302 Konark A Plus Sopan Baug

Pune

REPORT

(EDTA Mhala Blood)

Tel No: 919860155660

PID: 182043

Reference: Dr.--

SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

27-02-2021 05:27 PM

Biological Reference Interval

	Age:40.30 Years Sex:MALE	
Complete Blo	ood Count	

(EDTA Whole blood)		
Hemoglobin (Hb), EDTA whole blood	14.90	14.0 - 17.50 g/dL

Result

Method: Photometry

Total Leucocytes (WBC) count 5,700 4000-10000/µL

Method: Coulter Principle / Microscopy

344,000 Platelet count 150000 - 450000 /µL

Method: Coulter Principle / Microscopy

Red blood cell (RBC) count 5.09 4.52 - 5.90 x 10⁶ /µL

Method: Coulter Principle

PCV (Packed Cell Volume) 44.00 41.5 - 50.4 %

Method: Calculated

MCV (Mean Corpuscular Volume) 86.50 80.0 - 96.0 fL

Method: Derived from RBC histogram

MCH (Mean Corpuscular Hb) 29.20 27.5 - 33.2 pgms

Method: Calculated

33.4 - 35.5 g/dL MCHC (Mean Corpuscular Hb Conc.) 33.80

Method: Calculated

11.6 - 14.6 % RDW (RBC distribution width) 13.00

Method: Derived from RBC Histogram

WBC Differential Count

Method: VCSn / Microscopy / Calculated

Neutrophils	49	40 - 80 %
Absolute Neutrophils	2,793	2000 - 7000 /μL

Eosinophils	4	1 - 6 %	
Absolute Fosinophils	228	20 - 500 /ul	

Basophils	0	0 - 2 %
Absolute Basophils	0	0 - 100 /uL

Lymphocytes	38	20 - 40 %

, , ,		
Absolute Lymphocytes	2,166	1000 - 3000 /µL

Monocytes	9	2 - 10 %

Absolute Monocytes 513 ##\$



200 - 1000 /µL

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Pune

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Age:40.30 Years Sex: MALE

Reference:Dr.--SID: 120501585

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Complete Blood Count Findings

R.B.C. Normocytic, Normochromic

W.B.C. No abnormality detected

Platelets Adequate

ON FOLLOW UP. Remark

REPORT



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Page 2 of 15 Pr.(Mrs.) Awanti Golwilkar Mehendale MBBS,MD(Path) Regn.No:2000/02/1052

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302 Konark A Plus Sopan Baug

Pune

Tel No: 919860155660

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Reference: Dr.--

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

SID: 120501585

Age:40.30 Years Sex:MALE 27-02-2021 05:27 PM

Test Description Observed Value Biological Reference Interval

<u>Lipid Profile Maxi :</u>

REPORT

Serum Appearance Clear

Cholesterol (Total), serum by Enzymatic method 274 Desirable: < 200 mg/dL

Borderline high: 200 - 239 mg/dL

High : >/= 240 mg/dL

Triglycerides, serum by Enzymatic method 144 Normal : < 150 mg/dL

Borderline high: 150-199 mg/dL

High: 200-499 mg/dL Very high: >/= 500 mg/dL

HDL Cholesterol, serum by Enzymatic method 43 Men: > 40 mg/dL

Women: > 50 mg/dL

VLDL Cholestrol, serum by calculation 29 < 30 mg/dL

LDL Cholesterol, serum by calculation 202 Optimal: <100 mg/dL

Near optimal/above optimal: 100-129

mg/dL

Borderline high: 130-159 mg/dL

High: 160-189 mg/dL Very high: >/= 190 mg/dL

Cholesterol(Total)/HDL Cholesterol Ratio <u>6.37</u> Males : Acceptable ratio </= 5.00

Females : Acceptable ratio </= 4.50

LDL Cholesterol/HDL Cholesterol Ratio 4.70 Males: Acceptable ratio <= 3.60

Females : Acceptable ratio </= 3.20

Apolipoprotein A1, serum by Nephelometry 130 Male: 110 to 205 mg/dL

Apolipoprotein B, serum by Nephelometry <u>150</u> 55 to 140 mg/dL

Reference: ATP III, NCEP Guidelines and National Lipid Association (NLA) 2014 Recommendations

As per most international and national guidelines including Lipid Association of India 2016:

- 1. Lipoprotein and lipid levels should be considered in conjunction with other atherosclerotic cardiovascular disease (ASCVD) risk determinants to assess treatment goals and strategies.
- 2. Non-fasting lipid levels can be used in screening and in general risk estimation.



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REPORT

HIREN PARMAR 302 Konark A Plus Sopan Baug Pune

Tel No: 919860155660

PID: 182043

Reference:Dr.--

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am

SID: 120501585

Report Date: 27-02-2021 05:27 PM

Age:40.30 Years Sex: MALE

Test Description Liver Function Test :	Observed	Biological Reference Interval
Bilirubin-Total, serum by Diazo method	1.10	0.10 - 1.20 mg/dL Neonates : Upto 15.0 mg/dL
Bilirubin-Conjugated, serum by Diazo method	0.30	Upto 0.5 mg/dL
Bilirubin-Unconjugated, serum by calculation	0.80	0.1 to 1.0 mg/dL
SGOT (AST), serum by Enzymatic method	25	>or= 14 years : 8 - 48 U/Lt
SGPT (ALT), serum by Enzymatic Method	37	7 to 55 U/Lt
Alkaline Phosphatase, serum by pNPP-kinetic	70	Adult Male: (Unit: U/Lt.) 15 - < 17 years: 82 - 331 17 - < 19 years: 55 - 149 > or = 19 years: 40 - 129
Protein (total), serum by Biuret method	6.90	6.4 to 8.2 g/dL
Albumin, serum by Bromocresol purple method	4.30	3.4 to 5.0 g/dL
Globulin, serum by calculation	2.60	2.3 - 3.5 g/dL



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Page 4 of 15 Mc-3143 Dr.(Mrs.) Awanti Golwilkar Mehendale MBBS,MD(Path) Regn.No:2000/02/1052
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Reference: Dr.--

SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date: 27-02-2021 05:27 PM

Age:40.30 Years Sex:MALE

Test Description TEST NAME

REPORT

Observed Value Biological Reference Interval

Glycated Hemoglobin (HbA1C), by HPLC 5.60 4.0 to 5.6 %

Interpretation:

HbA1C level reflects the mean glucose concentration over previous 8-12 weeks and provides better indication of long term glycemic control.

For diagnosis of Diabetes Mellitus (>/= 18 yrs of age) :

5.7 % - 6.4 % : Increased risk for developing diabetes.

>/= 6.5 % : Diabetes

Therapeutic goals for glycemic control:

Adults: < 7%

Toddlers and Preschoolers: < 8.5% (but > 7.5%)

School age (6-12 yrs): < 8%

Adolescents and young adults (13 - 19 yrs): < 7.5 %

Levels of HbA1C may be low as result of shortened RBC life span in case of hemolytic anemia. Increased HbA1C values may be found in patients with polycythemia or post splenectomy patients. Patients with Homozygous forms of rare variant Hb(CC,SS,EE,SC) HbA1c can not be quantitated as there is no HbA. In such circumstances glycemic control can be monitored using plasma glucose levels or serum Fructosamine.

The A1c target should be individualized based on numerous factors, such as age, life expectancy, comorbid conditions, duration of diabetes, risk of hypoglycemia or adverse consequences from hypoglycemia, patient motivation and adherence.

Ref: ADA (Standards of Medical Care in Diabetes - 2017)



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Dr. Awanti Golwilkar

Dr. Vinanti Golwilkar

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Pune

Tel No: 919860155660

PID: 182043

Age:40.30 Years Sex: MALE

Reference: Dr.--SID: 120501585

> Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

27-02-2021 05:27 PM

Test Description

REPORT

Observed Value Biological Reference Interval

Gamma Glutamyl Transferase (GGT)

Gamma GT(GGT), Serum by Carboxy substrate-kinetic Male: (Unit: U/Lt.)

13 - 17 years : < 43 >or= 18 years: 8 - 61

Interpretation

- * GGT is used to diagnose and monitor hepatobiliary diseases.
- * Increased GGT and Alkaline Phosphatase indicate hepatobiliary diseases.
- * Normal GGT activity and increased Alkaline Phosphatase is consistent with skeletal disease.
- * May be used a screening test for occult alcoholism.
- * Elevated GGT is seen in:
 - 1) Intra or post hepatic biliary obstruction (5 to 30 times normal)
 - 2) Infectious hepatitis (2 to 5 times normal)
 - 3) Alcoholism
 - 4) Sclerosing cholangitis
 - 5) Primary or secondary neoplasm
 - 6) Medications such as phenytoin and phenobarbitone

Reference: Mayo Medical Laboratories, 2018 Interpretive Handbook.

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Sopan Baug Pune

REPORT

Tel No: 919860155660

PID: 182043

Age:40.30 Years Sex: MALE

SID: 120501585 Reference: Dr.--

> Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

27-02-2021 05:27 PM

Test Description	Observed Value	Biological Reference Interval
Plasma Glucose :		
Plasma glucose fasting, by Hexokinase method	92	< 100 mg/dL 100 to 125 mg/dL: Impaired fasting

glucose tolerance / Prediabetes >/= 126 mg/dL : Suggestive of diabetes mellitus (On more than one occasion) American Diabetes Association

Guidelines 2020

Clinical Chemistry

Urea, serum by GLDH-urease	19	17 to 49 mg/dL
BUN-Blood Urea Nitrogen, serum by calculation	8.88	8 to 23 mg/dL
Creatinine, serum by Jaffe w/o deproteinization	0.83	0.6 to 1.2 mg/dL
Uric Acid, serum by Uricase method	<u>7.60</u>	Male: 3.50 to 7.20 mg/dL

^{*} Uric acid is useful for 1. Diagnosis and follow up of renal failure. 2. Monitoring patients receiving cytotoxic drugs and a variety of other disorders, including gout, leukemia, psoriasis, starvation and other wasting conditions . * Increased uric acid is seen in following conditions :

- 1. Increased purine synthesis 2. Inherited metabolic disorders 3. Excess dietary purine intake
- 4. Increased nucleic acid turnover 5. Malignancy, cytotoxic drugs 6. Decreased urinary excretion (due to CRF) 7. Increased renal reabsorption .
- * Uric acid is decreased in : 1. Hepatocellular disease with reduced purine synthesis
- 2. Defective renal reabsorption 3. Overtreatment of uricemia (allopurinol or cancer therpies like 6-mercaptopurine, etc).



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Pune

Tel No: 919860155660

PID: 182043

Age:40.30 Years Sex: MALE

Reference: Dr.--

SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date: 27-02-2021 05:27 PM

Test Description Observed Value Biological Reference Interval

Clinical Chemistry:

REPORT

Calcium, serum by OCPC method

8.80 Adult: 8.4 to 10.2 mg/dL

Method : Colorimetric (o-cresolpthalein substrate) .

1. Calcium is useful for diagnosis and monitoring of a wide range of disorders including diseases of bone, kidney, parathyroid gland, or gastrointestinal tract.

2. Calcium ions play an important role in blood clotting, bone mineralization, musculature contractility and CNS functioning. .

3. Hypocalcemia is due to the absence or impaired function of the parathyroid glands or impaired vitamin-D synthesis. Chronic renal failure is also frequently associated with hypocalcemia due to decreased vitamin-D synthesis as well as hyperphosphatemia and skeletal resistance to the action of parathyroid hormone (PTH). 4. Hypercalcemia is mainly due to primary hyperparathyroidism (pHPT), and bone metastasis of carcinoma of the breast, thyroid gland, or lung. Severe hypercalcemia may result in cardiac arrhythmia.



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REPORT

HIREN PARMAR 302 Konark A Plus Sopan Baug Pune

Tel No: 919860155660

PID: 182043

Age:40.30 Years Sex: MALE

Reference: Dr.--

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

SID: 120501585

27-02-2021 05:27 PM

Test Description	Observed Value	Biological Reference Interval
Clinical Chemistry:		

Hormones

Free T3, serum by CMIA	3.13	1.71 to 3.71 pg/mL
Free T4, serum by CMIA	1.20	0.71 to 1.85 ng/dL
TSH(Ultrasensitive), serum by CMIA	3.64	0.40 - 4.00 μIU/mL
Testosterone (Total), serum by CMIA	535.73	Male:

15 to 16 yrs : 100 - 1200 ng/dL 17 to 18 yrs: 300 - 1200 ng/dL > or = 19 yrs : 240 - 950 ng/dL

Tanner stages

I (prepubertal) : < 7 - 20

II:8-66 III: 26 - 800 IV: 85 - 1200

V (young adult): 300 - 950

Testosterone is a major androgenic hormone. Decreased levels-Males:Partial or complete hypogonadism due to primary/secondary or tertiary testicular failure. Females : Primary or secondary ovarian failure, post- oophorectomy. Increased levels- Pre-pubertal boys/girls: Precocious puberty Males: Testicular/adrenal tumors, androgen abuse. Females: Polycystic ovarian syndrome, congenital adrenal hyperplasia and ovarian/adrenal tumors.

Note: Early-morning testosterone levels in young males are around 50% higher than p.m. levels. Testosterone levels can fluctuate substantially between different days, hence androgen status assessment should be based on more than a single measurement.



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Pune

Tel No: 919860155660

PID: 182043

Age:40.30 Years Sex: MALE

Reference: Dr.-- SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

27-02-2021 05:27 PM

Test Description Hormones:

REPORT

Observed Value

Biological Reference Interval



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DIAGNOSTICS
BE SURE
BE WELL

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Dr. Vinanti Golwilkar

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PID: 182043

Reference:Dr.--

SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date: 27-02-2021 05:27 PM

Age:40.30 Years Sex:MALE 27

Test Description Observed Value Biological Reference Interval

TEST NAME

REPORT

Vitamin B12, serum by CMIA <148 187 - 883 pg/mL

Interpretation:

Carrying forward

Four Decades

Dr. Ajit Golwilkar's legacy of Over

- 1. Vitamin B12 (cobalamin) is necessary for hematopoiesis and normal neuronal function.
- 2. Vitamin B12 is decreased in

Decreased Serum B12	
Pregnancy	
Contraceptive hormones	
Malabsorption	
Ethanol ingestion	
Smoking	
Strict vegan diet	
Pernicious anemia	

- 3. Serum methylmalonic acid and homocysteine levels are also elevated in vitamin B12 deficiency states. Active B12 (Holotranscobalamin) is low in Vitamin B12 deficiency.
- 4. Please correlate in case of patients taking vitamin B12 supplementation.



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Pune

REPORT

Tel No: 919860155660

PID: 182043

Age:40.30 Years Sex: MALE

Reference: Dr.--

SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

27-02-2021 05:27 PM

Test Description Observed value **Biological Reference Interval**

HOMA Index Insulin Resistance Test

Plasma glucose fasting, by Hexokinase method 92 < 100 mg/dL

> 100 to 125 mg/dL: Impaired fasting glucose tolerance / Prediabetes >/= 126 mg/dL : Suggestive of

diabetes mellitus

(On more than one occasion) American Diabetes Association

Guidelines 2020

Insulin Fasting, Serum by CMIA 4.20 Fasting: 2.5 to 25 µU/mL

Peak upto 150 µU/mL

HOMA IR Index 0.95 > 2.5 indicates insulin resistance

Interpretation

- 1. As, the direct measurement of the insulin effect on the blood sugar concentration is not possible other indices are used for determining an insulin resistance.
- 2. One of the most common indices is the HOMA index (Homeostasis Model Assessment), which is calculated according to the following formula:

HOMA index = fasting insulin (µU/ml) X fasting blood sugar (mg/dl) /405

- 3. Indications:
 - * Adiposis (BMI > 28 kg/m²)
 - * Suspected insulin resistance (metabolic syndrome, diabetes mellitus type 2)
 - * Suspected polycystic ovary syndrome (PCO-S)
 - * Cycle disturbances (e. g. amenorrhea)
 - * Infertility
- 4. Reference ranges:
 - > 2.0 indication for insulin resistance
 - > 2.5 insulin resistance probable
 - > 5.0 average value in patients with diabetes mellitus type 2

Reference: https://www.bioscientia.de/en/files/2011/10/Marker



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27-02-2021 05:27 PM

Age:40.30 Years Sex:MALE

Test Description TEST NAME

REPORT

25 - OH Vitamin D, serum by CMIA

Observed Value

13.30

Biological Reference Interval

Severe deficiency : < 10 ng/mL

Mild to moderate deficiency: 10 to 19 ng/mL

Optimum levels: 20 to 50 ng/mL

Increased risk of hypercalciuria: 51 to 80

ng/mL

Toxicity possible: > 80 ng/mL Ref.: Mayo Medical Laboratories These reference ranges represent clinical decision values, based on the 2011 Institute of Medicine report

Interpretation:

Vitamin D is vital for strong bones. It also has important, emerging roles in immune function and cancer prevention.

Vitamin D compounds in the body are exogenously derived by dietary means; from plants as 25-hydroxyvitamin D2 (ergocalciferol or calciferol) or from animal products as 25-hydroxyvitamin D3 (cholecalciferol or calcidiol).

Vitamin D may also be endogenously derived by conversion of 7-dihydrocholesterol to 25-hydroxyvitamin D3 in the skin upon ultraviolet exposure.

The total 25-hydroxyvitamin D (25-OH-VitD) level (the sum of 25-OH-vitamin D2 and 25-OH-vitamin D3) is the appropriate indicator of vitamin D body stores.

Patients with renal failure can have very high 25-OH-VitD levels without any signs of toxicity, as renal conversion to the active hormone 1,25-OH-VitD is impaired or absent.

Kindly corelate clinically, with supplementation history & repeat with fresh sample if necessary.



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SID: 120501585

Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date: 27-02-2021 05:27 PM

Age:40.30 Years Sex:MALE **Urine Routine Examination** Result **Biological Reference Interval**

(Sample: Urine, Automated / Semiautomated)

Physical

REPORT

Quantity Examined 5.0 ml

Method: Visual

Clear **Appearance**

Method: Visual / Automated

Colour Pale yellow

Method: Visual / Automated

Chemical (Dipstick)

pН 6.0 4.6 - 8.0

Method: Indicator Principle

Absent Protein Absent

Method: Sulphosalycylic Acid/ pH Indicator

Glucose **Absent** Absent

Method: GOD-POD/Benedict's

Acetone **Absent** Absent

Method: Sodium Nitroprusside reaction

Absent Bile Pigments Absent

Method: Diazo Reaction / Fouchet's test

Urobilinogen Not significant Not Significant

Method: Modified Ehrlich / Watson Schwartz

Microscopy / Flow cytometry

R.B.Cs 1-2 0 - 2 per hpf

Pus cells 1-2 0 - 5 per hpf

Epithelial cells Occasional 0 - 5 per hpf

Casts **Not Detected**

Not Detected Crystals

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SID: 120501585 Reference: Dr.--

> Collection Date: 27-02-2021 09:35 AM Registration Date: 27-02-2021 11:21 am Report Date:

27-02-2021 05:27 PM

Age:40.30 Years Sex: MALE **Test Description**

CRP(hs) - C- Reactive Protein high sensitivity

Observed Value Biological Reference Interval

See clinical information below

Method: Nephelometry / Immunoturbidimetry

Clinical Information:

REPORT

1. C-reactive protein (CRP) is a biomarker of inflammation. Plasma CRP concentrations increase rapidly and dramatically (100-fold or more) in response to tissue injury or inflammation.

3.84

2. High-sensitivity CRP (hs-CRP) is more precise than standard CRP when measuring baseline (i.e. normal) concentrations and enables a measure of chronic inflammation. It is recommended for cardiovascular risk assessment. Atherosclerosis is an inflammatory disease and hs-CRP has been endorsed by multiple guidelines as a biomarker of atherosclerotic cardiovascular disease risk.

Low cardiovascular risk : < 2.0 mg/L High cardiovascular risk : >/= 2.0 mg/L Acute inflammation : > 10.0 mg/L

3. A single test for high-sensitivity CRP (hs-CRP) may not reflect an individual patient's basal hs-CRP level. Repeat measurement may be required to firmly establish an individual's basal hs-CRP concentration. The lowest of the measurements should be used as the predictive value.

Reference: Mayo Medical Laboratories

End of Report

Dr.(Mrs.) Awanti Golwilkar Mehendale MBBS,MD(Path) Regn.No:2000/02/1052 A.G Diagnostics Pvt. Ltd.

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