



SAMEDAYLABORATORY

Patient

Name David Eidenfield
Birth Date 09/19/2001
Sex Male
Phone (912)-239-0352
Passport # United States of America (the) 659052820

Specimen

Order ID MMT7GR
Collected at 01/04/2022 17:57
Received at 01/04/2022 21:56
Reported at 01/05/2022 02:32
Report Status FINAL

Provider

Name Dr. Hirenkumar Italia
Contact team@sameday-testing.com
Address 7101 Democracy Boulevard, Bethesda, MD

Covid-19 PCR Test

SARS-CoV-2 RT-PCR Nasal Swab

Result

Not Detected (Negative)

The COVID-19 RT-PCR test is a real-time reverse transcription polymerase chain reaction (rRT-PCR) test, also known as a nucleic acid amplification test (NAAT), for the qualitative detection of nucleic acid from SARS-CoV-2 in upper respiratory specimens (such as nasopharyngeal or oropharyngeal swabs, saliva, and nasal swabs) collected from individuals suspected of COVID-19. This test has been validated for performance by Quickmed Diagnostic, Inc. that is certified under the Clinical Laboratory Improvement Amendment 2003 (CLIA), 42 U.S.C. §263a, to perform high complexity tests. This test has been authorized only for the detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens. Detected result is considered a positive test result for COVID-19. This indicates that RNA from SARS-CoV-2 (formerly 2019-nCoV) was detected, and the patient is infected with the virus and presumed to be contagious. If requested by public health authority, specimen will be sent for additional testing. A Not Detected (negative) test result for this test means that SARS-CoV-2 RNA was not present in the specimen above the limit of detection. A negative result does not rule out the possibility of COVID-19 and should not be used as the sole basis for treatment or patient management decisions. If COVID-19 is still suspected, based on exposure history together with other clinical findings, retesting should be considered in consultation with public health authorities. Laboratory test results should always be considered in the context of clinical observations and epidemiological data in making a final diagnosis and patient management decisions. Please review the "Fact Sheets" and FDA authorized labeling available for health care providers and patients. For details visit <https://www.cdc.gov/coronavirus/2019-ncov/hcp/index.html>.

All patient management decisions should be based on clinical judgement of a qualified health care professional.

These results are not intended to be used as the sole means for clinical diagnosis or patient management decisions.

Performing Laboratory

Quickmed Diagnostic Inc.
7600 Leesburg Pike Suite 110, Falls Church,
VA
(424) 250-6633

Lab Director Signature

Lindsay Stevenson

CLIA#

49D2231959

