



Tests you can trust

Pco g : XXXXXXXXXXXXXXXXXXXX

F cvg : XXXXXXXXXX

VguV'Cungf : Jaanch Rheumatoid Arthritis Package

T gr qtv"Uvcwu : Complete Report



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NABL From 2005[#]



ISO 9001: 2015 - From 2015



CAP From 2011⁻



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98% Reports
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of sample reaching the lab^{*}



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that Thyrocare
reports are
Accurate & Reliable^{*}



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Unique Barcode
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Reports Verified By Expert
MD Pathologists Stationed
at Every Lab

Patient Name : XXXXXXXXXXXXX
 Referred By : XXXXXXXXXXXXX
 Address : XXXXXXXXXXXXX

Tests Done : JAANCH RHEUMATOID ARTHRITIS
 PACKAGE



Report Availability Summary

Note: Please refer to the table below for status of your tests.

 **11** Ready
  **0** Ready with Cancellation
  **0** Processing
  **0** Cancelled in Lab

TEST DETAILS

REPORT STATUS

JAANCH RHEUMATOID ARTHRITIS PACKAGE	Ready 
CALCIUM	Ready 
C-REACTIVE PROTEIN (CRP)	Ready 
25-OH VITAMIN D (TOTAL)	Ready 
ERYTHROCYTE SEDIMENTATION RATE (ESR)	Ready 
PHOSPHOROUS	Ready 
RHEUMATOID FACTOR (RF)	Ready 
URIC ACID	Ready 
HEMOGRAM - 6 PART (DIFF)	Ready 
ANTI CCP (ACCP)	Ready 
ALKALINE PHOSPHATASE	Ready 
ANTI NUCLEAR ANTIBODIES (ANA)	Ready 

Patient Name : XXXXXXXXXXXXX
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 Address : XXXXXXXXXXXXX

 Tests Done : JAANCH RHEUMATOID ARTHRITIS
 PACKAGE


Tests Outside Reference Range

Note: Please refer to the table below for tests outside reference range.

Test Name	Observed Value	Units	Bio. Ref. Interval.
ARTHRITIS			
RHEUMATOID FACTOR (RF)	18.66	IU/mL	<=18
COMPLETE HEMOGRAM			
HEMATOCRIT(PCV)	33.6	%	40.0-50.0
HEMOGLOBIN	10.4	g/dL	13.0-17.0
LYMPHOCYTE	17.4	%	20-40
MEAN CORP.HEMO.CONC(MCHC)	31	g/dL	31.5-34.5
MEAN CORPUSCULAR HEMOGLOBIN(MCH)	26.8	pq	27.0-32.0
RED CELL DISTRIBUTION WIDTH (RDW-CV)	15.4	%	11.6-14
RED CELL DISTRIBUTION WIDTH - SD(RDW-SD)	48.9	fL	39-46
TOTAL RBC	3.88	X 10 ⁶ /μL	4.5-5.5
VITAMIN			
25-OH VITAMIN D (TOTAL)	11.62	ng/mL	30-100



Patient Name : XXXXXXXXXXXXX
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Address : XXXXXXXXXXXXX

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

TEST NAME	METHODOLOGY	VALUE	UNITS	Bio. Ref. Interval.
HEMOGLOBIN	SLS-Hemoglobin Method	10.4	g/dL	13.0-17.0
Hematocrit (PCV)	CPH Detection	33.6	%	40.0-50.0
Total RBC	HF & EI	3.88	X 10⁶/μL	4.5-5.5
Mean Corpuscular Volume (MCV)	Calculated	86.6	fL	83.0-101.0
Mean Corpuscular Hemoglobin (MCH)	Calculated	26.8	pq	27.0-32.0
Mean Corp.Hemo. Conc (MCHC)	Calculated	31	g/dL	31.5-34.5
Red Cell Distribution Width - SD (RDW-SD)	Calculated	48.9	fL	39-46
Red Cell Distribution Width (RDW - CV)	Calculated	15.4	%	11.6-14
RED CELL DISTRIBUTION WIDTH INDEX (RDWI)	Calculated	343.7	-	*Refer Note below
MENTZER INDEX	Calculated	22.3	-	*Refer Note below
TOTAL LEUCOCYTE COUNT (WBC)	HF & FC	7.77	X 10³ / μL	4.0 - 10.0
DIFFERENTIAL LEUCOCYTE COUNT				
Neutrophils Percentage	Flow Cytometry	75.2	%	40-80
Lymphocytes Percentage	Flow Cytometry	17.4	%	20-40
Monocytes Percentage	Flow Cytometry	2.7	%	2-10
Eosinophils Percentage	Flow Cytometry	3.9	%	1-6
Basophils Percentage	Flow Cytometry	0.5	%	0-2
Immature Granulocyte Percentage (IG%)	Flow Cytometry	0.3	%	0-0.5
Nucleated Red Blood Cells %	Flow Cytometry	0.1	%	0.0-5.0
ABSOLUTE LEUCOCYTE COUNT				
Neutrophils - Absolute Count	Calculated	5.84	X 10 ³ / μL	2.0-7.0
Lymphocytes - Absolute Count	Calculated	1.35	X 10 ³ / μL	1.0-3.0
Monocytes - Absolute Count	Calculated	0.21	X 10 ³ / μL	0.2 - 1.0
Basophils - Absolute Count	Calculated	0.04	X 10 ³ / μL	0.02 - 0.1
Eosinophils - Absolute Count	Calculated	0.3	X 10 ³ / μL	0.02 - 0.5
Immature Granulocytes (IG)	Calculated	0.02	X 10 ³ / μL	0-0.3
Nucleated Red Blood Cells	Calculated	0.01	X 10 ³ / μL	0.0-0.5
PLATELET COUNT	HF & EI	260	X 10³ / μL	150-410
Mean Platelet Volume (MPV)	Calculated	10.3	fL	6.5-12
Platelet Distribution Width (PDW)	Calculated	11.2	fL	9.6-15.2
Platelet to Large Cell Ratio (PLCR)	Calculated	26.6	%	19.7-42.4
Plateletcrit (PCT)	Calculated	0.27	%	0.19-0.39

*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

(Reference : *FC- flowcytometry, *HF- hydrodynamic focussing, *EI- Electric Impedence, *Hb- hemoglobin, *CPH- Cumulative pulse height)

Tests Done : ESR,HEMOGRAM - 6 PART (DIFF)

Doctor 1 Sign

Doctor 2 Sign

Report Remarks : LabCode:2406003451/IT011



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Sample Collected on (SCT): XXXXXXXXXXXXX
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Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS
ERYTHROCYTE SEDIMENTATION RATE (ESR) Bio. Ref. Interval. :-	MODIFIED WESTERGREN	< 7.5	mm / hr

<50 yr : Male: 0-15 mm/hr Female: 0-20 mm/hr
>50 yr : Male: 0-20 mm/hr Female: 0-30 mm/hr
Children: <=10 mm/hr

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

Tests Done : ESR, HEMOGRAM - 6 PART (DIFF)

Report Remarks : LabCode:2406003451/IT011

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TEST NAME	TECHNOLOGY	VALUE	UNITS
25-OH VITAMIN D (TOTAL) Bio. Ref. Interval. :-	C.L.I.A	11.62	ng/mL

DEFICIENCY : <20 ng/ml || INSUFFICIENCY : 20-<30 ng/ml
SUFFICIENCY : 30-100 ng/ml || TOXICITY : >100 ng/ml

Clinical Significance:

Vitamin D is a fat soluble vitamin that has been known to help the body absorb and retain calcium and phosphorous; both are critical for building bone health. Decrease in vitamin D total levels indicate inadequate exposure of sunlight, dietary deficiency, nephrotic syndrome. Increase in vitamin D total levels indicate Vitamin D intoxication.

Specifications: Precision: Intra assay (%CV):5.3%, Inter assay (%CV):11.9% ; Sensitivity:3.2 ng/ml.

Kit Validation Reference: Holick MF. Vitamin D Deficiency. N Engl J Med. 2007;357:266-81.

Please correlate with clinical conditions.

Method:- Fully Automated Chemi Luminescent Immuno Assay

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

Report Remarks : LabCode:2406003452/IT011

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Report Released on (RRT) : XXXXXXXXXXXXX
Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
URIC ACID	PHOTOMETRY	6.8	mg/dL	4.2 - 7.3
CALCIUM	PHOTOMETRY	9.62	mg/dL	8.8-10.6

Please correlate with clinical conditions.

Method :

URIC - Uricase / Peroxidase Method
CALC - Arsenazo III Method, End Point.

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

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TEST NAME	TECHNOLOGY	VALUE	UNITS
RHEUMATOID FACTOR (RF)	IMMUNOTURBIDIMETRY	18.66	IU/mL

Bio. Ref. Interval. :
ADULT : <= 18

Clinical Significance:

Rheumatoid factor is an anti IgG autoimmune antibody. There are high concentration of rheumatoid factor in the serum of some disease, especially rheumatoid arthritis patients. It helps to diagnose rheumatism ,systematic lupus erythematosus, chronic hepatitis etc.

Specifications:

Precision %CV :- Intra assay %CV- 1.38% , Inter assay %CV-2.88%, Sensitivity :- 40 IU/mL.

Kit Validation Reference:

Anderson, S.G., Bentzon, M.W., Houba, V. and Krag, P. Bull. Wld. Hlth. Org. 42: 311-318 (1970).

Method : LATEX ENHANCED IMMUNOTURBIDIMETRY

Please correlate with clinical conditions.

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

Report Remarks : LabCode:2406003452/IT011

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Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS
PHOSPHOROUS	PHOTOMETRY	4.24	mg/dL

Bio. Ref. Interval. :

Adults : 2.4 - 5.1 mg/dL

Children : 4.0 - 7.0 mg/dL

Clinical Significance:

In plasma and serum the majority of phosphate exists in the inorganic form (Pi), approximately 15% bound to protein and the remainder in complexes and free forms. Serum phosphate concentrations are dependent on diet and variation in the secretion of hormones such as Parathyroid Hormone (PTH).

Specifications:

Precision %CV :- Intra assay %CV- 1.55% , Inter assay %CV-2.99% , Sensitivity:-0.10 mmol/L

Kit Validation Reference:

Young DS. Effects of drugs on clinical laboratory tests, 5th ed. AACC Press, 2000.

Method : UNREDUCED PHOSPHOMOLYBDATE METHOD

Please correlate with clinical conditions.

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

Report Remarks : LabCode:2406003452/IT011

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Sample Received on (SRT) : XXXXXXXXXXXXX
Report Released on (RRT) : XXXXXXXXXXXXX
Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS
C-REACTIVE PROTEIN (CRP) Bio. Ref. Interval. :-	IMMUNOTURBIDIMETRY	1.33	mg/L

Acute phase determination : < 5 mg/L

Clinical Significance:

It's a protein present in the sera of acutely ill patients that bound cell wall C-polysaccharide of streptococcus pneumoniae and agglutinates the organisms. CRP is one of the strongest acute -phase reactants, with plasma concentrations rising up after myocardial infarction, stress, trauma, infection, inflammation, surgery, or neoplastic proliferation.

Concentrations >5 mg/L suggest the presence of an infection or inflammatory process. Concentrations are generally higher in bacterial than viral infection. The increase in peak is proportional to tissue damage. Determination of CRP is clinically useful to screen activity of inflammatory diseases such as rheumatoid arthritis; SLE; Leukemia; after surgery; to detect rejection in renal allograft recipients; to detect neonatal septicemia and meningitis. However, it is a nonspecific marker and cannot be interpreted without other clinical information.

Reference:

Tietz Textbook of clinical chemistry and molecular diagnosis fifth edition chapter 21 P538-539

Please correlate with clinical conditions.

Method:- FULLY AUTOMATED LATEX AGGLUTINATION – BECKMAN COULTER

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

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Report Released on (RRT) : XXXXXXXXXXXXX
Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS
ANTI NUCLEAR ANTIBODIES (ANA)	E.L.I.S.A	0.88	OD Ratio

Bio. Ref. Interval. :

Negative: <0.90
Equivocal : 0.90 - 1.10
Positive: >1.11

Clinical Significance:

Anti-nuclear antibodies (ANA) are autoantibodies directed against nuclear components and are commonly associated with autoimmune diseases such as systemic lupus erythematosus (SLE), mixed connective tissue disease (MCTD), systemic sclerosis, Sjögren's syndrome, and rheumatoid arthritis.

However, ANA positivity is not specific to any one condition and may also be seen in healthy individuals, especially the elderly, or in cases of infections, malignancies, or drug-induced autoimmunity.

Equivocal results may require repeat testing or additional evaluation with specific antibody panels (e.g., anti-Sm, anti-RNP, anti-Ro, anti-La) for diagnostic clarification. A negative ANA test does not completely rule out autoimmune disease, particularly in early stages or when non-ANA-specific autoantibodies are involved. The final diagnosis should be made by the physician based on the overall clinical picture and appropriate correlation with additional investigations.

Note: Hemolysis, lipemia, icterus, microbial contamination, high-dose biotin, or heterophile antibodies may interfere with test accuracy and may lead to false-positive or false-negative results.

Method : INDIRECT SOLID PHASE IMMUNOASSAY

Please correlate with clinical conditions.

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

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Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS	Bio. Ref. Interval.
ALKALINE PHOSPHATASE	PHOTOMETRY	105.1	U/L	45-129

Please correlate with clinical conditions.

Method :

ALKP - Modified IFCC method

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

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Sample Type | Barcode : XXXXXXXXXXXXX

TEST NAME	TECHNOLOGY	VALUE	UNITS
ANTI CCP (ACCP)	E.L.I.S.A	3.33	AU/mL

Reference Range :-

BRI	Interpretation
Negative < 15	Absence of IgG autoantibodies to cyclic citrullinated peptides (CCP)
Borderline 15-25	Indeterminate results. Additional testing and clinical correlation recommended
Positive > 25	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. If results are inconsistent with clinical history additional testing is suggested to confirm the results.
8. 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
- Vossenar ER et al. Arthritis Rheum. 2004
- Masson Bessire C et al. J Immunol. 2001

Please correlate with clinical conditions.

Method:- INDIRECT SOLID PHASE ENZYME IMMUNOASSAY

Tests Done : JAANCH RHEUMATOID ARTHRITIS PACKAGE

Doctor 1 Sign

Doctor 2 Sign

Report Remarks : LabCode:2406003452/IT011

CONDITIONS OF REPORTING

- ✓ The reported results are for information and interpretation of the referring doctor only.
- ✓ It is presumed that the tests performed on the specimen belong to the patient; named or identified.
- ✓ Results of tests may vary from laboratory to laboratory and also in some parameters from time to time for the same patient.
- ✓ Should the results indicate an unexpected abnormality, the same should be reconfirmed.
- ✓ Only such medical professionals who understand reporting units, reference ranges and limitations of technologies should interpret results.
- ✓ This report is not valid for medico-legal purpose.
- ✓ 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.
- ✓ Thyrocare Discovery video link :- <https://youtu.be/nbdYeRgYyQc>

EXPLANATIONS

- ✓ Majority of the specimen processed in the laboratory are collected by Pathologists and Hospitals we call them as "Clients".
- ✓ **Name** - The name is as declared by the client and recored by the personnel who collected the specimen.
- ✓ **Ref.Dr** - The name of the doctor who has recommended testing as declared by the client.
- ✓ **Labcode** - This is the accession number in our laboratory and it helps us in archiving and retrieving the data.
- ✓ **Barcode** - This is the specimen identity number and it states that the results are for the specimen bearing the barcode (irrespective of the name).
- ✓ **SCP** - Specimen Collection Point - This is the location where the blood or specimen was collected as declared by the client.
- ✓ **SCT** - Specimen Collection Time - The time when specimen was collected as declared by the client.
- ✓ **SRT** - Specimen Receiving Time - This time when the specimen reached our laboratory.
- ✓ **RRT** - Report Releasing Time - The time when our pathologist has released the values for Reporting.
- ✓ **Reference Range** - Means the range of values in which 95% of the normal population would fall.

SUGGESTIONS

- ✓ Values out of reference range requires reconfirmation before starting any medical treatment.
- ✓ Retesting is needed if you suspect any quality shortcomings.
- ✓ Testing or retesting should be done in accredited laboratories.
- ✓ For suggestions, complaints, clinical support or feedback, write to us at customersupport@thyrocare.com or call us on **022-3090 0000**

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* T&C Apply, #As on 5th December 2024 (Applicable for all company owned labs except Bhagalpur & Vijayawada),

* As per survey on doctors' perception of laboratory diagnostics (IJARIIT, 2023), -Mumbai Reference Lab is CAP Accredited