FACT SHEET FOR PATIENTS
VITROS Immunodiagnostic Products Anti-SARS-CoV2 Total Reagent Pack
Ortho-Clinical Diagnostics, Inc.

You are being given this Fact Sheet because your sample(s) was tested for the Coronavirus Disease 2019 (COVID-19) using the VITROS Immunodiagnostic Products Anti-SARS-CoV-2 Total Reagent Pack test.

This Fact Sheet contains information to help you understand the risks and benefits of using this test for the diagnosis of COVID-19. After reading this Fact Sheet, if you have questions or would like to discuss the information provided, please talk to your healthcare provider.

• For the most up to date information on COVID-19 please visit the CDC General webpage:
  • https://www.cdc.gov/COVID19

What is COVID-19?
COVID-19 is caused by the SARS-CoV-2 virus. The virus, which can cause mild to severe respiratory illness, was first identified in Wuhan, China, and has now spread globally, including the United States. There is limited information available to characterize the spectrum of clinical illness associated with COVID-19 but it likely spreads to others when a person shows signs or symptoms of being sick (e.g., fever, coughing, difficulty breathing, etc.).

What is the VITROS Immunodiagnostic Products Anti-SARS-CoV-2 Total Reagent Pack test?
The test is designed to detect antibodies against the virus that causes COVID-19 in serum and plasma.

Why was my sample tested?
You were tested because your healthcare provider believes you may have been exposed to the virus that causes COVID-19 based on your signs and symptoms (e.g., fever, cough, difficulty breathing), and/or because:

• You live in or have recently traveled to a place where transmission of COVID-19 is known to occur, and/or
• You have been in close contact with an individual suspected of or confirmed to have COVID-19.

Testing of the specimens will help find out if you may have COVID-19.

What are the known and potential risks and benefits of the test?

Potential risks include:

• Possible discomfort or other complications that can happen during specimen collection.
• Possible incorrect test result (see below for more information).

Potential benefits include:

• The results, along with other information, can help your healthcare provider make informed recommendations about your care.
• The results of this test may help limit the spread of COVID-19 to your family and others in your community.

What does it mean if I have a positive test result?
If you have a positive test result, it is very likely that you have or had COVID-19 and you have developed an antibody response to the virus. You may need isolation to avoid spreading the virus to others, or you

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may have been previously infected. Your healthcare provider will work with you to determine how best to care for you based on the test results along with other factors of your medical history, and your symptoms, possible exposures, and geographic location of places you have recently traveled. There is also the small chance that this test can give a positive result that is wrong (a false positive result).

What does it mean if I have a negative test result?
A negative test result means that the antibodies to the virus that causes COVID-19 was not found in your specimen. However, it is possible for this test to give a negative result that is incorrect (false negative) in some people with COVID-19. A negative result may occur if you are tested early in your illness and your body hasn’t had time to produce antibodies to infection. This means that you could possibly still have COVID-19 even though the test is negative. If this is the case, your healthcare provider will consider the test result together with all other aspects of your medical history (such as symptoms, possible exposures, and geographical location of places you have recently traveled) in deciding how to care for you.

It is important that you work with your healthcare provider to help you understand the next steps you should take.

Is this test FDA-approved or cleared?
No. This test is not yet approved or cleared by the United States FDA. When there are no FDA-approved or cleared tests available, and other criteria are met, FDA can make tests available under an emergency access mechanism called an Emergency Use Authorization (EUA). The EUA for this test is supported by the Secretary of Health and Human Service’s (HHS’s) declaration that circumstances exist to justify the emergency use of in vitro diagnostics for the detection and/or diagnosis of the virus that causes COVID-19. This EUA will remain in effect (meaning this test can be used) for the duration of the COVID-19 declaration justifying emergency of IVDs, unless it is terminated or revoked by FDA (after which the test may no longer be used).

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