

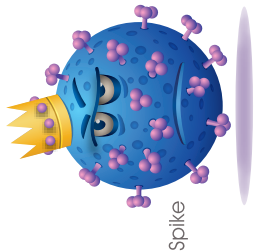
osom[®] COVID-19 Antigen Home Test

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2 TESTS

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Rapid Results in 15 Minutes

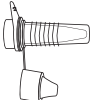
- KIT CONTENTS:**
- 2 x Individually pouched Test Devices
 - 2 x Tubes containing 300 µl of buffer liquid
 - 2 x Sterile Swabs
 - 4 x Tube holders (backside of carton)
 - 1 x Quick Reference Instructions

MADE IN THE USA

CONTENTS



Single-use test device in a foil pouch



Collection tube with buffer liquid



Nasal swab package



Quick Reference Instructions



IVD



15°C 59°F



1067-2

REF

3459-3

Distributed by:

SEKISUI Diagnostics, LLC

6659 Top Gun Street

San Diego, CA 92121 USA

30°C 86°F



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osom[®] COVID-19 Antigen Home Test

For an Emergency Use Authorization (EUA) only

The OSOM COVID-19 Antigen Home Test is intended for the qualitative detection of SARS-CoV-2 nucleocapsid protein antigen in anterior nasal (nares) swab samples. This test does NOT determine if you had COVID-19 in the past or if you have immunity.

Determining a negative result requires multiple tests. You may need to purchase additional tests to perform serial (repeat) testing. This test is more likely to give you a false negative result when you have COVID-19 than a lab-based molecular test.

If you test positive for COVID-19 please contact your doctor/primary care physician or your local health authority immediately and adhere to the local guidelines regarding self-isolation.

For home testing (ages 2 and up).



Scan QR Code to view video instructions.

TUBE HOLDERS

A

B

C

D

SEKISUI

DIAGNOSTICS

Because every result matters™

sekisuidiagnostics.com



Item necessary to use the test but not provided in the test kit is a timer. Use within 30 minutes after opening the foil pouch. Avoid contact of the extraction liquid in Tube with skin and eyes.



For more information on expiration dating for COVID-19 antigen tests, please refer to <http://www.fda.gov/covid-tests>

The emergency use of this product is 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. § 360bbb-3(b)(1), unless the declaration is terminated or authorization is revoked sooner.

The OSOM COVID-19 Antigen Home Test is a lateral flow immunoassay that uses antibodies to detect nucleocapsid protein antigen from SARS-CoV-2 in anterior nasal swabs. The test is authorized for individuals with symptoms of COVID-19 within the first five (5) days of symptom onset, when tested twice over three days with at least 48 hours between tests or individuals without symptoms or other epidemiological reasons to suspect COVID-19, when tested at least three (3) times over five (5) days and at least 48 hours between tests.

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2024-10-31