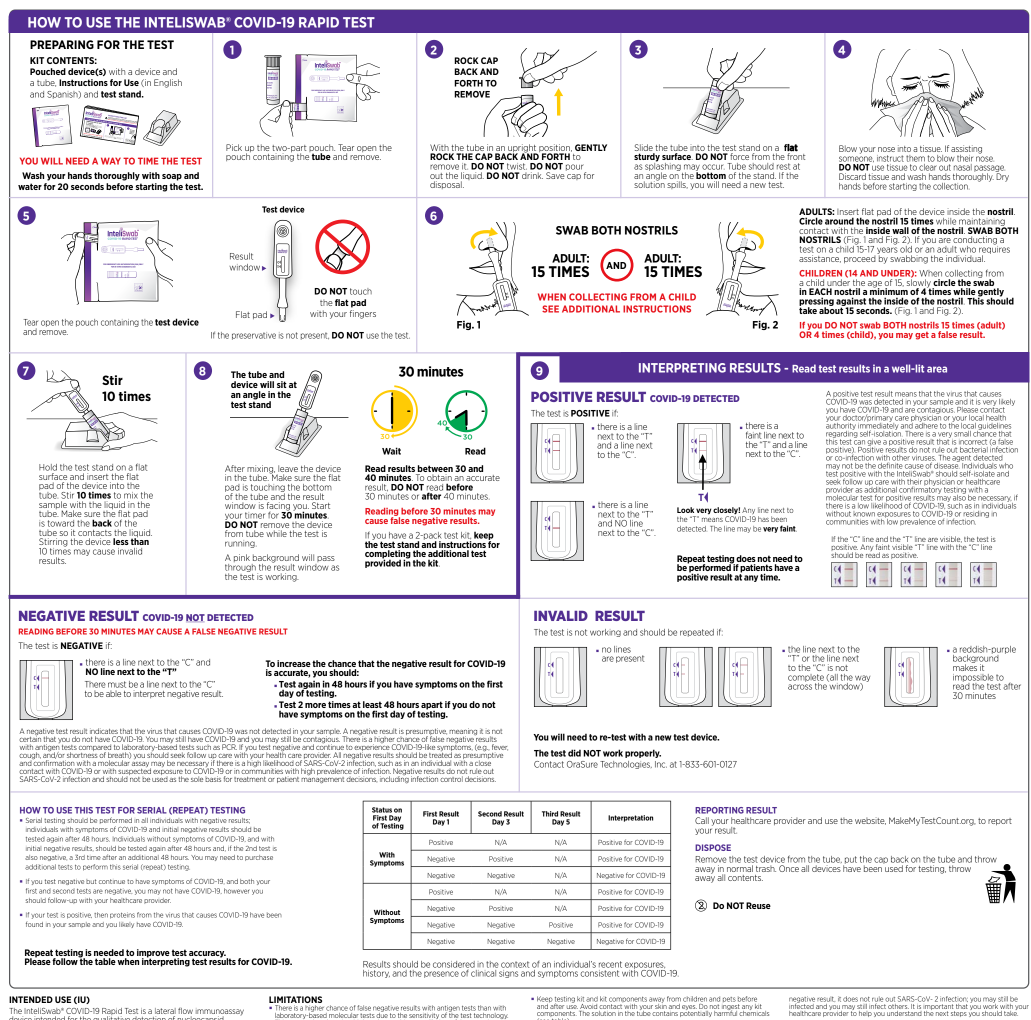


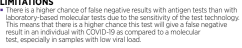
INSTRUCTIONS FOR USE

You must follow the test directions carefully to get an accurate result. Call OraSure Technologies at 1-833-601-0127 or visit www.InteliSwab.com to obtain the complete instructions for use. FOR USE UNDER EMERGENCY USE AUTHORIZATION (EUA) ONLY. IMPORTANT: Swabbing the nostrils is critical for obtaining an accurate result. If you do not swab your nose, the device will produce a false negative result.



The InteliSwab[®] COVID-19 Rapid Test is a lateral flow immunoassay device intended for the qualitative detection of nucleocapsid protein antigen from the SARS-CoV-2 virus.

This test is authorized for non-prescription home use with self-This test is authorized for non-prescription home use with self-collected anterior nasal (nares) swab samples from individuals aged 18 years or older, or adult-collected anterior nasal (nares) swab samples from individuals aged 2 years or older. This test is authorized for individuals with symptoms of COVID-19 within the first seven (7) days of symptom onset when tested at least twice over three days with at least 48 hours between tests, and for individuals without symptoms or other epidemiological reasons to suspect COVID-19 when tested at least three times over five days with at least 48 hours between tests. with at least 48 hours between tests.



The performance of this test was established based on the evaluation of a limited number of clinical specimens collected between February through September 2021. The clinical performance has not been established for all circulating variants but is anticipated to be reflective of the prevalent variant in circulation at the time and location of the clinical evaluation. Performance at the time of testion may use depending on the variants circulation of the testion of the testion of the clinical evaluation. at the time of testing may vary depending on the variants circulating, including newly emerging strains of SARS-CoV-2 and their prevalence, which change over time.

All COVID-19 antigen test negative results are presumptive and confirmation with a molecular assay may be necessary. If you continue to have symptoms of COVID-19, and both your first and second tests are negative. you may not

Keep testing kit and kit components away from children and pets before and after use. Avoid contact with your skin and eyes. Do not ingest any kit components. The solution in the tube contains potentially harmful chemic (see table)

(see table).		
Chemical Name	GHS Code for each Ingredient	Concentrations
Triton X-100	H302, harmful if swallowed H315, skin irritation H318, serious eye damage H400, short-term (acute) aquatic hazard H410, long-term (chronic) aquatic hazard	0.2%
ProClin 950	H302, harmful if swallowed H332, harmful if inhaled H314, causes severe skin burns and eye damage H317, may cause an allergic skin reaction H335, respiratory inflation H410, long-term (chronic) aquatic hazard	O.1%

The InteliSwab® COVID-19 Rapid Test does not differentiate between SARS-CoV-1 and SARS-CoV-2

Results are for the identification of SARS-CoV-2 nucleocapsid results are for the identification of SARS-COV-2 Mucleocapsid protein antigen. Antigen is generally detectable in anterior nasal swab specimens during the acute phase of infection. Positive results indicate the presence of viral antigens, but clinical correlation with past medical history and other diagnostic information is necessary to determine infection status. Positive results do not rule out bacterial infection or co-infection with other viruses and the agent detected may not be the definite cause of disease. Individuals who test positive with the InteliSwab* COVID-19 Rapid Test should self-isolate and seek follow-up care with their physician or healthcare provider as additional testing may be necessary.

All negative results are presumptive and confirmation with a molecular assay, if necessary, for patient management, may be performed. Negative results do not rule out COVID-19 and should not be used as the sole basis for treatment or patient management not be used as the sole basis for treatment of patient management decisions, including infection control measures such as isolating from others and wearing masks. Negative results should be considered in the context of an individual's recent exposures, history and the presence of clinical signs and symptoms consistent with COVID-19.

Individuals who test negative and continue to experience COVID-19 like symptoms of fever, cough and/or shortness of breath may still have SARS-CoV-2 infection and should seek follow up care with their healthcare provider.

Individuals should report all results obtained with this product to their healthcare provider and using the MakeMyTestCount.org website. MakeMyTestCount.org will report all test results received from individuals who use the authorized product to relevant public health authorities in accordance with local state, and federal requirements using appropriate LOINC and SNOMED codes, as defined humber and strate in the construction of the state of defined by the Laboratory In Vitro Diagnostics (LVID) Test Code Mapping for SARS-CoV-2 Tests provided by CDC.

The InteliSwab® COVID-19 Rapid Test is authorized for nonprescription self-use and/or as applicable an adult lay user testing another person 2 years of age or older in a non-laboratory setting.

The InteliSwab® COVID-19 Rapid Test is only for use under the Food and Drug Administration's Emergency Use Authorization. This product has not been FDA cleared or approved.

have COVID-19, however you should follow-up with a healthcare provid

- If the test is positive, then proteins from the virus that causes COVID-19 have
- been found in the sample and you likely have COVID-19. This test is read visually and has not been validated for use by those with impaired vision or callor impaired vision.
- ed vision or color-impaired vision. ct test results may occur if a specimen is incorrectly collected or

WARNINGS, PRECAUTIONS, AND SAFETY INFORMATION

- Read all instructions carefully before performing the test. Failure to follow the instructions may result in inaccurate test results
- In the USA, this product has not been FDA cleared or approved, but has been authorized by FDA under an Emergency Use Authorization. This product has been authorized only for the detection of proteins from SARS-CoV-2, only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 300bb-3(b)(1), unless the declaration is terminated or authorization is revoked sooner.

Serial testing should be performed in individuals with negative results at least twice over three days (with 48 hours between tests) for symptomatic individuals and three times over five days (with at least 48 hours between tests) for asymptomatic individuals. You may need to purchase additional tests to perform this serial (repeat) testing. If you have had symptoms longer than 7 days, you should consider testing at least three times over five days with at least 48 hours between tests.

- An anterior nasal swab sample can be self-collected by an individual age 18 years and older. Children age 2 to 17 years should be tested by an adult.
- Do not use on anyone under 2 years of age.
- Wear a safety mask or other face-covering when collecting a specimen from a child or another individual.
- Do not use if any of the test kit contents or packaging is damaged.
- Test components are single-use. Do not re-use
- Do not use kit past its expiration date.
- Do not touch the swab tip.
- Once opened, the test swab should be used immediately
- Do not read test results before 30 minutes or after 40 minutes. Results read before 30 minutes or after 40 minutes may lead to a false positive, false negative, or invalid result.

If the solution contacts the skin or eyes, flush with conious amounts of wate If irritation persists, seek medical advice: https://www.poisonhelp.org or 1-800-222-1222.

- · For more information on EUAs please visit: https://www.fda.gov, emergency-preparedness-and-response/mcm-legal-regulatory-and-policy-framework/emergency-use-authorization
- For the most up to date information on COVID-19, please visit: www.cdc.gov/COVID19

FREQUENTLY ASKED QUESTIONS

What are the known and potential risks and benefits of this test? Potential risks include

- Possible discomfort during sample collection
- Possible incorrect results (see Warnings and Interpreting Results sections for more information)

Potential benefits include

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

For more information on EUAs go here: https://www.fda.gov/emergency-preparedness-and-response/mcm-legal-regulatory-and-policy-framework/ emergency-use-authorization

What is the difference between a COVID-19 antigen test and a m

There are different kinds of tests for the SARS-CoV-2 virus that causes COVID-19. Nolecular tests detect genetic material from the virus. Antigen tests, such as the InteliSwab[®] COVID-19 Rapid Test, detect proteins from the virus. Due to the lower sensitivity of antigen tests, there is a higher chance this test will give you a false negative result when you have COVID-19 than a molecular test would.

What if you test positive? A positive result means that it is very likely you have COVID-19 because proteins from the virus that causes COVID-19 were found in your sample. You should selfisolate from others and contact a healthcare provider for medical advice about your positive result.

What if you test negative?

A negative test result indicates that antigens from the virus that causes COVID-19 were not detected in your sample. However, if you have symptoms of COVID-19, and your first test is negative, you schould test again in 48 hours since antigen tests are not as sensitive as molecular tests. If you do not have symptoms and received a negative result, you schould test at least two more times with 48 hours in between tests for a total of three tests. If you have a

This serial testing approach is recommended to minimize the risk of incorrect results. For more information on the performance of the test and how the performance may apply to you, please refer to the performance data in the Healthcare Provider Instructions for Use (IFU), available at www.inteliswab.com

hat antigen tests more accurately determ h the virus that causes COVID-19 when ta

What does an invalid test result mean? An invalid result means the test was not able to tell if you have COVID-19 or not. If the test is invalid, a new test device should be used to collect a new nasal specimen and you should test again.

If you have a test line and no control line? If you have a test line and no control line, your test is positive. When the level of virus in the sample is high, the line next to the "C" may not be present or may be very faint. The line next to the "C" must be visible to read a negative test result.

Why do I have a test line and no control line?

How accurate is the InteliSwab* COVID-19 Rapid Test?

IMPORTANT Do not use this test as the only guide to manage your illness. Consult your healthcare provider if your symptoms persist or become more severe. Individuals should provide all results obtained with this product to their

For information about current expiration dates for at-home OTC COVID-19 diagnostic tests, visit http://www.fda.gov/covid-tests

EXPLANATION OF SYMBOLS

REF Batch Code	Use By
Do Not Reuse	Caution, Consult Accompanying Documents
Temperature Limitation	Manufacturer
LOT Catalog Number	Consult Instructions for Use
IVD In Vitro Diagnostic Medical Device	

MORE QUESTIONS ABOUT THE INTELISWAB® COVID-19 RAPID TEST?

-601-0127 or visit www.InteliSwab.com

The InteliSwab[®] COVID-19 Rapid Test Letter of Authorization, authorized Fact Sheets and authorized labeling are available on the FDA website and www.InteliSwab.com

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