



Laboratory Investigation Report

Patient Name	Centre
Age/Gender	OP/IP No/UHID
MaxID/Lab ID	Collection Date/Time
Ref Doctor	Reporting Date/Time
Passport No.	

Molecular Diagnostics



Test Name	Result	Unit	Bio Ref Interval
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COVID-19 RTPCR (SARS COV-2)THROAT/NASAL SWAB- Uttar Pradesh H/C*

COVID-19 (SARS CoV-2) Real Time PCR Negative

Comment Positive result does not necessarily indicate the presence of an active, viable virus as RTPCR only detects the presence of viral RNA (dead or alive).

In clinically suspected patients, a single negative test result does not exclude infection. Presence of inhibitors, mutations and insufficient RNA can influence the test results.

In case of clinical discrepancy with RTPCR test results, please feel free to contact us for further course of action.

Please correlate the test result with Clinical & Radiological findings.

CT Value Literature: -


1. There are no reliable studies to definitively prove a direct correlation between disease severity / infectiousness and CT values. Viral load does not have much role in patient management.
2. CT values differ from one kit to the other. Comparability of CT values among different kits is a challenge as different labs are using a mixed basket of kits with different CT cut-offs and different gene targets.
3. Samples from asymptomatic / mild cases show CT values similar to those who develop severe disease.
4. Patients in early symptomatic stage may show a high CT value which may subsequently change. In such cases, high Ct values will give a false sense of security.
5. Severity of COVID-19 disease largely depends on host factors besides the viral load. Some patients with low viral load may land up in very severe disease due to triggering of the immunological responses. Hence, again high CT value may give a false sense of security.
6. Negative result shows no CT value

ICMR Registration Number: MAXDL001

SRF No.:0708304413506

Kindly correlate with clinical findings

*** End Of Report ***


Dr. Bansidhar Tarai, M.D
 Associate Director
 Microbiology & Molecular Diagnostics


Dr. Sonu Kumari Agrawal, MD
 Associate Consultant
 Microbiology


Dr Nidhi Malik, MD
 Consultant Microbiology



Patient Ref. No. 9000012180693

CLIENT CODE : C000060706

CLIENT'S NAME AND ADDRESS :

CROSSLEY REMEDIES LIMITED
 A UNIT OF MAX HOSPITAL, W-3, SECTOR-1,
 VAISHALI,
 GHAZIABAD 201012
 UTTAR PRADESH INDIA
 9650111522

SRL Ltd
 PRIME SQUARE BUILDING,PLOT NO 1,GAIWADI INDUSTRIAL
 ESTATE,S.V. ROAD,GOREGAON (W)
 MUMBAI, 400062
 MAHARASHTRA, INDIA
 Tel : 9111591115, Fax : 022 - 67801212
 CIN - U74899PB1995PLC045956

Test Report Status	Final	Results	Biological Reference Interval	Units
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SEROLOGY

INFLUENZA VIRUS A IGG (SERUM, ELISA)

INFLUENZA VIRUS A IGG (SERUM, ELISA)	2.8	Negative: < 9.0 Indeterminate: 9.0 - 11.0 Positive: > 11.0	NTU
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Comments

The sample for Influenza A Virus-IgG(Serum,ELISA), was referred to an external laboratory (code no.20002) for performance of the test requested.



Scan to View Details



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Patient Ref. No. 9000012180693

CLIENT CODE : C000060706

CLIENT'S N.
CROSSLEY RI
A UNIT OF M,
VAISHALI,
GHAZIABAD
UTTAR PRADE
9650111522

SRL Ltd
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ESTATE,S.V.
MUMBAI, 40
MAHARASH
Tel : 9111591115, Fax : 022 - 6/801212
CIN - U74899PB1995PLC045956

STRIAL

Test Report Status	Final	Results	Biological Reference Interval	Units
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EIA - INFECTIOUS SECTION

INFLUENZA VIRUS B IGG (SERUM, ELISA)

INFLUENZA VIRUS B IGG (SERUM, ELISA)	2.1	Negative: < 9.0 Indeterminate: 9.0 - 11.0 Positive: > 11.0	NTU
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Comments

The sample for Influenza B Virus-IgG(Serum,ELISA), was referred to an external laboratory (code no.20002) for performance of the test requested.

****End Of Report****

Please visit www.srlworld.com for related Test Information for this accession

Dr. Ekta Patil,MD
Microbiologist



Scan to View Details



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CONDITIONS OF LABORATORY TESTING & REPORTING

- | | |
|---|---|
| <ol style="list-style-type: none"> 1. It is presumed that the test sample belongs to the patient named or identified in the test requisition form. 2. All tests are performed and reported as per the turnaround time stated in the SRL Directory of Services. 3. Result delays could occur due to unforeseen circumstances such as non-availability of kits / equipment breakdown / natural calamities / technical downtime or any other unforeseen event. 4. A requested test might not be performed if: <ol style="list-style-type: none"> i. Specimen received is insufficient or inappropriate ii. Specimen quality is unsatisfactory iii. Incorrect specimen type iv. Discrepancy between identification on specimen container label and test requisition form | <ol style="list-style-type: none"> 5. SRL confirms that all tests have been performed or assayed with highest quality standards, clinical safety & technical integrity. 6. Laboratory results should not be interpreted in isolation; it must be correlated with clinical information and be interpreted by registered medical practitioners only to determine final diagnosis. 7. Test results may vary based on time of collection, physiological condition of the patient, current medication or nutritional and dietary changes. Please consult your doctor or call us for any clarification. 8. Test results cannot be used for Medico legal purposes. 9. In case of queries please call customer care (91115 91115) within 48 hours of the report. |
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SRL Limited
Fortis Hospital, Sector 62, Phase VIII,
Mohali 160062

