



Fact Sheet for Recipients: CovWipe-19 SARS-CoV-2 Oral Rapid Test

This fact sheet has been provided to inform you on the potential risks and benefits of the CovWipe-19 Oral, a rapid test for the qualitative detection of SARS-CoV-2 IgG and IgM antibodies in human oral fluid. It may also help you understand when, why, and how a test can be applied to monitor your COVID-19 status. The CovWipe-19 was developed by Oranoxis for emergency use to identify or confirm SARS-CoV-2 infection. T

Please read the full fact sheet, and if you have any additional questions feel free to reach out to Oranoxis or your healthcare provider for more information.

What are the symptoms of SARS-CoV-2 (COVID-19)?

According to the World Health Organization (WHO) and the Centers for Disease Control and Prevention (CDC), the most common symptoms for COVID-19 are fever, dry cough, and fatigue. Other symptoms such as loss of taste or smell, sore throat, and difficulty breathing were reported. Once infected it may take 2-14 days before an individual demonstrates symptoms.

Please check the CDC webpage for the most up-to-date information regarding COVID-19.

What do I need to know about COVID-19 testing?

The CovWipe-19 can be used to test human oral fluid. The test measures human IgM and IgG that are generated by the immune system in response to the virus.

Positive results can be indicative of active infection with SARS-CoV-2, but does not rule out co-infection with other diseases. The agent detected may not be the definite cause of disease. Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for patient management decisions. All results must be combined with clinical observations, patient history, and epidemiological information.

All specimens should be collected with appropriate infection control precautions and personal protective equipment (PPE).

For the current guidance on COVID-19 infection control precautions please visit the CDC website and search for the "Where Can I go for updates and more information."

Why & when should I take a CovWipe-19 Oral Test?

If you are exhibiting symptoms of COVID-19 please contact your healthcare provider and discuss options for testing. CovWipe-19 is a noninvasive oral fluid test that can act as a preliminary screening device that can monitor SARS-CoV-2 antibody responses at variable stages of the infection.

If you are not exhibiting symptoms and would like to test or regularly monitor your COVID-19 status, CovWipe-19 offers a quick and affordable solution. Acting in parallel with other existing testing methods, the test device can be implemented as an initial filter, a regularly applied test for monitoring, a primary form of identification of infected individuals, or secondary form of confirmation for a nucleic acid test.

Consider taking a CovWipe-19 if you have come in contact with individuals who are suspected or confirmed to have been infected with COVID-19. CovWipe-19 can also be taken to consider your status before returning to the work force or coming in contact with the public, family, or friends. Positive results should always be reported to your healthcare provider and related authorities

What do the test results mean?

For information on how to interpret test results please see the CovWipe-19 IFU.

A positive test result on the CovWipe-19 indicates that the antibodies to COVID-19 were detected, and the individual is possibly infected and can transmit COVID-19 to other individuals.

When IgM or IgG antibodies are present it can indicate that a subject has had a recent or prior infection with COVID-19. Positive results for IgM or IgG does not mean that a subject's existing symptoms are a result of COVID-19 infection.

A negative test result with the CovWipe-19 means that the COVID-19 specific antibodies were not present in the sample provided by the donor. The donor may have been infected recently and antibodies to COVID-19 are yet to be detectable.

The CovWipe-19 has been optimized to minimize the rates of false positives and false negatives. Other confirmation methods should always be used to reduce risk.

Report Adverse Events, such as problems with test performance or results, to Oranoxis at info@oranoxis.com and to MedWatch by submitting Form 3500 to the FDA. (<https://www.accessdata.fda.gov/scripts/medwatch/index.cfm?action=reporting.home>) or by calling 1-800-FDA-1088.

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Risks to an individual of a false negative result include: delayed or lack of supportive treatment, lack of monitoring of infected individuals and their household or other close contacts for symptoms resulting in increased risk of spread of COVID-19 within the community, or other unintended adverse events.

What is an EUA?

The United States FDA has made this oral test available under an emergency access mechanism called the Emergency Use Authorization (EUA). The EUA is supported by the Secretary of Health and Human Service's declaration that circumstances exist to justify the emergency use of in vitro diagnostics (IVDs) for detection and/or diagnosis of the virus that causes COVID-19.

Any device or test kit made available under an EUA has not undergone the same type of review as an FDA approved or FDA cleared IVD. An EUA may be issued when criteria are met when no other approved alternatives exist. Based on existing scientific evidence, it is reasonable to believe that this IVD may be effective in the detection of COVID-19.

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).

More Information and Resources

CDC webpages:

General: <https://www.cdc.gov/COVID19>

Healthcare Professionals: <https://www.cdc.gov/coronavirus/2019-nCoV/guidance-hcp.html>

Information for Laboratories: <https://www.cdc.gov/coronavirus/2019-nCoV/guidance-laboratories.html>

Laboratory Biosafety: <https://www.cdc.gov/coronavirus/2019-nCoV/lab-biosafety-guidelines.html>

Isolation Precautions in Healthcare Settings: <https://www.cdc.gov/coronavirus/2019-ncov/infection-control/controlrecommendations.html>

Specimen Collection: <https://www.cdc.gov/coronavirus/2019-nCoV/guidelines-clinical-specimens.html>

Infection Control: <https://www.cdc.gov/coronavirus/2019-ncov/infection-control/index.html>

FDA webpages:

General: www.fda.gov/novelcoronavirus

EUAs: <https://www.fda.gov/medicaldevices/emergency-situations-medical-devices/emergencycuse-authorizations>

Manufacturer Information

Oranoxis Inc., located in San Diego, CA, is specialized in the development, manufacture, and marketing of rapid immunoassay diagnostic products for rapid testing biochemical samples in oral fluid/saliva.

The CovWipe-19 Oral is designed, laboratory tested, and manufactured by the following:

Oranoxis Inc.
7438 Trade St.
San Diego, CA 92121

Contact Information:

(Email) info@oranoxis.com

(Website) www.oranoxis.com

Report Adverse Events, such as problems with test performance or results, to Oranoxis at info@oranoxis.com and to MedWatch by submitting Form 3500 to the FDA. ((<https://www.accessdata.fda.gov/scripts/medwatch/index.cfm?action=reporting.home>) or by calling 1-800-FDA-1088.