





CLIENT CODE: C000114814

CLIENT'S NAME AND ADDRESS :

SANTOSHI DIAGNOSTIC CENTRE SHOP NO. 1 MANGAL NIWAS SURVEY NO. 25 MUNDHWA, MANJARI ROAD SURVEY NO 25/3 KESHAV NAGAR LONKAR WASTI, OPP. SHREE LAXMI PLY & HARDWARE MUNDHWA, PUNE 411036 MAHARASHTRA INDIA 9322188588

SRL Ltd 301-305, 3RD FLOOR, KAMLA ARCADE, J M ROAD, 3RD FLOOR, KAMLA ARCADE, J M ROAD, OPP- BALGANDHARVA RANG MANDIR PUNE, 411004 MAHARASHTRA, INDIA Tel : 9111591115

Test Report Status <u>Fi</u>	nal Results	Biological Reference Interval Units
ICMR Registration No: SRLI AADHAAR NO 9726 1782		
CLINICAL INFORMATION :		
REFERRING DOCTOR : SEL	F	CLIENT PATIENT ID :
DRAWN : 25/11/2021 17:3	RECEIVED : 25/11/2021 22:21	REPORTED : 26/11/2021 10:57
ACCESSION NO : 0285UK	001486 AGE : 40 Years SEX : Male	
PATIENT NAME : VISHA	L VINAYAKRAO ASTUNKAR	PATIENT ID : VISHM301119800A

MOLECULAR BIOLOGY

SARS COV -2 REAL TIME PCR

SARS-COV-2 RNA

NEGATIVE

Interpretation(s) SARS COV -2 REAL TIME PCR-SARS-CoV-2, formerly known as 2019-nCoV, is the causative agent of the coronavirus disease 2019 (COVID-19). Main symptoms of the disease include fever, cough and shortness of breath. SARS-CoV-2 transmission occurs primarily via respiratory droplets. SARS-CoV-2 is likely to be at the highest concentrations in the nasopharynx during the first 3 to 5 days of symptomatic illness. Real Time PCR assay targets specific genes and can be used for diagnosis of SARS-CoV-2 virus infection.

Positive result indicates that RNA from SARS-CoV-2 was detected in the specimen, and the patient is considered infected with the virus and presumed to be contagious. Negative test result for this test means that SARS-CoV-2 RNA was not detected in the specimen

Limitations: Negative results do not preclude COVID-19 and must be correlated with clinical observations, patient history, and epidemiological information.

· Positive results do not rule out bacterial infection or co-infection with other viruses.

• The sensitivity of the assay is dependent on the timing of the specimen collection (in relation to symptom onset/stage of infection), quality, and type of the specimen submitted for testing

• Follow-up testing may particularly be important if patient has a clinical picture of viral pneumonia, a potential exposure history, and/or radiographic findings (chest CT or MRI scan) consistent with COVID -19 pneumonia. However repeat testing in the near-term after clearance (within 90 days) should be avoided as prolonged shedding of non-viable virus is not uncommon

• Ct values generated from different assay systems within the same laboratory, or from different laboratories, are not directly comparable and do not necessarily reflect the same viral load due to inter-assay and inter-laboratory variability.

Variation in timing of sample collection, fluctuations in virus shedding, and difference between detection limit of different testing methods within same or different labs could lead to variation in results particularly during initial phase of infection.
If the virus mutates in the rRT-PCR target region, 2019-nCoV may not be detected or may be detected less predictably. Inhibitors or other types of interference may

The performance of this test has not been established for monitoring treatment of 2019-nCoV infection.

Note: Test is performed using ICMR approved Kit targeting any of these genes - E/RDRP/N/ORF1AB

References: 1. Euro Surveill 2020 25, 2. Druce et al. JCM. 2011, 3. N. Engl. J. Med. 2020, 382, 929-936

End Of Report

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

Dr.Swati Pravin Mulani Lab Head





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SRL Diagnostics

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Test Report Status <u>Final</u>	Results	Biological Reference Interval Units
ICMR Registration No: SRLLIPMH AADHAAR NO 9726 1782 9686		
CLINICAL INFORMATION :		
REFERRING DOCTOR : SELF		CLIENT PATIENT ID :
DRAWN : 16/11/2021 17:33	RECEIVED : 16/11/2021 19:21	REPORTED : 16/11/2021 21:57
ACCESSION NO : 0285UK001486	AGE : 40 Years SEX : Male	
PATIENT NAME : VISHAL VINAY	AKRAO ASTUNKAR	PATIENT ID : VISHM301119800A

CONDITIONS OF LABORATORY TESTING & REPORTING		
 It is presumed that the test sample belongs to the patient named or identified in the test requisition form. All Tests are performed and reported as per the turnaround time stated in the SRL Directory of services (DOS). SRL confirms that all tests have been performed or assayed with highest quality standards, clinical safety & technical integrity. A requested test might not be performed if: a. Specimen received is insufficient or inappropriate specimen quality is unsatisfactory b. Incorrect specimen type c. Request for testing is withdrawn by the ordering doctor or patient d. There is a discrepancy between the label on the specimen container and the name on the test requisition 	 The results of a laboratory test are dependent on the quality of the sample as well as the assay technology. Result delays could be because of uncontrolled circumstances. e.g. assay run failure. Tests parameters marked by asterisks are excluded from the "scope" of NABL accredited tests. (If laboratory is accredited). Laboratory results should be correlated with clinical information to determine Final diagnosis. Test results are not valid for Medico- legal purposes. In case of queries or unexpected test results please call at SRL customer care (91115 91115). Post proper investigation repeat analysis may be carried out. 	
form	SRL Limited	
	Fortis Hospital, Sector 62, Phase VIII, Mohali 160062	







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