised in infectious hepatitis and other inflammatory conditions affecting the liver along with ALT, though ALT levels are higher. The ALT:AST ratio which is normally <1 is reversed in these conditions and becomes >1. AST levels are usually raised before clinical signs and symptoms of disease appear. AST and ALT also rise in primary or metastatic carcinoma of the liver, with AST usually being higher than ALT. Elevated AST values may also be seen in disorders affecting the heart, skeletal muscle and kidney, such as myocardial infarction, muscular dystrophy, dermatomyositis, acute pancreatitis and crushed muscle injuries."

SGPT / ALT

Clinical Significance:

Elevated alanine aminotransferase (ALT) values are seen in parenchymal liver diseases characterized by a destruction of hepatocytes. Values are at least 10 times higher the normal range and may reach up to 100 times the upper reference limit. Commonly, values are seen to be 20 - 50 times higher than normal. In infectious hepatitis and other inflammatory conditions affecting the liver, ALT levels rise more than aspartate aminotransferase (AST), and the ALT/AST ratio, which is normally <1, is reversed and becomes >1. ALT levels usually rise before clinical signs and symptoms of disease appear.

Alkaline Phosphatase (ALP)

Clinical Significance:

Alkaline Phosphatase levels can be elevated in both liver related as well as bone related conditions. ALP levels are raised (more than 3













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Uttar Pradesh-211003

: Mrs. PRATIMA VIASHYA REG-315077 ECHS 16/01/202311:24:26 Name Billing Date : 33 Yrs 16/01/2023 14:41:00 Age Sample Collected on Sex : Female Sample Received on 16/01/2023 15:02:21 P. ID No. : P1212100008608 Report Released on 16/01/2023 15:23:53 **Accession No** : 121222028678 Barcode No. 994858100, 994858151,

994858102, 994858091,

994858101 Ref no.

Report Status - Preliminary Report

Test Name Result **Biological Ref. Interval** Unit

fold) in extrahepatic biliary obstruction (eg, by stone or by cancer of the head of the pancreas) than in intrahepatic obstruction, and is directly proportional to the level of obstruction. Levels may rise up to 10 to 12 times the upper limit of normal range and returns to normal on surgical removal of the obstruction. ALP levels rise together with GGT levels and If both GGT and ALP are elevated, a liver source of the ALP is likely. Among bone diseases, ALP levels rise in Paget disease (up to 25 fold), osteomalacia, rickets, primary and secondary hyperparathyroidism and osteogenic bone cancer. Elevated ALP is seen in children following accelerated bone growth. Also, a 2 to 3fold elevation may be observed in women in the third trimester of pregnancy, although the interval is very wide and levels may not exceed the upper limit of the reference interval in some cases.

Total Protein

Clinical Significance:

High levels of Serum Total Protein is seen in increased acute phase reactants in inflammation, late-stage liver disease, infections, multiple myeloma and other malignant paraproteinemias.n. Hypoproteinemia is seen in hypogammaglobulinemia, nephrotic syndrome and protein-losing enteropathy.

Albumin

Clinical Significance:

"Hypoalbuminemia can be caused by impaired synthesis due to liver disease (primary) or due to diminished protein intake (secondary), increased catabolism due to tissue damage and inflammation; malabsorption of amino acids; and increased renal excretion (eg, nephrotic syndrome). Hyperalbuminemia is seen in dehydration."

Lipid Profile

Proposed LDL-C goals in very high risk and extreme risk group patients by the Lipid Association of India.

Very High Risk group(VHRG)	Extreme Risk group		
	Category A	Category B	
LDL-C goal of <50 mg/dl	LDL-C goal of <50 mg/dl (recommended) LDL-C goal of ≤30 mg/dl (optional)	LDL-C goal of ≤30 mg/dl	
High-risk conditions		CAD with ≥ 1 of following:	













Sex

Jeevan Jyoti HLM

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Plot No. 55-56, Udyog Vihar, Phase IV, Sector-18, Gurugram-122015 E-Mail: care@pathkindlabs.com | Website: www.pathkindlabs.com Customer Care: 75000 75111

Processed By

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Uttar Pradesh-211003

Name : Mrs. PRATIMA VIASHYA REG-315077 ECHS
Age : 33 Yrs

: 33 Yrs : Female

P. ID No. : P1212100008608

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Referring Doctor: MR SELF

Referred By :

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Any one of following:

1. ASCVD (CAD/PAD/TIA or stroke)

- 2. Homozygous familial
- 3. hypercholesterolemia
- 4. Diabetes with ≥2 major ASCVD risk factors*/target organ damage

CAD with ≥ 1 of following:

- Diabetes without target organ damage/≤1 major
- 2. ASCVD risk factors
- 3. Familial hypercholesterolemia
- 4. ≥3 major ASCVD risk factors
- 5. CKD stage 3B and 4
- 6. ≥2 major ASCVD risk factors with ≥1 moderate
- 7. non-conventional risk factor#
- 8. $Lp(a) \ge 50 \text{ mg/dl}$
- 9. Coronary calcium score ≥300 HU
- 10. Extreme of a single risk factor
- 11. PAD
- 12. H/o TIA or stroke
- 13. Non-stenotic carotid plaque

- 1. Diabetes + polyvascular disease/≥2
- 2. major ASCVD risk factors*/target organ
- 3. damage
- 4. Recurrent ACS (within 12 months)
- 5. despite on LDL-C goal
- 6. Homozygous familial
- 7. Hypercholesterolemia

The LDL-C goal of ≤30 mg/dl must be pursued after detailed risk-benefit discussion between physician and patient.

Clinical judgment to be used in decision making if the patient has disease/risk factors not covered in the table, eg. peripheral arterial disease or cerebrovascular disease.

*Major ASCVD risk factors: 1. Age- male ≥45 years, female ≥55 years, 2. Family h/o premature CAD- male <55 years, female <65 years, 3.

Uric Acid

Clinical Significance:

Uric acid is the final product of purine metabolism. Serum uric acid levels are raised in case of increased purine synthesis, inherited metabolic disorder, excess dietary purine intake, increased nucleic acid turnover, malignancy and cytotoxic drugs. Decreased levels are seen in chronic renal failure, severe hepatocellular disease with reduced purine synthesis, defective renal tubular reabsorption, overtreatment of hyperuricemia with allopurinol, as well as















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some cancer therapies.

Referred By

Urine Routine & Microscopic Examination

Urine routine examination and microscopy comprises of a set of screening tests that can detect some common diseases like urinary tract infections, kidney disorders, liver problems, diabetes or other metabolic conditions. Physical characteristics (colour and appearance), chemical composition (glucose, protein, ketone, blood, bilirubin and urobilinogen) and microscopic content (pus cells, epithelial cells, RBCs, casts and crystals) are analyzed and reported.

** End of Report**

Dr. Ankit Singh

MBBS, MD (Pathologist) Lab Head







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HEART	Lipid Profile	Lipid Profile with Direct LDL	Lipid Profile with Direct LDL	
DIABETES	FBS, HbA1c	FBS, HbA1c, Microalbumin	FBS, HbA1c, Microalbumin	
KIDNEY	BUN, Creatinine, Bun/Creatinine Ratio, Electrolytes, Uric Acid, Urine R/E	BUN, Creatinine, BUN/Creatinine Ratio, Electrolytes, Uric Acid, Urine R/E	BUN, Creatinine, BUN/Creatinine Ratio, Electrolytes, Uric Acid, Urine R/E	
BONES	Vitamin D, Calcium	Vitamin D, Calcium, Phosphorus	Vitamin D, Calcium, Phosphorus, Rheumatoid Factor	
THYROID	T3, T4, TSH	T3, T4, TSH	FT3, FT4, TSH	
NERVES	Vitamin B12	Vitamin B12	Vitamin B12	
LIVER	Bilirubin (Total, Direct, Indirect), SGOT, SGPT, ALP, Protein, Albumin, Globulin, A:G Ratio, HBsAg	Bilirubin (Total, Direct, Indirect), SGOT, SGPT, ALP, GGT, LDH, Protein, Albumin, Globulin, A:G Ratio, HBsAg	Bilirubin (Total, Direct, Indirect), SGOT, SGPT, ALP, GGT, LDH, Protein, Albumin, Globulin, A:G Ratio, HBsAg	
ANAEMIA	Iron, TIBC, UIBC, % Saturation	Iron, TIBC, UIBC, % Saturation, Ferritin	Iron, TIBC, UIBC, % Saturation, Ferritin, Folic Acid	
INFECTION	CBC, ESR	CBC, ESR	CBC, ESR	

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