

in a new normal. NAVICA displays results from the 15-minute Abbott BinaxNOW™ COVID-19 Ag Card, an antigen rapid test, to help you and others make

informed decisions.

NAVICA-enabled COVID-19 test sites are coming soon. Visit **NAVICA.ABBOTT** to sign up to be one of the first to know when and where the COVID-19 tests are being offered in your community.

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NAVICA" AND THE NAVICA PASS

SHARE ENCRYPTED DIGITAL COVID-19 RESULTS

How to get a NAVICA Pass

1 Download the NAVICA app and create your secure profile

Show your profile's unique NAVICA ID to get tested at a NAVICA-enabled test site Receive a NAVICA Pass when you test negative for COVID-19





WHAT IS NAVICA?

NAVICA is an app that allows you to receive and store your encrypted BinaxNOW™ COVID-19 Ag Card test results and manage your NAVICA Pass.

WHAT IS THE NAVICA PASS?

The NAVICA app will display a digital NAVICA Pass via a QR code, similar to an airline boarding pass, for anyone who has received a negative result from a NAVICA-enabled test site using the BinaxNOW COVID-19 Ag Card test. Sharing your pass with NAVICA-enabled organizations such as employers and schools verifies authenticity of your negative COVID-19 test results.

HOW CAN I GET NAVICA?

NAVICA, the first-of-its-kind app, can be downloaded from both the App Store® and Google Play™ at no charge. Go to your app store, search NAVICA and download the blue NAVICA app.









HOW DO I FIND A NAVICA-ENABLED TEST SITE?

NAVICA-enabled test sites administering the 15-minute BinaxNOW COVID-19 Ag Card test are coming soon. Visit **NAVICA.ABBOTT** to sign up to be one of the first to know when and where the COVID-19 tests are being offered in your community.

WHAT KIND OF TEST IS USED?

The BinaxNOW COVID-19 Ag Card is a rapid lateral flow antigen test administered by a healthcare professional or a trained operator. It tests for active infection using a direct nasal swab. It provides a positive or negative result in 15 minutes with no instrument required.

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The BinaxNOWTM COVID-19 Ag Card EUA has not been FDA cleared or approved. It has been authorized by the FDA under an emergency use authorization for use by authorized laboratories. The test has been authorized only for the detection of proteins from SARS-CoV-2, not for any other viruses or pathogens, and is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of *in vitro* diagnostic tests for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Act, 21 U.S.C. § 360bbb-3(b) (1), unless the authorization is terminated or revoked sooner.

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