



Name	Sujith Nivedh	Registered Date	12-Feb-2022 06:52 AM
Age/Gender	13 Years/Male	Collection Date	12-Feb-2022 08:20 AM
Registration No	376790	Reported	12-Feb-2022 09:18 AM
Barcode No	RD16128160	Panel	Walk-In
Referred By	SELF	Passport No	V1170437
SRF No	2999900319971	Adhaar No	
		Date of Birth	17-Apr-2009

RAPID PCR COVID-19

Final Result	Negative	
Name of Assay	SARS-CoV-2(COVID-19) RAPID-PCR Assay.	
Name of Technology	Abbott ID NOW.	
Specimen Type	Nasopharyngeal swab.	
ICMR Registration Number for COVID-19: AURPLLUBK		

Interpretation:

Positive result is considered a positive test result for nCoV-19(COVID-19). This shows that RNA from novel corona virus (SARS-CoV-2) was detected and patient should be considered infected with corona virus.

Negative result for nCoV-19(COVID-19) means that nCoV-19(COVID-19) RNA was not present in the specimen.

The result (negative or positive) of this test must always be correlated with clinical status and history of the patient and other relevant data and should not be used alone for the interpretation.

Positive results but do not rule out bacterial infection or co-infection with other viruses& Negative results do not preclude COVID- 19 and should not be used as the sole basis for patient management decisions.

If the virus mutates in RT-PCR target region, nCoV-19 may not be detected or may be detected less predictably. Inhibitors or other types of interference may produce a false negative result. Kindly correlate the test results with clinical findings.

The performance of this test has not been established for monitoring treatment of nCoV-19(SARS-CoV-19) Infection.

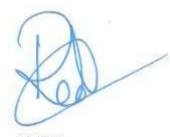
False positive results may happen from cross-contamination between patient samples, specimen mix-up and RNA contamination during product handling.

Possible cause of false negative results - Inadequate specimen quality. Specimen collected too early or too late. Specimens improperly handled or transported. Occurrence of viral genetic mutation. Presence of PCR inhibitors. Antiviral administration prior to testing. Note: Test has been performed using ICMR approved kit.

D Block
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*** End Of Report ***





Dr.Radhika Clinical Microbiologist

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