





Dr. Vinay Chopra MD (Pathology & Microbiology) Chairman & Consultant Pathologist

Dr. Yugam Chopra MD (Pathology) CEO & Consultant Pathologist

**NAME** : Mrs. ARCHANA

**AGE/ GENDER** : 30 YRS/FEMALE **PATIENT ID** : 1701577

**COLLECTED BY** : SURJESH REG. NO./LAB NO. :012412170043

REFERRED BY : LOOMBA HOSPITAL (AMBALA CANTT) **REGISTRATION DATE** : 17/Dec/2024 02:15 PM BARCODE NO. :01522590 **COLLECTION DATE** : 17/Dec/2024 02:16PM CLIENT CODE. : KOS DIAGNOSTIC LAB REPORTING DATE : 17/Dec/2024 02:46PM

**CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT

**Value** Unit **Biological Reference interval Test Name** 

#### **HAEMATOLOGY HAEMOGLOBIN (HB)**

12.6 HAEMOGLOBIN (HB) 12.0 - 16.0gm/dL

by CALORIMETRIC

**INTERPRETATION:-**Hemoglobin is the protein molecule in red blood cells that carries oxygen from the lungs to the bodys tissues and returns carbon dioxide from the tissues back to the lungs.

A low hemoglobin level is referred to as ANEMIA or low red blood count.

#### ANEMIA (DECRESED HAEMOGLOBIN):

- 1) Loss of blood (traumatic injury, surgery, bleeding, colon cancer or stomach ulcer)
- 2) Nutritional deficiency (iron, vitamin B12, folate)
- 3) Bone marrow problems (replacement of bone marrow by cancer)
- 4) Suppression by red blood cell synthesis by chemotherapy drugs
- 5) Kidney failure
- 6) Abnormal hemoglobin structure (sickle cell anemia or thalassemia).

#### POLYCYTHEMIA (INCREASED HAEMOGLOBIN):

- 1) People in higher altitudes (Physiological)
- 2) Smoking (Secondary Polycythemia)
- 3) Dehydration produces a falsely rise in hemoglobin due to increased haemoconcentration
- 4) Advanced lung disease (for example, emphysema)
- 5) Certain tumors
- 6) A disorder of the bone marrow known as polycythemia rubra vera,
- 7) Abuse of the drug erythropoetin (Epogen) by athletes for blood doping purposes (increasing the amount of oxygen available to the body by chemically raising the production of red blood cells).

NOTE: TEST CONDUCTED ON EDTA WHOLE BLOOD



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: 17/Dec/2024 02:50PM

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#### **BLOOD GROUP (ABO) AND RH FACTOR TYPING**

REPORTING DATE

ABO GROUP
by SLIDE AGGLUTINATION
RH FACTOR TYPE
by SLIDE AGGLUTINATION

CLIENT CODE.

O

**POSITIVE** 



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**BLEEDING TIME (BT)** 

BLEEDING TIME (BT) 2 mts 35 secs MINS 1 - by DUKE METHOD



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**CLOTTING TIME (CT)** 

CLOTTING TIME (CT) 5 mts 50 secs MINS 4 - by CAPILLARY TUBE METHOD



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CLIENT CODE.



### **KOS Diagnostic Lab**

(A Unit of KOS Healthcare)



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: 17/Dec/2024 05:04PM

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### **CLINICAL CHEMISTRY/BIOCHEMISTRY GLUCOSE RANDOM (R)**

GLUCOSE RANDOM (R): PLASMA 132.15 NORMAL: < 140.00 mg/dL

by GLUCOSE OXIDASE - PEROXIDASE (GOD-POD) PREDIABETIC: 140.0 - 200.0 DIABETIC: > 0R = 200.0

INTERPRETATION

IN ACCORDANCE WITH AMERICAN DIABETES ASSOCIATION GUIDELINES:

1. A random plasma glucose level below 140 mg/dl is considered normal.

2. A random glucose level between 140 - 200 mg/dl is considered as glucose intolerant or prediabetic. A fasting and post-prnadial blood test (after consumption of 75 gms of glucose) is recommended for all such patients.

3. A random glucose level of above 200 mg/dl is highly suggestive of diabetic state. A repeat post-prandial is strongly recommended for all such patients. A fasting plasma glucose level in excess of 125 mg/dl on both occasions is confirmatory for diabetic state.



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0.35 - 5.50

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# ENDOCRINOLOGY THYROID STIMULATING HORMONE (TSH)

THYROID STIMULATING HORMONE (TSH): SERUM 0.424 µIU/mL

by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

3rd GENERATION, ULTRASENSITIVE

#### **INTERPRETATION:**

| AGE                 | REFFERENCE RANGE (μIU/mL) |  |  |
|---------------------|---------------------------|--|--|
| 0 – 5 DAYS          | 0.70 - 15.20              |  |  |
| 6 Days – 2 Months   | 0.70 - 11.00              |  |  |
| 3 – 11 Months       | 0.70 - 8.40               |  |  |
| 1 – 5 Years         | 0.70 - 7.00               |  |  |
| 6 – 10 Years        | 0.60 - 5.50               |  |  |
| 11 - 15             | 0.50 - 5.50               |  |  |
| > 20 Years (Adults) | 0.27 - 5.50               |  |  |
|                     | PREGNANCY                 |  |  |
| 1st Trimester       | 0.10 - 3.00               |  |  |
| 2nd Trimester       | 0.20 - 3.00               |  |  |
| 3rd Trimester       | 0.30 - 4.10               |  |  |

NOTE:-TSH levels are subjected to circardian variation, reaching peak levels between 2-4 a.m and at a minimum between 6-10 pm. The variation is of the order of 50 %. Hence time of the day has influence on the measured serum TSH concentration.

**USE**:- TSH controls biosynthesis and release of thyroid harmones T4 & T3. It is a sensitive measure of thyroid function, especially useful in early or subclinical hypothyroidism, before the patient develops any clinical findings or goitre or any other thyroid function abnormality.

#### **INCREASED LEVELS:**

- 1. Primary or untreated hypothyroidism, may vary from 3 times to more than 100 times normal depending on degree of hypofunction.
- $2. \\ Hypothyroid\ patients\ receiving\ insufficient\ thyroid\ replacement\ the rapy.$
- 3. Hashimotos thyroiditis.
- 4.DRUGS: Amphetamines, Iodine containing agents and dopamine antagonist.
- 5. Neonatal period, increase in 1st 2-3 days of life due to post-natal surge.

#### **DECREASED LEVELS:**

- 1. Toxic multi-nodular goitre & Thyroiditis.
- 2. Over replacement of thyroid harmone in treatment of hypothyroidism.
- 3. Autonomously functioning Thyroid adenoma
- 4. Secondary pituatary or hypothalmic hypothyroidism
- 5. Acute psychiatric illness
- 6. Severe dehydration.
- 7.DRUGS: Glucocorticoids, Dopamine, Levodopa, T4 replacement therapy, Anti-thyroid drugs for thyrotoxicosis.



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8.Pregnancy: 1st and 2nd Trimester

LIMITATIONS:

 $1. TSH\ may\ be\ normal\ in\ central\ hypothyroidism,\ recent\ rapid\ correction\ of\ hyperthyroidism\ or\ hypothyroidism,\ pregnancy,\ phenytoin\ the rapy.$ 

2. Autoimmune disorders may produce spurious results.



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#### ANTI MULLERIAN HORMONE (AMH) GEN II

ANTI MULLERIAN HORMONE (AMH) GEN II: SERUM 1.33

ng/mL 0.05 - 11.00

: 17/Dec/2024 09:15PM

by ECLIA (ELECTROCHEMILUMINESCENCE IMMUNOASSAY)

**INTERPRETATION:-**

CLIENT CODE.

#### A Correlation of FERTILITY POTENTIAL and AMH levels are:

| OVARIAN FERTILITY POTENTIAL | AMH VALUES IN (ng/mL)                   |  |
|-----------------------------|---|--|
| OPTIMAL FERTILITY:          | 4.00 – 6.80 ng/mL                       |  |
| SATISFACTORY FERTILITY:     | 2.20 – 4.00 ng/mL                       |  |
| LOW FERTILITY:              | 0.30 – 2.20 ng/mL                       |  |
| VERY LOW/UNDETECTABLE:      | 0.00 – 0.30 ng/mL                       |  |
| HIGH LEVEL:                 | >6.8 ng/mL (PCOD/GRANULOSA CELL TUMOUR) |  |

Anti Mullerian Hormone (AMH) is also known as Mullerian Inhibiting Substance provided by sertoli cells of the testis in males and by ovarian granulose cells in females upto antral stage in females.

1.It is used to evaluate testicular presence and function in infants with intersex conditions or ambiguous genitalia, and to distinguish between cryptorchidism and anorchia in males

#### IN FEMALES:

- 1.During reproductive age, follicular AMH productionbegins during the primary stage, peaks in preantral stage & has influence on follicular sensitivity to FSH which is impoetant in selection for follicular dominance. AMH levels thus represents the pool or number of primordial follicles but not thequality of oocytes. AMH does not vary significantly during menstrual cycle & hence can be measured independently of day of cycle.
- 2. Polycystic ovarian syndrome can elevate AMH 2 to 5 fold higher than age specific reference range & predict anovulatory, irregular cycles, ovarian tumours like Granulosa cell tumour are often associated with higher AMH levels.
- 3. Obese women are often associated with diminished ovarian reserve and can have 65% lower mean AMH levels than non-obese women.
- 4.In females, AMH levels do not change significantly throughout the menstrual cycle and decrease with age.
- 5. Assess Ovarian Reserve correlates with the number of antral follicies in the ovaries.
- 6.Evaluate fertility potential and ovarian response in IVF- Women with low AMG levels are more likely to the poor ovarian responders.
- 7. Assess the condition of Polycystic Ovary and premature ovarian failure.

A combination of Age, Ultrasound markers-Ovarian Volume and Antral Follicle Count, AMH and FSH levels are useful for optimal assessment of ovarian reserve. Studies in various fertility clinics are ongoing to establish optimal AMH concentretaion for predicting response to invitro fertilization, however, given below is suggested interpretative reference.

AMH levels (ng/mL) Suggested patient **Anticipated Antral** 



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| Test Name    |   | Value             | Unit  | Biological Reference interva |
|--------------|---|-------------------|---|------------------------------|
|              | Categorization for<br>fertility based on AMH<br>for age group (20 to 45<br>yrs) | Follicle counts   | (day 3)   | to IVF/COH cycle             |
| Below 0.3    | Very low  | Below 4           | Above 20  | Negligible/Poor              |
| 0.3 to 2.19  | Low   | 4 - 10            | Usually 16 - 20   | Reduced                      |
| 2.19 t0 4.00 | Satisfactory  | 11 - 25           | Within reference range<br>or between 11 - 15                | Safe/Normal                  |
| Above 4.00   | Optimal   | Upto 30 and Above | Within reference range<br>or between 11 – 15 or<br>Above 15 | Possibly Excessive           |

#### **INCREASED:**

- 1.Polycystic ovarian syndrome (most common)
- 2. Ovarian Tumour: Granulosa cell tumour

#### **DECREASED:**

- 1. Anorchia, Abnormal or absence of testis in males
- 2.Pseudohermaphroditism
- 3.Post Menopause

#### NOTE:

1.AMH measurement alone is seldom suffcient for diagnosis and results should be interpreted in the light of clinical finding and other relevant test such as ovarian ultrasonography(In fertility applications); abdominal or testicular ultrasound(intersex or testicular function applications); measurement of sex steroids (estradiol, Progesterone, Testosterone), FSH, Inhibin B (For fertility), and Inhibin A and B (for tumour work up).

2. Conversion of AMH grom ng/mL to pmol/L can be performed by using equation 1 ng/mL = 7.14 pmol/L



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Test Name Value Unit Biological Reference interval

# IMMUNOPATHOLOGY/SEROLOGY HEPATITIS C VIRUS (HCV) ANTIBODY: TOTAL

HEPATITIS C ANTIBODY (HCV) TOTAL: SERUM

0.06

S/CO

NEGATIVE: < 1.00

by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

POSITIVE: > 1.00

HEPATITIS C ANTIBODY (HCV) TOTAL

NON - REACTIVE

RESULT

by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

#### INTERPRETATION:-

| MATERIA REPORTORI |  |  |  |  |
|-------------------|--|--|--|--|
| RESULT (INDEX)    | REMARKS  |  |  |  |
| < 1.00            | NON - REACTIVE/NOT - DETECTED                        |  |  |  |
| >=1.00            | REACTIVE/ASYMPTOMATIC/INFECTIVE STATE/CARRIER STATE. |  |  |  |

Hepatitis C (HCV) is an RNA virus of Favivirus group transmitted via blood transfusions, transplantation, injection drug abusers, accidental needle punctures in healthcare workers, dialysis patients and rarely from mother to infant. 10 % of new cases show sexual transmission. As compared to HAV & HBV, chronic infection with HCV occurs in 85 % of infected individuals. In high risk population, the predictive value of Anti HCV for HCV infection is > 99% whereas in low risk populations it is only 25 %.

- 1. Indicator of past or present infection, but does not differentiate between Acute/ Chronic/Resolved Infection.
- 2. Routine screening of low and high prevelance population including blood donors.

#### NOTE:

- 1. False positive results are seen in Auto-immune disease, Rheumatoid Factor, HYpergammaglobulinemia, Paraproteinemia, Passive antibody transfer, Anti-idiotypes and Anti-superoxide dismutase.
- 2. False negative results are seen in early Acute infection, Immunosuppression and Immuno—incompetence. 3. HCV-RNA PCR recommended in all reactive results to differentiate between past and present infection.



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S/CO

: 17/Dec/2024 05:04PM

NEGATIVE: < 1.00

POSITIVE: > 1.00

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#### ANTI HUMAN IMMUNODEFICIENCY VIRUS (HIV) DUO ULTRA WITH (P-24 ANTIGEN DETECTION)

HIV 1/2 AND P24 ANTIGEN: SERUM

by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

HIV 1/2 AND P24 ANTIGEN RESULT

**NON - REACTIVE** 

by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

INTERPRETATION:-

CLIENT CODE.

| RESULT (INDEX) | REMARKS                |
|----------------|------------------------|
| < 1.00         | NON - REACTIVE         |
| > = 1.00       | PROVISIONALLY REACTIVE |

Non-Reactive result implies that antibodies to HIV 1/2 have not been detected in the sample. This menas that patient has either not been exposed to HIV 1/2 infection or the sample has been tested during the "window phase" i.e. before the development of detectable levels of antibodies. Hence a Non Reactive result does not exclude the possibility of exposure or infection with HIV 1/2. RECOMMENDATIONS:

1. Results to be clinically correlated

2. Rarely falsenegativity/positivity may occur.



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#### HEPATITIS B SURFACE ANTIGEN (HBsAg) ULTRA

HEPATITIS B SURFACE ANTIGEN (HBsAg):

0.18

NEGATIVE: < 1.0 POSITIVE: > 1.0

by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

NON REACTIVE

HEPATITIS B SURFACE ANTIGEN (HBsAg)

RESULT by CMIA (CHEMILUMINESCENT MICROPARTICLE IMMUNOASSAY)

#### INTERPRETATION:

| INVERTICAL.           |                |  |  |  |
|-----------------------|----------------|--|--|--|
| RESULT IN INDEX VALUE | REMARKS        |  |  |  |
| < 1.30                | NEGATIVE (-ve) |  |  |  |
| >=1.30                | POSITIVE (+ve) |  |  |  |

Hepatitis B Virus (HBV) is a member of the Hepadna virus family causing infection of the liver with extremely variable clinical features. Hepatitis B is transmitted primarily by body fluids especially serum and also spread effectively sexually and from mother to baby. In most individuals HBV hepatitis is self limiting, but 1-2 % normal adolescent and adults develop Chronic Hepatitis. Frequency of chronic HBV infection is 5-10% in immunocompromised patients and 80 % neonates. The initial serological marker of acute infection is HBsAg which typically appears 2-3 months after infection and disappears 12-20 weeks after onset of symtoms. Persistence of HBsAg for more than 6 months indicates carrier state or Chronic Liver disease.



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: 17/Dec/2024 03:08PM

**NAME** : Mrs. ARCHANA

**AGE/ GENDER** : 30 YRS/FEMALE **PATIENT ID** : 1701577

**COLLECTED BY** : SURJESH REG. NO./LAB NO. :012412170043

REFERRED BY : LOOMBA HOSPITAL (AMBALA CANTT) **REGISTRATION DATE** : 17/Dec/2024 02:15 PM BARCODE NO. :01522590 **COLLECTION DATE** : 17/Dec/2024 02:16PM

: KOS DIAGNOSTIC LAB **CLIENT ADDRESS** : 6349/1, NICHOLSON ROAD, AMBALA CANTT

**Test Name Value** Unit **Biological Reference interval** 

**VDRL** 

REPORTING DATE

**VDRL** NON REACTIVE NON REACTIVE

by IMMUNOCHROMATOGRAPHY

#### **INTERPRETATION:**

CLIENT CODE.

1. Does not become positive until 7 - 10 days after appearance of chancre.

- 2. High titer (>1:16) active disease.
- 3.Low titer (<1:8) biological falsepositive test in 90% cases or due to late or late latent syphillis.
- 4.Treatment of primary syphillis causes progressive decline tonegative VDRL within 2 years.
- 5. Rising titer (4X) indicates relapse, reinfection, or treatment failure and need for retreatment.
- 6. May benonreactive in early primary, late latent, and late syphillis (approx. 25% ofcases).
- 7. Reactive and weakly reactive tests should always be confirmed with FTA-ABS (fluorescent treponemal antibody absorption test).

#### SHORTTERM FALSE POSITIVE TEST RESULTS (<6 MONTHS DURATION) MAY OCCURIN:

- 1. Acute viral illnesses (e.g., hepatitis, measles, infectious mononucleosis)
- 2.M. pneumoniae; Chlamydia; Malaria infection.
- 3. Some immunizations
- 4. Pregnancy (rare)

#### LONGTERM FALSE POSITIVE TEST RESULTS (>6 MONTHS DURATION) MAY OCCUR IN:

- 1. Serious underlying disease e.g., collagen vascular diseases, leprosy, malignancy.
- 2.Intravenous drug users.
- 3. Rheumatoid arthritis, thyroiditis, AIDS, Sjogren's syndrome.
- 4.<10 % of patients older thanage 70 years.
- 5. Patients taking some anti-hypertensive drugs.

\*\*\* End Of Report \*



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