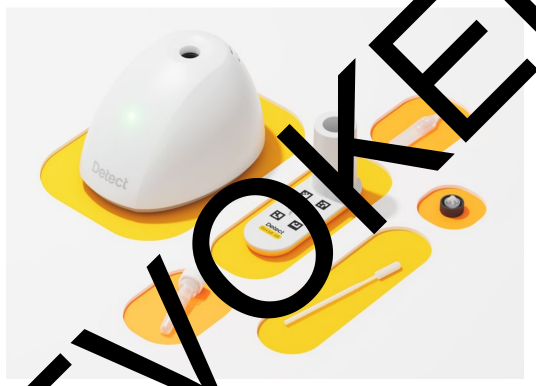
Detect

Detect Covid-19 Test™

Covid-19 Molecular Home Tea



Instructions for Use For mealthcare Providers

For in vitro diagnostic use.

For use under Emergency Use Authorization only.

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1. Intended Use

The Detect Covid-19 Test™ (the Detect test) is a molecular in vitro diagnostic test for the qualitative detection of nucleic acid from the novel coronavirus SARS-CoV-2 that causes Covid-19.

This test is authorized for non-prescription home use with self-collected erior nasal (nasal) swab samples from individuals aged 14 years or older suspected Covid-19. This test is also authorized for non-prescription home use with adulacollect nterior nasal swab samples from individuals aged 2 years or older susp ed of Covi

This test is also authorized for non-prescription home use y self-colle d anteri nasal (nasal) swab samples from individuals aged 14 year r older, dult collecte anterior nasal swab samples from individuals aged 2 years lder thout symmoms wice over or other epidemiological reasons to suspect Covid-1 e days with at least 24 hours (and no more than 48 hours) t ween tests

SARS-CoV-2 viral RNA is generally detectal le in an rior nasal swab cimens during the acute phase of infection. Positive results see of viral RNA, but diagnosus... rmation is clinical correlation with past medical history and necessary to determine infection state s do not rule out bacterial infection or co-infection with other idua bo test positive with the Detect uses. Covid-19 Test should self-isola and seek for with their physician or up ca healthcare provider as additio testing may be ecessa v.

e treated as presumptive and may be confirmed with a d in a coratory, if no essary for patient management. Negative results should molecular assay perform Negative results do not p COV clude S infection and should not be used as the sole basis for treatment or control decisions. Negative anagement decisions for the individual, including infection sults should be considered in the context of an individual's recent exposu istory an he presence of clinical signs and symptoms consistent with Ca **-**19.

duals who test negative and continue to experience Covid-19 like symptoms of d/or shortness of breath may still have SARS-CoV-2 infection and should p care with their physician or healthcare provider.

Test uts will be ported to relevant public health authorities in accordance with local, tate, a requirements, using appropriate LOINC and SNOMED codes, as aboratory In Vitro Diagnostics (LIVD) Test Code Mapping for SARS-CoV-2 Tests provided by the CDC. Automatic test result reporting will be performed by App™ and the Detect secure cloud server.

The Detect Covid-19 Test is authorized for non-prescription self-test by individuals aged 14 years or older and/or, as applicable, for an adult lay user testing another person aged 2 years or older in a non-laboratory setting. The Detect Covid-19 Test is only for use under the Food and Drug Administration's Emergency Use Authorization.

2. **Summary and Explanation of the Test**

An outbreak of pneumonia of unknown etiology in Wuhan City, Hubei Province, China was initially reported to the World Health Organization (WHO) in December 2019. Chinese authorities identified a novel coronavirus SARS-CoV-2 (cause of Co respiratory disease) which has resulted in confirmed human infections work including the United States. Cases of severe respiratory illness and deat have been reported. Patients can become infected with SARS-CoV-2 virus through tact with a contaminated environment or person.

The Detect Covid-19 Test is a molecular in vitro diagnostic te of Covid-19 through the identification of the SARS-CoV-2 specimens.

In asymptomatic individuals (those without Covid-19 e Detect C mptoms Test should be used as a serial test.

What is serial testing?

Serial testing involves testing the same person e times within a few days. Such testing for Covid-19 increases the g infections earlier and should be used for people who are not exp

How do I use the Detect Cov testing? -19 Test for se

sposed to Covice 9 <u>and</u> your first test is negative, you hours but within 48 hours. If your first test is positive, If you're asymptomatic d no take a second test after least then you are likely to hav and should consult with a healthcare Covid-1 provider without waiting to te the second test. If only your second test is positive, then (id-19 and should consult with a healthcare provider. vou are also kely to have



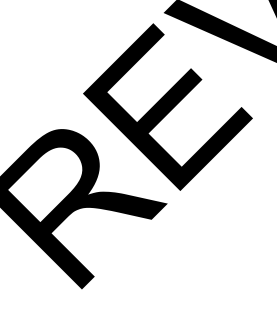
3. **Principles of the Procedure**

The Detect test uses RT-LAMP (Reverse Transcription Loop-mediated Isothermal Amplification) and lateral flow strip technologies to recognize nucleic acids from the Open Reading Frame 1ab (ORF1ab) region of the SARS-CoV-2 genome. The test also identifies nucleic acids from a human gene that serves as a contra sample collection, extraction, reagent integrity, and test execution.

Isothermal amplification occurs at elevated temperature within a dimosable e placed into the reusable Detect Hub™. After amplification, the tube is in ed into th and the tube's liquid wicks onto the lateral flow strip. On the la al flow strip's sa pad, SARS-CoV-2 and control amplicons bind colored partial and flow lateral flow strip's membrane, where they are captured by mobilized itibodies at distinct lines on the strip. A valid negative result must how Sar e Processi Control line. A positive result must show the SARS-1 V-2 line may or ma ot also show the Sample Processing Control line.

In asymptomatic patients, serial testing is re ssist in identifying individuals and facilitate timely infection conctive test result does not rule out infection but repeat testing done on a rease the risks of asis may false negative results.

negative te d be the first of a minimum of For asymptomatic users, an ini sult two tests. An asymptomatic in ∕idual undergoii serial testing with two or more negative results may require oing serial testir or confirmatory testing with this or a different SARS-CoV-2 t, de ding on patient story and potential exposures. An with one or more positive results asymptomatic individua nderd serial testi oes not rule out coinfection with other indicates that SARS-CoV RNA I pathogens.



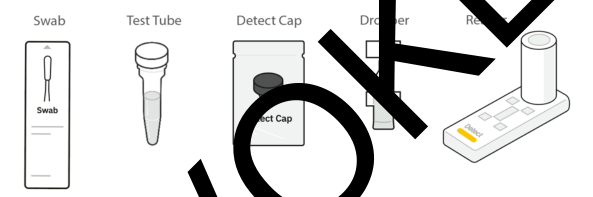
Assay/Reagents 4.

4.1 Materials

The Detect Covid-19 Test contains enough reagents to process on collected sample.

Materials Provided

- Swab (sterilized)
- Test Tube (contains Collection Buffer)
- Detect Cap™ (contains lyophilized reagent bead
- Dropper (contains buffer)
- Reader (contains lateral flow strip inside of



Required but Not Prov

- Detection (Model 2 101, device sold arately
- e of the following: st instructions
 - Detect App (free) and a smartphone visit com/app to see a list of compatible ices and download the app
 - Detect Cover 19 Test Instructions for Use visit de et.com/ifu to download



artphone requirements for the Detect App:

OS Models released after 2018 and using the operating system iOS 13 or Men

Android Models released after 2018 using the Android API level 29 (version 10 or higher) including models from Samsung, Google, and OnePlus. The Detect App is not compatible with Windows smartphones, Android tablets, or iPads.

Warnings and Precautions 5.

5.1 General



- For in vitro diagnostic use.
- For use under FDA Emergency Use Authorization only.
- has been This product has not been FDA cleared or approved authorized by FDA under an EUA.
- cleic acid This product has been authorized only for the detail from SARS-CoV-2, not for any other viruses
- The emergency use of this product is only a horized for the declaration that circumstances exist just ring the author emergency use of in vitro diagnostic and/or diagnosis of Covid-19 under Section 564(b)(1) of the ral Food, ag and less the declaration is Cosmetic Act, 21 U.S.C. § 360111-3(b)(1) terminated, or authorizati is revol
- For more information of EUAs please V https://www.fda.gov/en_gencyprepared essand-response/mcm-legal-regulatory-and-pacy-fra ework/emerge cy-useauthorization
- For the most up to the info Covid-19, please visit: https://www.cdc.gov_pronavirus/zu19-ncov/index.html
- The interest of the tendine does not necessarily correlate to the amount of RS-Covers in the sample.
- samples should be tested as quickly as possible after sample collection.
 - to follow the instructions for use may adversely affect test rmance are or invalidate the test result.
- use the lest components provided. Do not use swabs from other ot re-use any of the components included in the Detect Covid-19 Test. Only the Detect Hub may be re-used for testing other samples.
- Positive results are indicative of the presence of SARS-CoV-2 RNA.



Treat all biological specimens, including used test components, as if capable of transmitting infectious agents. Because it is often impossible to know which might be infectious, all biological specimens should be handled using standard precautions. Guidelines for specimen handling

are available from the U.S. Centers for Disease Control and Prevention [1, 2] and the Clinical and Laboratory Standards Institute [3].

- Do not ingest.
- Keep out of reach of children.
- Avoid contact with skin and eyes.
- Do not apply the Test Tube buffer directly onto the skin or muco membranes or ingest. If contact with the body occurs, r lf irritation persists, seek medical advice. If swallowed a poison center/doctor if you feel unwell.
- Do not apply the buffer contained in the Dropper of the skin mucous membranes or ingest. If contact wit curs, rinse the book water. If irritation persists, seek medical advee. If swalls d: cal poison center/doctor if you feel unwell.

5.2 Storage & Handling

- 9 °F to 86 °F 5°C Store all components at
- until you are ready to perform testing. Do not open componer
- Open all package careful to avoid long small components.
- Do not use Test Tule's that are not or have leaked or spilled.
- Detection ap if its storage pouch is punctured or not fully ontai a freeze-dried bead of reagents that is sensitive d. The noisture.
- the Detect Covid-19 Test past the Use By date on the test box Do not labe
- other than the Detect Hub are single-use and should be componer ed_ after use.
- Follow instructions carefully as shown in the Detect App or the Detect ▶19 Test Instructions for Use.
- Touch only the plastic handle of the Swab with your hands to avoid contaminating the soft tip of the Swab.
- Do not insert the Swab deeper than 1-2 cm into your nose. A deeper swabbing will not yield more accurate results.

See Section 8 for detailed instructions on the control included with the test. It helps indicate whether the reaction is taking place correctly.

5.3 Components & Reagents

- Do not remove the Detect Cap after screwing it onto the Test T
- loving it fr Do not use a Detect Cap that has been dropped after the packaging.
- Section Step 4-5 Do not shake the Test Tube except as described (Test Tube Preparation).
- Begin Test Tube processing for each samp within 1 he the sample.
- Place swab immediately into the Tes bе Failure to do so may result in dried swall l yield an incorrect test result.
- Do not put anything into ie chimney o e Re r until instructed to do so. Doing so may lead indeterminate sults.
- sed for one te Do not reuse Swabs or use a Each single-use yab il Swab other than t one vided in the
- Each single-use Tes Tube is used for one test. Do not reuse Test Tubes.
- se Dete Cap is used for one test. Do not reuse Detect Each sine
- Each single-use Dropper is used for one test. Do not reuse Droppers.
- ngle-use Reader is used for one test. Do not reuse Readers. Each
 - the samples using any protocol other than the one Section 7, Step 4-6 as other protocols have not been tested.
 - Do not tamper with the Reader or attempt to remove the Test Tube once ted into the Reader.

6. Operating Conditions

- The test should be used between 59 °F and 86 °F (15 °C and 30 °C).
 Failure to do so may yield invalid or inaccurate results.
- The test is best used in a room with adequate lighting and away for glare. Failure to do so may result in an inability to see the rest.
- The Hub must be run on a level surface and should run be moved due operation. Failure to do so may yield invalid or inact arate results.
- If a power failure occurs or if the Hub is unplugged table. Test Tubo is in the Hub, the test result is invalid and the per should be retested following the retest procedure described in action 9.2.



7. Procedure

Step 1: Obtain items required but not provided in the test

You will need the items listed below to run your Detect Covid-19 Test. They kems are not included in the Detect test.

- Detect Hub. You can purchase the Hub from Detect, Inc_at detection.
- Smartphone. Go to <u>detect.com/app</u> for the list of commutable smartphones.
- The Detect App installed on your smartphone. Down add a Detect App from detect.com/app.

Instead of using a smartphone with the Detect App, yo can also do not the Detect Covid-19 Test Instructions for Use from <u>Netect</u>, <u>om/ifu</u>. If this is our first time taking the Detect Covid-19 Test, using the latest commended.

Step 2: Prepare to run your test

Carefully read the Detect Covid-1 Test Getting Stated instructions (on the inner lid of the test box) before you run you test. These directions will help you download the Detect App and complete the tracerrectly any safely.

If you do not understand the instructions, the fund that the customer support at support or call toll-free at 855-322-3692 for help.

Step 3: Quant the Detect Approximately our smartphone and follow the instructions

The Forect App X raise pictures, videos, and on-screen instructions to guide you through collecting your naval sample and running the Detect Covid-19 Test. Be sure to call any follow at the instructions given within the Detect App. If you do not, lest make ive approvalid result.

If yo do not understand the directions, contact Detect customer support at sect.com or call toll-free at 855-322-3692 for help.

- You may need to update your Detect App to the latest version before running your test. Follow any on-screen prompts to update the Detect App.
- Within the Detect App, tap on the "Start new test" button.

Step 4: Run the Detect test

Follow on-screen instructions to run the Detect test.

Step 4-1: Get ready to run the Detect test

Review and complete the instructions on the "Getting ready" screens

- Check that the test has not expired by reviewing the Use By on the box.
- Wipe down your work surface and wash your hands, rinse them thoroughly.
- Plug in the Detect Hub.
- Set aside time (approximately 65 minutes) t run the t
- Tap "Next step" to continue to the test active on step.

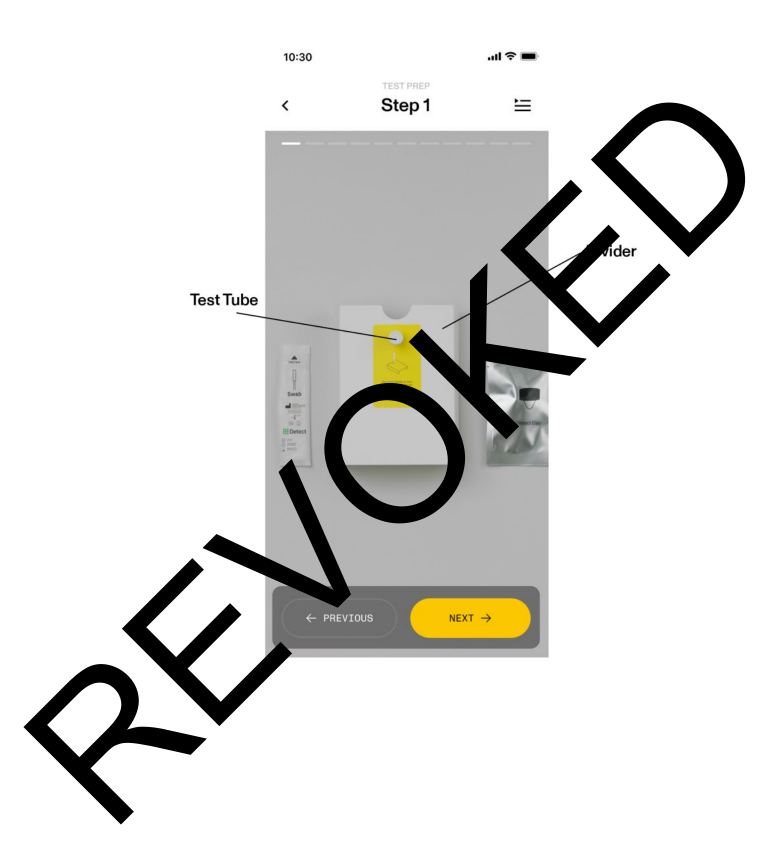
Step 4-2: Activate the Detect to

Open your test and find the divation code of the er lid of the box. Use code or manually enter the 7your smartphone camera to can the actival word activation code printe your test box. n the inner lid

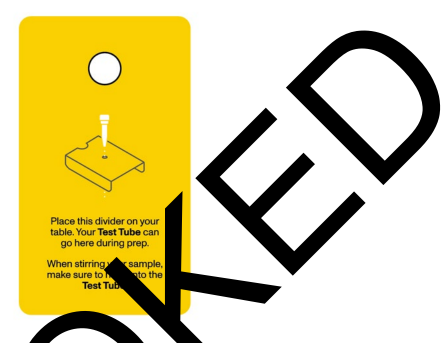
Step 4-3: Gather the components required to run the test

Pack puch and the divider on a flat surface. See the Place the P ving \ aterials that you need to run a test as shown Detect p video here

Swab, Detect Cap, and Test Tube from the Prepare Pack Unpac pou



Follow the directions printed on the divider to place the Test Tube in the divider.



REMINDER: Do not open the Detect Cap and the



Step 4-4: Prepare and collect a nasal sample

Unscrew the cap from the Test Tube. Be careful since the Test Tube contains liquid. Place the Test Tube back into the divider as shown in the Detect App video.



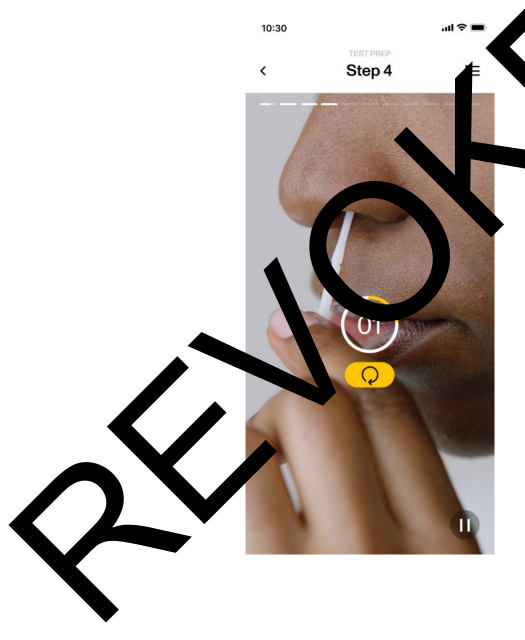
Remove the Swab on the side that says "peel here". Make sure that the up of the Swab does not touch anything besides your nose.

The Detect App video will demonstrate how to collect a nasal sample.

REMINDER: To properly collect a nasal sample, swab both of your nostrils with the same Swab following the directions below.

The Detect test is for use with self-collected samples for individuals aged 14 years and older and for samples collected and tested by an adult caregiver for individuals aged 2 years and older.

• For adult collection, insert the Swab into the nostril until just the soft tip is completely inside (about 1 inch). For pediatric collection insert the into the child's nostril only ½ inch. Swab in a circle around the incae wall of the nostril 5 times. Then gently remove the Swab, insert into our other nostril, and repeat. Make sure the Swab stays in full contact with the inside of the nostril.



 Hold the Test Tube with one hand and fully submerge the Swab tip in the Test Tube and vigorously twirl the Swab for 15 seconds, as shown in the Detect App video. Do not break off the Swab tip. Discard the Swab by placing it back inside its package.



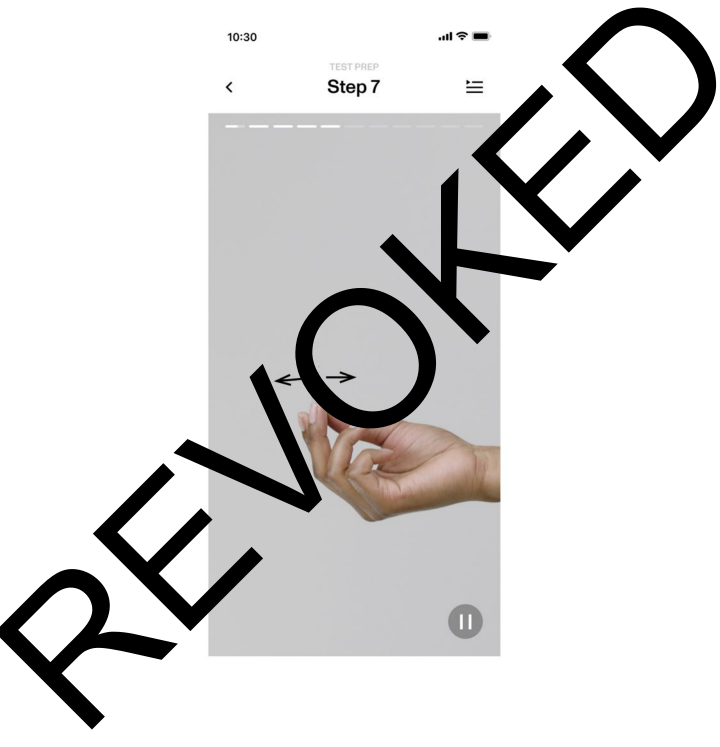
Step 4-5: Test Tube Preparation

REMINDER: Pick up the Detect Cap pouch. While it's closed, gently push the contents inside down toward the bottom of the pouch. The Detect Cap can fly out of the pouch if opened too quickly.

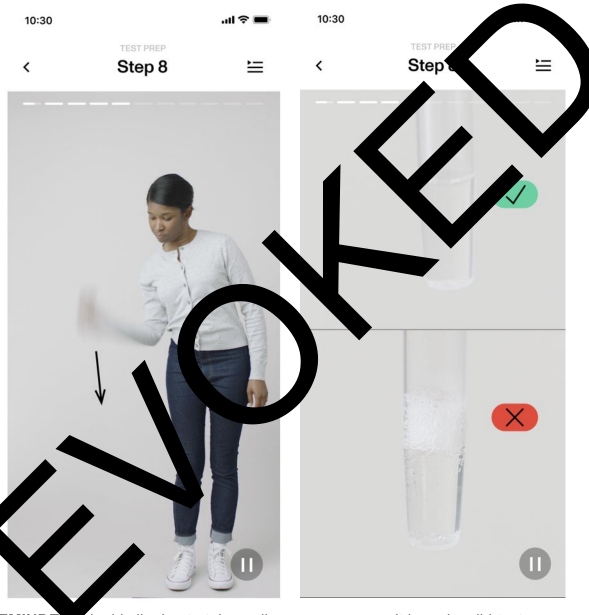
- Slowly and carefully open the pouch and gently remove the Detail Cap.
 You may need to remove the silica packet first.
- On the inside of the Detect Cap you'll see a small white reagent and—this contains the reagents to run your test.
- Screw the Detect Cap onto the Test Tube, tightening as much possible. Do <u>not</u> open the Test Tube after this st
- Hold the Test Tube by the cap, turn it upside down, a make it vigorously side to side for 10 seconds, as so wn in the steet Art video. Then turn the tube right side up again, continuing to hold have cap.



REMINDER: Do not shake the Test Tube up and down. Shake it side to side to ensure the reagent bead contained within the Detect Cap is always in contact with the liquid. This will help dissolve the reagents needed to run the Detect test.



Forcefully bring your whole arm downward to move the liquid to the bottom of the tube. Do this a few times to ensure the liquid is not stuck in the cap or against the walls of the tube.



REMINDER Liquid clinging to tube walls or cap may result in an invalid test.

Step 4-6: Process the sample

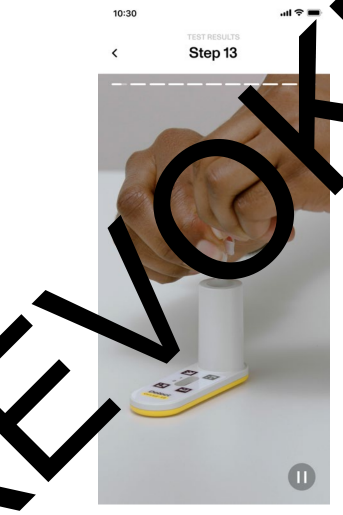
• Place the Test Tube into the well of the Detect Hub, pushing all the way in, as shown in the Detect App video. The Hub will beep once and green light will start blinking. Your sample will start processing automatically and will take 55 minutes. The blinking green light All turn solid when it's complete.



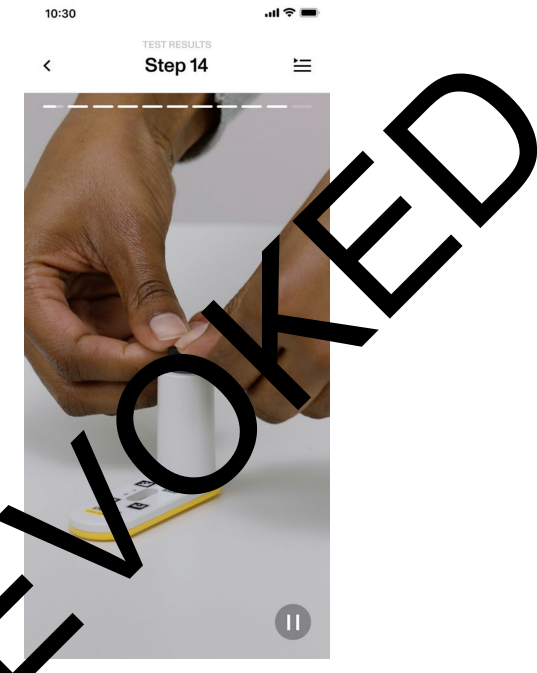
REMINDER: Do not remove the Test Tube from the Hub until it is time to use it.

Step 4-7: Results

- Pick up the Dropper by the rectangular tip and snap your wrist downward to collect the liquid at the bottom as shown in the Detect App video Carefully twist off the tip of the Dropper. Hold the Dropper gently avoid squeezing it and spilling the liquid.
- Insert the dropper as far as it will go into the Reader chimney to spilling, and using both hands, squeeze it firmly to disp as shown in the Detect App video. Discard the Drop

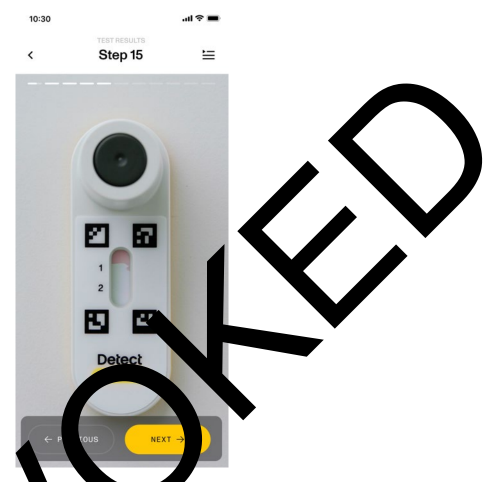


Remove the Test Tube from the Detect Hub, and place it in the chimney of the reader. Using both thumbs, press down on the tube, as shown in the Detect App video. You may hear a pop as the blade at the bottom of the chimney opens the tube, allowing the liquid inside to flow through the Reader.



EMINDE Leep pressing until the tube is completely flush with the lip of Reader chimney. Liquid should begin to flow through the Reader and visible within the Reader's window in about 10 seconds. If there's no flow, firmly tap the Reader against a hard surface 3 times to help trigger flow.

22

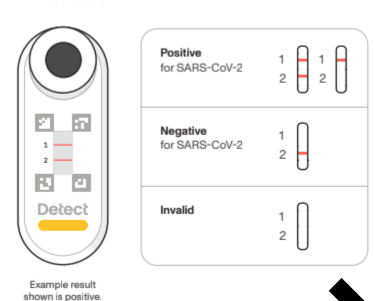


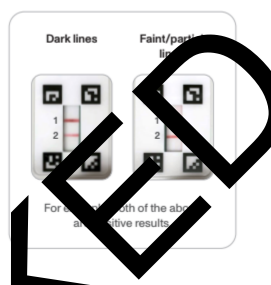
The red lines will take approach darkness and fullness may vary. minutes to fully develop. Line

Do no camper with the Reader or try to remove the est Tube from the pader.

- The Det App will then guide you through identifying the visible lines on the Resear to interpret the result of the test.
- 1 is the to line, which tells you whether SARS-CoV-2 (the virus that ed Cov (19) was identified in your sample.
- Line 2 the control line, which tells you if the test was performed errectly for a negative test result.

As shown below on the right, lines can be faint or incomplete.



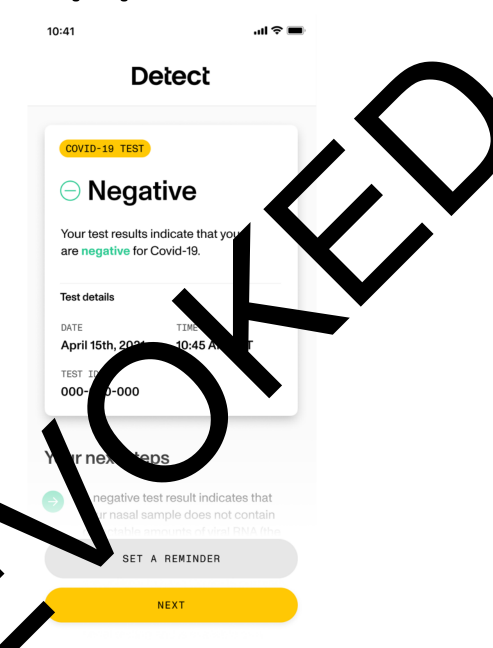


sult Step 5: Understanding the test

gative, Posji e, or Invalid. Please see The app will show res s as Detect's Fact Sheet for dividu (resou s.detect.com) for more information on understating the teamsult.

- of the test result, it is important that when you are sick you Regard tanci and good hygiene. ce socia.
- you develop symptom. or your symptoms persist or become more you are concerned about your health, or if you develop one of ergency warning signs (www.cdc.gov/coronavirus), then you and seek manical attention immediately.

Step 5-1: Understanding a Negative result



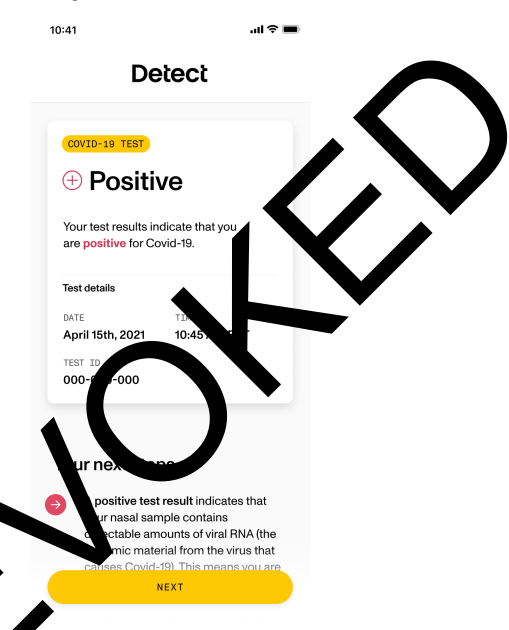
A Negativ alt means that the Detect Covid-19 Test did not identify the SARS-CoV-virus that causes Covid-19 in your sample and it is unlikely that ntly have a Covid-19 infection.

If you are symptomatic and your symptoms persist or become more severe, and you obtained a negative result for your sample, seek help from your HCP even if the results remain negative based on serial testing.

- If you have no symptoms or reasons to suspect Covid-19 infection, this was your first result, and your result was negative, you must collect and test a second sample at least 24 hours after your first test and within the next 48 hours.
- In the following scenarios it is possible that the test may give a neg result even if you have Covid-19 (called a false negative result)
 - A false negative can occur if the sample was not colle d or processed properly.
 - if you are too early or late in your Covid-19 accurately identify a low amount of SAR ∠oV-2 vir sample.
 - A false negative can also occur if t SARS √-2 virus genetic material changes (mutates such that to Dete 19 Test cannot identify the virus.
- For serial testing programs, additional ben a negative result is obtained for the first sample. As al testing may also be necessary if the individual meone who tested positive ed to that can b for SARS-CoV-2 (the vir e Co. 19), or in communities with high numbers of partive cases (high prevalance of infection).
- Even if you do not have tovid-19, you may still have another type of infection. Many over virtues can cause milar symptoms to Covid-19, f vour ent illness. and these may be e caus
- Tell your healthcare ovider whether or not you have symptoms.



Step 5-2: Understanding a Positive result



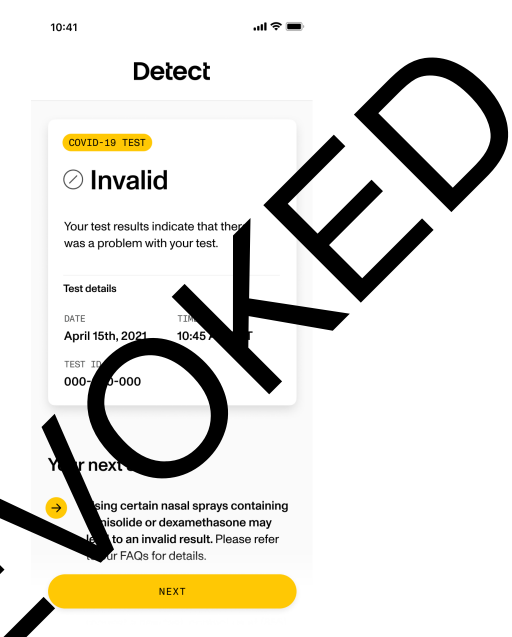
heans the Detect Covid-19 Test identified the SARS-CoV-2 es Covid-19 in your sample, and it is very likely that you rirus that currently have a Covid-19 infection.

- You should self-isolate at home per CDC recommendations to stop spreading the virus to others. Consult the CDC recommendations regarding self-isolation at cdc.gov/coronavirus.
- Consult your healthcare provider as soon as possible and tell him or her that you tested positive for Covid-19 using the Detect Covid-19 Test.

- Tell your healthcare provider if you have symptoms or no symptoms.
- Tell your healthcare provider to view the Detect Covid-19 Test Fact Sheet for Healthcare Professionals at <u>resources.detect.com</u>.
- There is a small possibility that this test can give a positive result the
 wrong (a false positive result) particularly when used in a population
 without many cases of COVID-19 infection. Your healthcare product will
 work with you to determine how best to care for you based on
 results along with medical history, and your symptoms
- There is still a chance of co-infection with another type of illness.
- If you do not have any symptoms, particularly if you live in a carea with low numbers of Covid-19 infections and have had have used to any be diagnosed with Covid-19, additional testing confirm for result may be required.



Step 5-3: Understanding an Invalid result



means that there was an error, and the Detect Covid-19 Test provide a result. You will need to perform a retest.

mmon causes of Invalid results are:

The sample was improperly collected and did not include enough nasal material. Make sure to thoroughly swab both nostrils as directed in Step 4-4.

- The Test Tube was not prepared correctly (either the reagent was not properly dissolved or the liquid was not fully collected into the bottom of the Test Tube). See Step 4-5 for details.
- The processing step was run incorrectly. Make sure you fully insert the Test Tube into the Detect Hub until it beeps and do not remove the Tube until the processing step is complete. See Step 4-6 for detail power failure or interruption during test processing can also invalid result.
- If the result is invalid, retest. You must use a new Deta and a new Swab. Contact Detect customer support support@detect.com or call toll-free at 855-322-2 2 for a re acement test.

Quality Control 8.

8.1 Internal Control

CONTROL

SPC) that Each reaction includes a Sample Processing Contr is shown as a separate line in the reader.

Sample Processing Control (SPC, Line 2 Ensures that sample was processed correctly. The Sample Processing control is lesigne to amplify a human control gene that will be present in swab sa collected by a subject. The SPC determines whether sa ection was performed correctly and whether amplification re tion co ns were appropriate (temperature, time, and reagent mixi). The SP hould positive in a human sample that tests negative a positive in a n sample lacks the SPC still valid. that tests positive; however, a positive seemle the

9. Retests

9.1 Reasons to Retea

If an INVALID test resu occurs, e test once according to instructions in 9.2 (Retes Procedure). If the repeat test fails to produce a valid result, See contact Detect Customer Support at 855-322-3692 or support detect.

have no reason to suspect Covid-19 infection, this was your first result, R was negative, you must collect and test a second sample at ars after your first test and within the next 48 hours.

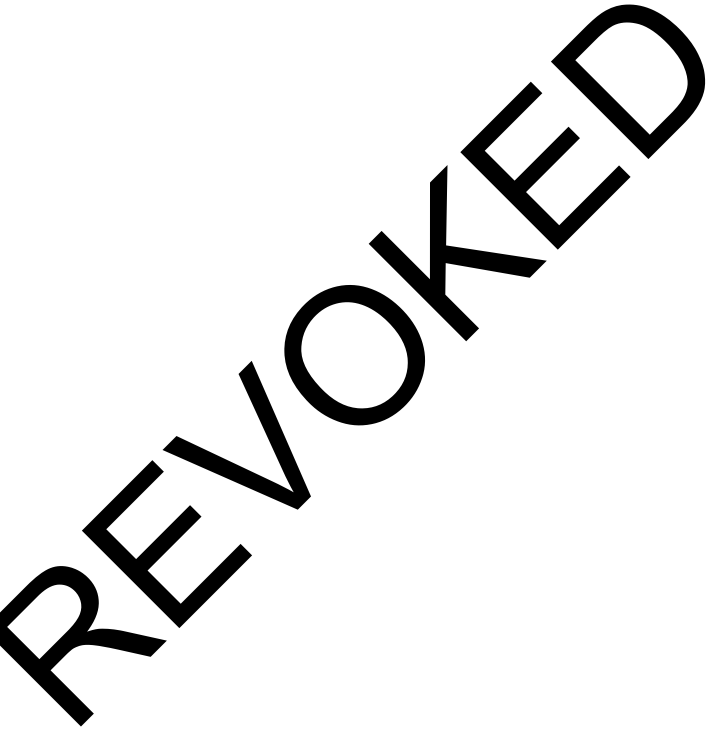
t Prog

- are the work surface as detailed in Section 7.1 (Setup).
- Obtain a new Detect Covid-19 Test.
- 3. Repeat the test procedure outlined in Section 7 (Procedure).

10. Limitations

- Performance has only been evaluated for self-collected nasal swab samples. Use of the Detect Covid-19 Test with other specimen types (such as saliva) has not been evaluated.
- A false negative result may occur if the individual's nose is swale incorrectly. False negative results may also occur if the amour €SARS-CoV-2 virus present on the swab is below the test's limit of detection
- False negative results may occur in patients currently biotin (vitamin B7) supplements. Biotin levels of 0.82 g/mL or higher in the nasal sample may result in incorrect test result false ne
- Invalid results may occur if:
 - The patient is currently using certa nasal spra fiaht allergies, such as dexamethasone flunisolide-c ning lide levels higher than products. Dexamethasone d flu 0.25 mg/mL and 7.5% volume e nasal sample Clivery may result in invalid test results me cases.
 - tions hand soap comes in moisturizin High amounts direct contact th the swab a is introduced into the test should therefo be washed and well-rinsed reaction. Han before from a this test.
- The performance of this test as estates and based on the evaluation of limited number of clinical specimens. Clinical performance has not been s estate shed based on the evaluation of a established with all circulating variants but is anticipated to be reflective of the commerciants in inculation at the time and location of the clinical evaluation. Penetropic at the time of testing may vary depending on the lants circulating. Including newly emerging strains of SARS-CoV-2 and ng newly emerging strains of SARS-CoV-2 and lants circulating, inc. heir prevence, which change over time.
- st cannot rule out diseases caused by other bacterial or viral ogens.
 - should be interpreted together with the patient's medical history, mical signs and symptoms, and the results of other diagnostic ts performed.
- As with other tests, negative results do not rule out SARS-CoV-2 infections and should not be used as the sole basis for patient management decisions.
- This is a qualitative test. Test line intensity is not related to the quantity of virus in the sample.

• The material identified by this test–viral nucleic acid–may persist in the body, even after the patient is not infectious anymore. A positive result on this test does not imply that the patient is infectious, or that the virus is causing the subject's clinical symptoms.



11. Performance Characteristics

11.1 Analytical Sensitivity (Limit of Detection)

The Limit of Detection (LoD) is the lowest amount of virus that car determined in 95% of samples and was determined by testing the test's analytical sensitivity with heat-inactivated SARS-CoV-2 WA1/2020) in pooled nasal matrix. Over two lots of tests, LoD of the Detect Covid-19 Test was determined to be 313 copies r swab, which is from the equivalent to 800 copies per mL if all virus is transfer vab to the buffer.

Viral Load (genomic copies/swab)	Lot 1 SARS- CoV-2 Detection Rate	% Detected	Lot 2 SA CoV-2 Detection	Detected
313	20/20	70%	20/20	100%
156	20/2	100	18/20	90%
78	15/2	75%	14/20	70%

11.2 Analytic Reactivity clusivity)

lico inclusivity ste. s performed to analyze the Detect test's primer ing sequences in the SARS-CoV-2 genome to demonstrate that the mers will zentify all variants of the SARS-CoV-2 virus identified to date and predict inclusivity of the Detect Covid-19 Test. A total of 9 sequences from the GISAID EpiCoV database (www.gisaid.org) Quated the study. 39 seguer were a

Based on it in silico analysis combined with laboratory testing, at least LU.S. viral genomes from the past 90 days are expected to be robustry identified by the Detect test's SARS-CoV-2 primer set.

11.3 Analytical Specificity/Exclusivity (Cross-Reactivity)

The Detect test's cross-reactivity with closely related pathogens, common disease agents, and normal and pathogenic flora that may be present in respiratory tract was tested in triplicate by spiking the organism or germalic material from the organism directly into Detect reactions at the copy attrations listed in the table below. The Detect test showed no interaction who may of the 31 organisms tested. The Detect test was also tested repeatedly with coled human nasal matrix (>500 replicates in all) and showed no interaction.

Organism	Target	Concentration Tested (in final reaction)	# te.	cross- reactivity with Detect
Human coronavirus 229E	Virus	1.00E+05 T^ID ₅₀ /mL	0/3	No
Human coronavirus OC43	Virus	1.00L 2 D ₅₀ /h	0/3	No
Human coronavirus HKU1	Synt dic	6.8% 05 copie 1L	0/3	No
Human coronavirus NL63	Vil	4.00E 4 TCID nL	0/3	No
MERS-coronavirus	Virus	TCID ₅₀ /mL	0/3	No
SARS-coronav.	Is	1.50E+03 TCID ₅₀ /mL	0/3	No
Adenoid 71	Vires	1.00E+05 TCID₅₀/mL	0/3	No
Me eumovirus (hMP	Virus	1.00E+05 TCID₅₀/mL	0/3	No
Parainfluerrus 1	Virus	5.00E+04 TCID ₅₀ /mL	0/3	No
Parainiiuenza virus 2	Virus	1.00E+05 TCID ₅₀ /mL	0/3	No
Parainfluenza virus 3	Virus	1.00E+05 TCID₅₀/mL	0/3	No

Parainfluenza virus 4	Virus	1.60E+04 TCID ₅₀ /mL	0/3	No
Influenza A	Virus	1.00E+05 CEID ₅₀ /mL	0/3	No
Influenza B	Virus	1.00E+05 TCID ₅₀ /mL	0/3	No
Enterovirus 68	Virus	1.00E+05 TCID ₅₀ /mL	0/3	No
Respiratory syncytial virus (Subgroup A)	Virus	1.00E+05 PFU/mL	1/3	
Rhinovirus 89	Virus	8.00E+04 TCID ₅₀ /mL	0/3	N
Chlamydia pneumoniae	Bacteria	1.00E+06 JFU/mL	0/3	No
Haemophilus influenzae	Bacteria	1.6 +06 CFU		No
Legionella pneumophila	Bacteri	C. nL	0/3	No
Mycobacterium tuberculosis	Bac ria	1.00E CFU/	0/3	No
Streptococcus pneumoniae	Bactel	8.8° , 04 J/mL	0/9	No
Streptococcus pyogenes	Leteria	1.00E+06 CFU/mL	0/3	No
Bor ella pertussis	a ja	1.00E+06 CFU/mL	0/3	No
Mycoplas aum e	Bacteria	1.00E+06 CFU/mL	0/3	No
Pne pcystis jiro di (PJP)- erevi e*	Yeast	1.00E+06 CFU/mL	0/3	No
Candida als Zans	Yeast	1.00E+06 CFU/mL	0/3	No
Pseudomonas aeruginosa	Bacteria	1.00E+06 CFU/mL	0/3	No
Staphylococcus epidermis	Bacteria	1.00E+06 CFU/mL	0/3	No

Streptococcus Bacteria 1.00E+06 0/3 No salivarius CFU/mL
--

Due to limited pathogen availability, cross-reactivity was tested with a recombinant version of S. cerevisiae containing genomic material from PJP.

In addition, in silico cross-reactivity analysis of Detect primer sequer was performed by comparing them to representative genomic sequen of the specific respiratory microorganisms below, downloaded from the N database. The table below details all instances of ≥80% hom en a primer and respiratory microorganism genome.

gle SAR Greater than 80% homology was only apparent for a primer with Pneumocystis jirovecii (PJP) and two prime with ndida ntification albicans. Further, none of the labelled primers re uired to amplified nucleic acid target showed ≥80% hom ogy with a of the respiratory microorganism genomes. Therefore, silico analy ufied no potential unintended cross-reactivity of test with the lis respiratory pathogens, including other co.

Organism	SS-CoV-2 primer set
Human coronavirus 229E	no alignment found
Human coronavirus OC43	no alignment found
Human coronavirus HKU	no alignment found
Human coronavirus NL63	no alignment found
MERS-CoV	no alignment found
S/ 3-CoV	no alignment found
Adenovirus .g. C1 Ad. 71)	no alignment found
A Metapneum us (hMPV)	no alignment found
Paraini, 12a 4s 1	no alignment found
Parainfluenza virus 2	no alignment found
Parainfluenza virus 3	no alignment found
Parainfluenza virus 4	no alignment found
Influenza A	no alignment found

Influenza B	no alignment found
Enterovirus (e.g. EV68)	no alignment found
Respiratory syncytial virus	no alignment found
Rhinovirus A	no alignment found
Rhinovirus B	no alignment found
Rhinovirus C	no alignment found
Chlamydia pneumoniae	no alignment found
Haemophilus influenzae	no lignme four
Legionella pneumophila	ne lignment for
Mycobacterium tuberculosis	ne lignment found
Streptococcus pneumoniae	lignment
Streptococcus pyogenes	ne ranment found
Bordetella pertussis	no alignment found
Mycoplasma pneumon	no alignment found
Pneumocystis jirovecii (F	single primer only, 83%
Candida albicans	Two primers at 82% and 89%
Pseugra onas aeras sa	no alignment found
hylococcus epidermidis	no alignment found
taphylo cus salivarius	no alignment found

11.4 Analytical Specificity (Interfering Substances)

Common endogenous and exogenous substances that might be present in clinical nasal swab samples were tested for interference with the Detect Each potentially interfering substance was spiked into both negative resided nasal matrix and contrived positive pooled nasal matrix spiked with eatinactivated SARS-CoV-2 virus at 2X LoD. From these pools, triplical swabs were tested using the Detect test. The interfering substances and the concentrations are listed in the table below. The results show that the Detect test is robust to a wide range of potentially interfering substances.

Interfering Substance	Final Concentration in Nasal Matrix Pool	Negative Sa ples # N gative /# sted	Po ave inples # sitive /# . **	Observed
Rhinocort Allergy	15% v/v		3/3	No
Afrin Nasal Congestion Relief Spray	15% v/v	3	3/3	No
Zicam Cold Remedy Nasal Spray	% v/v	3/5	3/3	No
Chloraseptic Sore Throat Spray	5% v/v	3/3	3/3	No
Flonase Allergy Relief Nasal Spray	7 V/V	3/3	3/3	No
Mupirocin	1 mg/mL	3/3	3/3	No
Neo-Synepm	15% v/v	3/3	3/3	No
Nas Saline Spray	5% v/v	3/3	3/3	No
bramycin	600 μg/mL	3/3	3/3	No
esh w e blood	15%	3/3	3/3	No
	3.5 μg/mL	3/3	1/3*	Yes
Biotin	0.875 μg/mL	3/3	3/3	No
	0.5 mg/mL	2/3**	3/3	Yes
Dexamethasone	0.25 mg/mL	3/3	3/3	No
	15%	2/3**	3/3	Yes
Flunisolide	7.5% v/v	3/3	3/3	No
Mucin	1 mg/mL	3/3	3/3	No

Triamcinolone	15% v/v	3/3	3/3	No
Mometasone nasal spray	1 mg/mL	3/3	3/3	No
Method All-Purpose Surface Cleaner	15% v/v	3/3	3/3	

^{*} The positive samples that could not be detected had false negative results.

Other potentially interfering substances (hand soap and lotion that have present in the home environment were tested for interferen under rea use cases. Hand soap and lotion were tested by having rs wash their diately p with hand soap without rinsing or apply hand lotion im the swab by the tip. These intentionally contaminated abs w then spiked with contrived positive nasal samples at 2x LoD a the Dete nd as Neither of the tested substances interfered with ty to ider e test's SARS-CoV-2.

Interfering Substance	C tion relivery reco	Samples # Positive /# tested	Interference Observed
Dial Antibacterial Liquid Hand Soap	n operator's hand	3/3	No
Aveeno Daily Moisturizin Lotion	On prator's by ds	3/3	No

11.5 mical Expluation

A part ctive, multiplenter clinical study was conducted in the United States in subject's home or a simulated home environment. Testing was performed by the targing subject or the subject's parent/guardian for children under 14. The sixt enrolled symptomatic subjects and asymptomatic subjects with expent exposure, each of whom self-collected two nasal swab samples. For each subject, one swab was collected and sent to a reference laboratory and tested using a high sensitivity FDA-authorized SARS-CoV-2 RT-PCR test by trained laboratory personnel as a comparator, while the other swab was run through the Detect test by the untrained subject or parent/guardian.

Comparing the Detect test's results to those produced by the high sensitivity FDA-authorized SARS-CoV-2 RT-PCR test, the Positive Percent Agreement (PPA) was 90.9.% (30/33) and the Negative Percent Agreement (NPA) was

^{**} The samples without a negative result had invalid results.

97.5% (77/79). Both apparent false positives were due to subject's misinterpreting the test result; the Detect App has since been modified to significantly reduce the frequency of this user error.

Clinical Study Results Summary

	FDA-authorized SARS-CoV-2	2 RT-PCR Assay
Detect	Positive	Negative
Positive	30	2*
Negative	3**	77

^{*}Both of the apparent false positive results were due to subject interpretation the Detect App. A modified version of the Detect App has since en developed an to be effective in reducing the frequency of subjects misinterpreting the **One of the three apparent false negative samples ga tested with a second highly nega

sensitive EUA SARS-CoV-2 RT-PCR assay.

% (3t **3**), (95% CI: 76.4%-96.9%) Positive Percent Agreement Negative Percent Agreeme (NPA): 97 (95% CI: 91.2%-99.3%)

A second evaluation was conducted testing t ability of untrained lay users to correctly identify and eterpris pear-cutoff policive samples with the Detect test. Twelve subjects were such growthree blinged contrived swab samples, a mix of st. Comparing their results to those negative and low-positi (1.9x L 4% (17/18) and the NPA was 100.0% (18/18), expected, the PPA was demonstrate that in the Linds of lay users the Detect Covid-19 Test can esitive amples. reliably

Cutoff Results Summary

	ontrived Sample Expected Result		
Detect	Positive	Negative	
itive	17	0	
Negative	1	18	

Positive Percent Agreement (PPA): 94.4% (17/18), (95% CI: 74.2%-99.0%) Negative Percent Agreement (NPA): 100.0% (18/18), (95% CI: 82.4%-100.0%)

12. Bibliography

- 1 Centers for Disease Control and Prevention. "Interim Guidelines for Collecting, Handling, and Testing Clinical Specimens from Persons for Coronavirus Disease 2019 (Covid-19)." (Refer to latest edition.) https://www.cdc.gov/coronavirus/2019-ncov/lab/guidelines-clinical-specimens.htm
- 2 Centers for Disease Control and Prevention. "Biosafety in Microbiological and Biomedia Laboratories (BMBL) 5th Edition." (Refer to latest edition.) https://www.cdc.gov/labs/PagL.html
- Clinical and Laboratory Standards Institute. "Protection of Laboratory Workers from Capationally Acquired Infections; Approved Guideline." Document M29
 https://clsi.org/standards/products/microbiology/documents/m29/ (Refer to less tedition).

13. Symbols and Abbreviations

The following symbols are used throughout this anual:

Symbol	Definition
À	Biohazard – Pot ly infectious materials. Production by but the beserved.
IVD	or <i>in vitro</i> diag_stic use
CONTROL	ternal control
2	Do h
	Manufacturer

Detect. Whitfield St.

Support: support@detect.com or call toll-free +1 855-322-3692

Detect

Detect[™] Hub User Marual

For a vitro cargnostic use.
For assemble the Detect Covid-19 Test.

The product is for use under Emergency Use Authorization (EUA) only.

REF 21101 MN-00005-US Effective Date: October 2021

Rev 5

Trademark, Patents and Copyright Statements

Detect™ is a trademark of Detect, Inc.

The purchase of this product conveys to the buyer the nontransferable right to use it in accordance with this user manual. No other rights are conveyed expressly, by implication or by estoppel. Furthermore, no rights for resale are conferred with the purchase of this product.

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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 nucleic acid 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 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.



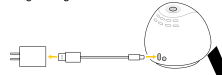
Detect, Inc. 530 Old Whitfield St Guilford, CT 06437 US +1 (855) 322-3692 detect.com

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1 Using the Hub for the Detect Covid-19 Test

1.1 To start your test, you need a Detect Covid-19
Test (sold separately) and the Detect App (free to download). Follow the instructions in the app to prepare your sample using the Detect test.
At the appropriate step, connect the Hub to a wall power outlet via the included USB-C cable and the USB power adapter. The Hub will beep when powered up, and it is ready to use when a solid green light is visible.



Caution: Do not insert the Test Tube into the Hub without powering up.

1.2 When ready, ensure that the Patect Capthe Test Tube is securely the kened. Place the Tube in the Hub until you lear a beat. This indicates that the precessing program has started.



1.3 As anking green as light all indicate that the processing progress in progress.



A solid green LED light and two audible beeps will indicate completion of the process.

This will take approximately 55 minutes.



1.5 Review the next steps in the Detect App before removing the Test Tube from the Hub.

2 Product Information

3 Hub Information

2.1 <u>Proprietary Name</u> Detect Hub

2 . 2 <u>Established Product Name</u> Detect Hub

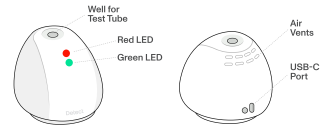
2.3 Intended Use

This Hub is intended for use in combination with the Detect Covid-19 Test for the purpose of detecting SARS-CoV-2 nucleic acids in human anterior nares specimens.

3.1 Ho Component ach Detect Hub co. with the following components:

Cox synent No.1e Detect.	Quantity 1
Cable	1
USB Power Adapter Model S010WU0500200 Input: 100-240V~50/60Hz 400mA Output: 5.0V=2000mA	1
User Manual	1

3.2 <u>Device Diagram</u>



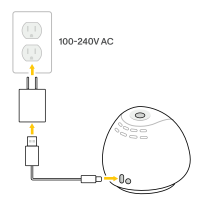
3.3 <u>Hub Specifications</u>

Model	21101
Input power	DC 5V==2A Max
Max. power	10 W
Operating environment	···Indoor Use Only
Operating temperature range	59 °F - 86 ° (15 °C to 30 °
Storage temperature range	32 °F \ 76 °F \ (0 °C to \ °C)
Operating humidity range	25%-8 %
Storage humidity range	0-86-
Max operating altitude	6500 ft (1981.2 m)
Dimension 2 W	4 x 3.4" x 2.6" x 85 × 65 mm)
Net weigh	3.3 oz (93 g)

3.4 Hanstallation

Inly use the USB-Course and USB power adapter colluded with this Detect Hub. Place the least a level, he contal surface in a dry place the strong sunlight and with good ventilating sound the device.

the USB power adapter to the USB-C cable at the back of the Hub, and plug the other end of the power cord into an electrical outlet. The voltage requirement is 100-240V AC.



4 Hub Safety Precautions

- Please read this manual carefully before using the device and follow all recommended safety precautions to avoid personal injury or damage
- → Only use with the included adapter.

to the device:

- → Do not place anything besides Detect Test Tubes into the Hub.
- → Do not open the Test Tube during or after the processing step in order to prevent Hub contamination with biohazardous substance.
- → Keep the Hub dry, dust-free and with good ventilation on all sides. Do not place near a heat source.
- → Unplug the Detect Hub before cleaning.
- → Do not attempt to open or repair the Hub. Cont Detect Customer Support for any or pions.
- → Power off the Hub when no a use by unplugging from the was sutlet.
- → If it will be left unuse for an extra ded period of time, keep the devike sove a with a cloth to protect it from dust.

5 Cleaning a aintenance



Power off the Humbefore Laning. Do not se corror to cleam Landids or fill the well with cleaning solutions.

Clearthe plane surface with cloth dampened with is a contact alcohol. Do not clean the interior wall of the Hub well.

Troubleshooting

If the problem per s, pleas ntact support at: support@r sct.com +1 (8 322-3692

PROBLEM

Hub does not power on (no LEDs are lit)



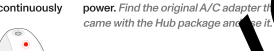
SOLUTION

Check the following connections:

- Hub and USB-C cable
- USB-C cable and A/C adapter
- A/C adapter to outlet If problem persists, contact support.

Cause: Hub is not receiving correct

Red LED fades in/ out continuously



Datacs

Both Red and Green LED fade in/ out continuously



Cause: Hub of sects that its internal and/or external operating temperature is outside the spect and range.

Relocate Hub and environment where to dem, mature is 5°C - °C (59°F - 26°F). Yo may need wait until Hub cools demonstrate before is erational and in.

PROBLEM

Red LED anks continually white Test is uside Hub



SOLL N

ause: Texture be placed in the Hub prior to the Hub being plugged in. Take out the Labe and re-insert. If power was a terrupted during processing, card the test as it is no longer valid—start a new test.

Red Labelinks times after Test ube removal Cause: Test Tube removed before processing step is complete. The test is no longer valid—do not proceed.

Solid Red LED



Cause: Hub detects internal error. Contact support.

Red LED blinks very rapidly, accompanied by a continuous series of beeps Cause: Hub detects a severe internal error. Remove Hub from power immediately. Contact support.

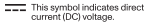
Legend of Symbols

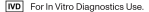


This symbol certifies that the electromagnetic interference from the device is under limits approved by the Federal Communications Commission.



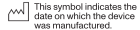
ւ(Սլ) us This symbol indicates that the device has been tested to applicable standards by Underwriters Laboratory (UL).

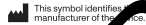






This symbol indicates that caution is necessary when operating the device in order to avoid undesirable consequences. See Section 4 of this manual to see the full list of safety precautions.







This symbol indicates that the instruction manual should be consulted.



This symbol indicates that the device should be kept in dry conditions.



This symbol indicates that the device is fragile and shall be handled with care.



This symbol indicates that the device must not be used if the package holding the device is damaged.



This symbol identified the manufacturer's catalogue number.



tifies This sym



identifies lectronics device the safety ried for irements si II equip C 61140.

Repeat_

reusable and ased wan any additional ct Test

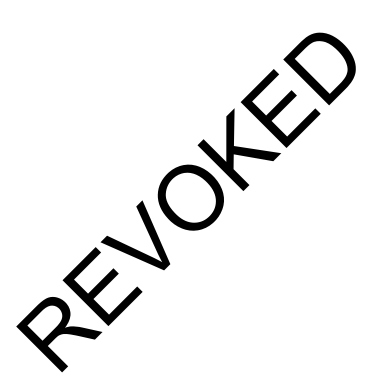
u need to dispose of it, please see the information below.

This electronics product contains substances which could damage the environment if not properly handled. As such, it is not possible to dispose of it in the regular trash.

Please check with your local government or electronics store to verify how to recycle your product. You can also check with your local electronics store to see if they will be able to recycle this product for you.

Depending on the solution chosen, fees may apply.

Detect



Contact support at: support@detect.com +1 (855) 322-3692