Humasis Co., Ltd.
Site 1: Rm. 114, 502, 504, 604, 604-1,

O CELUTION USA Distributed by CELLTRION Celltrion USA, Inc One Evertrust Plaza Suite 1207, Jersey City, New Jersey, 07302, USA Tel: (201) 499-1844

IVD OTC





EASY Sample Collection



Results in 15 Minutes

Celltrion DiaTrust[™] COVID-19 Ag
Home Test













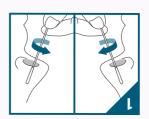












Only open the aluminum pouch when you are ready to do the test.

the test, pay particular attention to the instructions on how to swab your nose. This test involves taking a sample from deep inside your nose. When performing Wash or sanitize your hands and dry them thoroughly before starting the test.

Please carefully read the precautions outlined in the Instructions for Use manual prior to starting your test. Then please refer to the mobile app and follow the detailed instructions required to collect your sample.

PRECAUTIONS BEFORE THE TEST



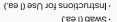






INTERPRETATION OF RESULT





· Filter Cap (1 ea.)

· Test Tube (Extraction Buffer) (1 ea.)

· Test Device (1 ea.)

CONTENTS

Do not use on children under 14 years of age.

This test is intended for individuals aged 14 years and older only. current infection with the virus that causes $\text{COVID-}\overline{\text{19}}$. This test is intended to be used as an aid in the diagnosis of a For in vitro diagnostic use. For use under the Emergency Use Authorization (EUA) only. $\hfill \Sigma$

COVID-19 Ag Home Test

Celltrion DiaTrust^{IM}







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Results in







15 Minutes

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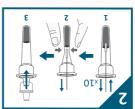


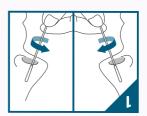












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PRECAUTIONS BEFORE THE TEST









Instructions for Use (1 ea.) · Swab (2 ea.)

· Filter Cap (2 ea.)

· Test Tube (Extraction Buffer) (2 ea.) Test Device (2 ea.)

CONTENTS

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COVID-19 Ag Home Test Celltrion DiaTrust





INTERPRETATION OF RESULT

BEE C1-P60 D-2 05







Collection **EASY** Sample



15 Minutes

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OTC **QVI**

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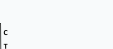












PRECAUTIONS BEFORE THE TEST

Please carefully read the precautions outlined in the Instructions for Use manual prior to starting your test. Then please refer to the mobile app and follow the detailed instructions required to collect your sample.

Wash or sanitize your hands and dry them thoroughly before starting the test.
Make sure they are completely dry.
This test involves taking a sample from deep inside your nose. When performing the test, pay particular attention to the instructions on how to swab your nose.



Scan the QR code through your smartphone (Android 10 or newer, iOS 14.2 or newer) camera to download the free Celltrion DiaTrust™ COVID-19 Ag Home Test App (CELLTRION SAFEKEY). Celltrion DiaTrust™ COVID-19 Ag Home Test App can also be accessed through https://celltrion.safekey.tools if any error occurs with QR code.

Please Follow the Step-by-Step Instructions Available on the Mobile App. Rev. 00 / 2021-07-26 TY / PB-228C2HU04-E005













If you have symptoms of COVID-19, you This test is more likely to give you a false negative result when you have COVID-19 than a lab-based molecular test.



COVID-19 Ag Home Test





- CONTENTS · Test Device (5 ea.) · Test Tube (Extraction Buffer) (5 ea.)
 - · Filter Cap (5 ea.) · Swab (5 ea.)
 - ·Instructions for Use (1 ea.)



The Celltrion DiaTrust™ COVID-19 Ag Home Test is for use under Emergency Use Authorization (EUA) only. This product has not been FDA cleared or approved, but has been authorized by FDA under an EUA. This product has been authorized only for the detection of proteins from FDA under an EUA. This product has been authorized only for the detection of proteins from SARS-COV-2, not for any other viruses or pathogens. The emergency use of this product is only authorized for the duration of the declaration that circumstances exist justifying the authoriza-tion of emergency use of *in vitro* diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(f) of the Federal Food, Drug and Cosmetic Act, 2 I U.S.C. § 360bbb-3(b)(f), unless the declaration is terminated, or authorization is revoked sooner.

