The novel coronavirus disease (COVID-19) is a new virus of global health significance caused by infection of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). On March 11, 2020, the World Health Organization (WHO) declared COVID-19 a global pandemic.

TRANSMISSION AND SYMPTOMS
According to the U.S. Centers for Disease Control and Prevention (CDC), COVID-19 is thought to spread from person to person in close contact through respiratory droplets. It is also possible that a person can catch COVID-19 by touching a surface or object that has the virus on it.

The CDC reports that the following symptoms may appear 2–14 days after exposure:

- Fever
- Cough
- Shortness of breath

DIAGNOSTIC TESTING FOR COVID-19
BioReference offers a real-time reverse-transcription polymerase chain reaction (real-time RT-PCR) assay, giving healthcare providers accurate and timely test results to ensure greater access to testing, promote earlier diagnosis and help limit the subsequent spread of infection. The test detects the presence of SARS-CoV-2 and is for use with patients who meet current guidance for evaluation of infection with COVID-19.

The test has been made available pursuant to the US Food and Drug Administration (FDA) Emergency Use Authorization (EUA) for diagnostic testing in CLIA Certified high-complexity laboratories. The test has been validated, and is being performed at BioReference.

Please refer to the ICD-10 Coding Guidelines for more information on coding encounters related to 2019 novel coronavirus (COVID-19).
**GUIDANCE FOR TESTING**

The CDC has released criteria for healthcare providers for the evaluation of patients under investigation (PUI) of COVID-19. As of March 4, 2020 recommendations include testing for a wider group of symptomatic patients. Providers should use their judgment to determine if a patient has signs and symptoms compatible with COVID-19 and whether the patient should be tested. Decisions on which patients receive testing should be based on the local epidemiology of COVID-19, as well as the clinical course of illness.


**Healthcare providers should notify their local or state health department immediately in the event of a patient under investigation for COVID-19.**

**COLLECTION INSTRUCTIONS**

Patients under investigation of COVID-19 and seeking evaluation of the disease will not be collected at BioReference Patient Service Centers. Specimen should be collected at physician offices, hospitals or other clinic settings. As of March 13, the CDC recommends collecting only the upper respiratory nasopharyngeal (NP) swab. Collection of an oropharyngeal (OP) specimen is a lower priority, and, if collected, should be combined in the same tube as the NP swab and immediately place in 2-3 mL of viral transport media. Refrigerate specimen at 2-8° C. Label with patient name. Place in specimen bag and label with “COVID-19” and submit to laboratory.

Please refer to the most current CDC guidelines for further information on collecting, handling, and testing clinical specimens. [www.cdc.gov/coronavirus/2019-nCoV/lab/guidelines-clinical-specimens.html](http://www.cdc.gov/coronavirus/2019-nCoV/lab/guidelines-clinical-specimens.html)

*The COVID-19 assay provides you with accurate and timely test results to provide proper care and reduce subsequent spread of infection.*
STANDARD PROCEDURES FOR NASOPHARYNGEAL COLLECTION

1. Tilt patients head so nasal passages are parallel to the palate.
2. Insert a swab into nostril. Leave the swab in place for several seconds to absorb secretions.
3. Slowly remove swab while rotating it.

STANDARD PROCEDURES FOR OROPHARYNGEAL COLLECTION

1. Swab the posterior pharynx, avoid touching the tongue, teeth, and gums.
2. Remove swab.
TEST DETAILS

Test Code and Name:
TH68 Novel Coronavirus COVID-19 Nasopharynx
TH69 Novel Coronavirus COVID-19 Oropharynx
TH71 Novel Coronavirus COVID-19 Pooled NP/OP

Primary Container: Dacron-tipped plastic swab with universal transport media (Speedy# 510)
MAY INCLUDE ONE OF THE FOLLOWING:
• M6 MicroTip Flock Swab
• M4 MicroTip Flock Swab
• M6 Universal Flock Swab
• Star Swab

Alternate Container: Swab Viral Culturette (Speedy# 509)

Turn Around Time*: 3 Days

Transportation Temperature: Refrigerate (2-8˚ C)

Stability: 48 Hours (Refrigerated) 30 Days (Frozen)

Methodology: Real Time RT-PCR

Reference Range: Not Detected

Result Comments:
• Positive 2019-nCoV: Critical
• Presumptive Positive 2019-nCoV: Critical.
  Re-collection of a new sample is advised.
• Inconclusive: Consider Re-Collection As Clinically Indicated
• Invalid: Consider Re-Collection As Clinically Indicated
• Not Detected: Consider Re-Collection As Clinically Indicated

NOTE: All results, positive, negative and inconclusive, will be reported to the respective state health departments through ELR (Electronic Laboratory Reporting).

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AOEs: Source: □ Oropharyngeal □ Nasopharyngeal


CPT: 87635**

Clinical Utility: For the detection of the novel COVID-19, Coronavirus

* TAT is based upon receipt of the specimen at the laboratory.
**CPT codes provided are based on AMA guidelines and are for informational purposes only. CPT coding is the sole responsibility of the billing party. Please direct any questions regarding coding to the payer being billed.
SOURCES


FOR MORE INFORMATION PLEASE VISIT

www.bioreference.com/coronavirus/
OR CALL 833-684-0508

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