

Phone: 0522-4062223, 9305548277, 8400888844 9415577933, 9336154100, Tollfree No.: 8688360360

E-mail: charak1984@gmail.com

CMO Reg. No. RMEE 2445133 NABLReg. No.MC-2491 Certificate No. MIS-2023-0218

: CHA250036865 Patient Name : Mr.AMAR TIWARI Visit No

Age/Gender : 35 Y/M Registration ON : 01/Mar/2025 11:16AM Lab No : 10134161 Sample Collected ON : 01/Mar/2025 11:42AM Referred By : Dr.KG1 Sample Received ON : 01/Mar/2025 11:52AM Refer Lab/Hosp : CHARAK NA Report Generated ON : 01/Mar/2025 02:11PM

. CRP (Quantitative),LIPASE,AMYLASE,CHEST PA,ECG,HBA1C (EDTA),HBSAg,HCV,HIV,CREATININE,UREA,NA+K+,LFT,CBC (WHOLE BLOOD) Doctor Advice

| Test Name | Result | Unit | Bio. Ref | . Range | Method |
|---------------------------------|--------|------|----------|-------------|--------|
| HBA1C | | | | | |
| Glycosylated Hemoglobin (HbA1c) | 5.0 | % | 4 - 5.7 | HPLC (EDTA) |) |

NOTE:-

P.R.

Glycosylated Hemoglobin Test (HbA1c) is performed in this laboratory by the Gold Standard Reference method, ie: HPLC Technology(High performance Liquid Chromatography D10) from Bio-Rad Laboratories. USA.

EXPECTED (RESULT) RANGE:

Bio system Degree of normal 4.0 - 5.7 % Normal Value (OR) Non Diabetic 5.8 - 6.4 % Pre Diabetic Stage > 6.5 % Diabetic (or) Diabetic stage 6.5 - 7.0 % Well Controlled Diabet 7.1 - 8.0 % **Unsatisfactory Control** > 8.0 % Poor Control and needs treatment

CHARAK



DR. NISHANT SHARMA DR. SHADAB **PATHOLOGIST PATHOLOGIST**

Dr. SYED SAIF AHMAD MD (MICROBIOLOGY)

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| T | | | | | |
|-----------------------|--------|------|-----------------|--------|--|
| Test Name | Result | Unit | Bio. Ref. Range | Method | |
| CRP-QUANTITATIVE | | | | | |
| CRP-OUANTITATIVE TEST | 2.6 | MG/L | 0.1 - 6 | | |

Method: Immunoturbidimetric

(Method: Immunoturbidimetric on photometry system)

SUMMARY: C - reactive protien (CRP) is the best known among the acute phase protiens, a group of protien whose concentration increases in blood as a response to inflammatory disorders.CRP is normally present in low concentration in blood of healthy individuals (< 1mg/L). It is elevated up to 500 mg/L in acute inflammatory processes associated with bacterial infections, post operative conditions tissue damage already after 6 hours reaching a peak at 48 hours.. The measurment of CRP represents a useful aboratory test for detection of acute infection as well as for monitoring inflammtory proceses also in acute rheumatic & gastrointestinal disease. In recent studies it has been shows that in apparrently healthy subjects there is a direct orrelation between CRP concentrations & the risk of developing oronary heart disease (CHD).

hsCRP cut off for risk assessment as per CDC/AHA

Level Risk <1.0 Low 1.0-3.0 Average >3.0 High

All reports to be clinically corelated

AMYLASE SERUM AMYLASE

Comments:

Amylase is produced in the Pancreas and most of the elevation in serum is due to increased rate of Amylase entry into the blood stream / decreased rate of clearance or both. Serum Amylase rises within 6 to 48 hours of onset of Acute pancreatitis in 80% of patients, but is not proportional to the severity of the disease. Activity usually returns to normal in 3-5 days in patients with milder edematous form of the disease. Values persisting longer than this period suggest continuing necrosis of pancreas or Pseudocyst formation. Approximately 20% of patients with Pancreatitis have normal or near normal activity. Hyperlipemic patients with Pancreatitis also show spuriously normal Amylase levels due to suppression of Amylase activity by triglyceride. Low Amylase levels are seen in Chronic Pancreatitis, Congestive Heart failure, 2nd & 3rd trimesters of pregnancy, Gastrointestinal cancer & bone fractures.

amylase amylase amylase

20.0-80.00

Enzymatic

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| Test Name | | Result | Unit Bio. Ref. Range | | Range | Method |
|-----------|--|--------|----------------------|---------|--------------|--------|
| LIPASE | | | | | | |
| LIPASE | | 25 | U/L | Upto 60 | colorimetric | |

COMMENTS:as, such as acute pancreatitis, chronic pancreatitis, and obstruction of the pancreatic duct. In acute pancreatitis serum lipase activity tends to become elevated & remains for about 7 - 10 days. Increased lipase activity rarely lasts longer than 14 days, and prolonged increases suggest a poor prognosis or the presence of a cyst. Serum lipase may also be elevated in patients with chronic pancreatitis, obstruction of the pancreatic duct and non pancreatic conditions including renal diseases, various abdominal diseases such as acute cholecystitis, intestinal obstruction or infarction, duodenal ulcer, and liver disease, as well as alcoholism & diabetic keto-acidosis & in patients who have undergone endoscopic r

Lipase measurements are used in the diagnosis and treatment of diseases of the pancre

etrograde cholangiopancreatography. Elevation of serum lipase activity in patients with mumps strongly suggests significant pancreatic as well as salivary gland involvement by the disease......





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Bio. Ref. Range Method **Test Name** Result Unit

HEPATITIS B SURFACE ANTIGEN (HBsAg)

Sample Type: SERUM

HEPATITIS B SURFACE ANTIGEN NON REACTIVE <1 - Non Reactive

CMIA

>1 - Reactive

Note: This is only a Screening test. Confirmation of the result (Non Reactive/Reactive) should be done by performing a PCR based test.

COMMENTS:

-HBsAg is the first serological marker after infection with Hepatitis B Virus appearing one to ten weeks after exposure and two to eight weeks before the onset of clinical symptoms. HBsAg persists during the acute phase and clears late in the convalescence phase. Failure to clear HBsAg within six months indicates a chronic HBsAg carrier state. HBsAg assays are used to identify the persons infected with HBV and to prevent transmission of the virus by blood and blood products as well as to monitor the status of infected individuals in combination with other hepatitis B serological markers.

-Borderline cases must be confirmed with confirmatory neutralizing assay

LIMITATIONS:

- -Results should be used in conjunction with patient history and other hepatitis markers for diagnosis of acute and chronic infections.
 -Specimens from patients who have received preparations of mouse monoclonal antibodies for diagnosis or therapy may contain human anti-mouse antibodies (HAMA) which may produce anomalous values when tested with assay kits that employs mouse monoclonal antibodies.
- -Heterophilic antibodies in human serum can react with reagent immunoglobulins, interfering with in vitro immunoassays. Patients routinely exposed to animals or animal serum products can be prone to this interference and anomalous results may be observed.

 -Cross reactivity for specimens from individual with medical conditions (Pregnancy, HIV etc) has been observed.
- -HBsAg mutations may result in a false negative result in some HBsAg assays
- -If HBsAg results are inconsistent with clinical evidence, additional testing is suggested to confirm the result

HIV-SEROLOGY

HIV

NON REACTIVE

< 1.0 : NON REACTIVE >1.0: REACTIVE

Done by: Vitros ECI (Sandwich Assay)

Note:-Elisa test is a screening method for HIV.It is known to give false Positive & Negative result.

Hence confirmation: "Western Blot" method is advised.

DR. NISHANT SHARMA DR. SHADAB **PATHOLOGIST**

PATHOLOGIST

Dr. SYED SAIF AHMAD MD (MICROBIOLOGY) Page 4 of 7



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Test Name Result Unit Bio. Ref. Range Method

Anti-Hepatitis C Virus Antibodies.

NON REACTIVE

< 1.0 : NON REACTIVE Sandwich Assay

> 1.0 : REACTIVE

Done by: Vitros ECI (Sandwich Assay)

Note: This is only a Screening test. Confirmation of the result (Non Reactive/Reactive) should be done by performing a PCR based test.





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| Test Name | Result | Unit | Bio. Ref. Range | Method |
|------------------------------|---------|---------|-----------------|----------------|
| CBC (COMPLETE BLOOD COUNT) | | | | |
| Hb | 15.0 | g/dl | 12 - 15 | Non Cyanide |
| R.B.C. COUNT | 4.30 | mil/cmm | 3.8 - 4.8 | Electrical |
| | | | | Impedence |
| PCV | 46.2 | % | 36 - 45 | Pulse hieght |
| | | | | detection |
| MCV | 107.7 | fL | 80 - 96 | calculated |
| MCH | 35.0 | pg | 27 - 33 | Calculated |
| MCHC | 32.5 | g/dL | 30 - 36 | Calculated |
| RDW | 13.6 | % | 11 - 15 | RBC histogram |
| | | | | derivation |
| RETIC | 1.0 % | % | 0.5 - 2.5 | Microscopy |
| TOTAL LEUCOCYTES COUNT | 8340 | /cmm | 4000 - 10000 | Flocytrometry |
| DIFFERENTIAL LEUCOCYTE COUNT | | | | |
| NEUTROPHIL | 77 | % | 40 - 75 | Flowcytrometry |
| LYMPHOCYTES | 15 | % | 25 - 45 | Flowcytrometry |
| EOSINOPHIL | 4 | % | 1 - 6 | Flowcytrometry |
| MONOCYTE | 4 | % | 2 - 10 | Flowcytrometry |
| BASOPHIL | 0 | % | 00 - 01 | Flowcytrometry |
| PLATELET COUNT | 304,000 | /cmm | 150000 - 450000 | Elect Imped |
| PLATELET COUNT (MANUAL) | 304000 | /cmm | 150000 - 450000 | Microscopy. |
| Absolute Neutrophils Count | 6,422 | /cmm | 2000 - 7000 | Calculated |
| Absolute Lymphocytes Count | 1,251 | /cmm | 1000-3000 | Calculated |
| Absolute Eosinophils Count | 334 | /cmm | 20-500 | Calculated |
| Absolute Monocytes Count | 334 | /cmm | 200-1000 | Calculated |
| Mentzer Index | 25 | | | |
| Peripheral Blood Picture | : | | | |

Red blood cells are macrocytic. Platelets are adequate. No immature cells or parasite seen.







PR.

292/05, Tulsidas Marg, Basement Chowk, Lucknow-226 003

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| nit | Bio. Ref. Range | Method |
|-----|-----------------|--------|

| Test Name | Result | Unit | Bio. Ref. Range | Method |
|--------------------------------|--------|-------|-----------------|-------------------|
| NA+K+ | | | | |
| SODIUM Serum | 138.0 | MEq/L | 135 - 155 | ISE Direct |
| POTASSIUM Serum | 4.4 | MEq/L | 3.5 - 5.5 | ISE Direct |
| BLOOD UREA | | | | |
| BLOOD UREA | 19.90 | mg/dl | 15 - 45 | Urease, UV, Serum |
| SERUM CREATININE | | | | |
| CREATININE | 0.60 | mg/dl | 0.50 - 1.40 | Alkaline picrate- |
| | | | | kinetic |
| LIVER FUNCTION TEST | | | | |
| TOTAL BILIRUBIN | 1.77 | mg/dl | 0.4 - 1.1 | Diazonium Ion |
| CONJUGATED (D. Bilirubin) | 0.30 | mg/dL | 0.00-0.30 | Diazotization |
| UNCONJUGATED (I.D. Bilirubin) | 1.47 | mg/dL | 0.1 - 1.0 | Calculated |
| ALK PHOS | 103.80 | U/L | 30 - 120 | PNPP, AMP Buffer |
| SGPT | 42.0 | U/L | 5 - 40 | UV without P5P |
| SGOT | 27.0 | U/L | 5 - 40 | UV without P5P |

*** End Of Report ***

CHARAK





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Patient Name

: Mr.AMAR TIWARI

Age/Gender

: 35 Y/M

Lab No

PR.

: 10134161

Referred By

: Dr.KG1

Refer Lab/Hosp

: CHARAK NA

Visit No

: CHA250036865

Registration ON

: 01/Mar/2025 11:16AM

Sample Collected ON

: 01/Mar/2025 11:16AM

Sample Received ON

Report Generated ON

: 01/Mar/2025 12:31PM

ECG-REPORT

RATE

82 bpm

* RHYTHM

: Normal

* P wave

Normal

* PR interval

Normal

* QRS

Axis

Normal

Duration

Normal

Configuration

Normal

* ST-T Changes

None

_

* QT interval

* QTc interval

: Sec.

* Other

ner :

OPINION:

ECG WITH IN NORMAL LIMITS

(FINDING TO BE CORRELATED CLINICALLY)

[DR. PANKAJ RASTOGI, MD, DM]



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Referred By : Dr.KG1 Sample Received ON

Refer Lab/Hosp : CHARAK NA Report Generated ON : 01/Mar/2025 12:47PM

SKIAGRAM CHEST PA VIEW

- Both lung fields are clear.
- Bilateral hilar shadows are normal.
- Cardiac shadow is within normal limits.
- Both CP angles are clear.
- Soft tissue and bony cage are seen normally.
- Both domes of diaphragm are sharply defined.

IMPRESSION:

• NO ACTIVE LUNG PARENCHYMAL LESION IS DISCERNIBLE.

Clinical correlation is necessary.

[DR. RAJESH KUMAR SHARMA, MD]

transcribed by: anup

*** End Of Report ***

