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QIAreach® Anti-SARS-CoV-2 Total Test Instructions for Use (Handbook)

For *in vitro* diagnostic use For Emergency Use Authorization Only Rx Only

Version 1





Sample to Insight

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Intended Use

The QIAreach[™] Anti-SARS-CoV-2 Total Test is a rapid, digital lateral flow serological test, using nanoparticle fluorescence, intended for qualitative detection of total antibodies to SARS-CoV-2 in human serum and plasma (sodium heparin, lithium heparin, dipotassium EDTA, and tripotassium EDTA). The QIAreach[™] Anti-SARS-CoV-2 Total Test is intended for use as an aid in identifying individuals with an adaptive immune response to SARS-CoV-2, indicating recent or prior infection. The QIAreach[™] Anti-SARS-CoV-2 Total Test should not be used to diagnose or exclude acute SARS-CoV-2 infection. At this time, it is unknown for how long antibodies persist following infection and if the presence of antibodies confers protective immunity. Testing is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C §263a, that meet requirements to perform moderate or high complexity tests.

Results are for the detection of SARS CoV-2 antibodies. Antibodies to SARS-CoV-2 are generally detectable in blood several days after initial infection, although the duration of time antibodies are present post-infection is not well characterized. Individuals may have detectable virus present for several weeks following seroconversion.

Laboratories within the United States and its territories are required to report all results to the appropriate public health authorities.

The sensitivity of the QIAreach[™] Anti-SARS-CoV-2 Total Test early after infection is unknown. Negative results do not preclude acute SARS-CoV-2 infection. If acute infection is suspected, direct testing for SARS-CoV-2 is necessary.

False-positive results for the QIAreach[™] Anti-SARS-CoV-2 Total Test may occur due to cross-reactivity from pre-existing antibodies or other possible causes. Due to the risk of false positive results, confirmation of positive results should be considered using a second, different SARS-CoV-2 antibody test.

The QIAreach™ Anti-SARS-CoV-2 Total Test is only for use under the Food and Drug Administration's Emergency Use Authorization.

Summary and Explanation of the Test

COVID-19 (coronavirus disease 2019) is the disease caused by SARS-CoV-2 (severe acute respiratory syndrome coronavirus 2) viral infection.¹ The virus is readily transmitted from both symptomatic and presymptomatic²⁻⁴ individuals via respiratory droplets, aerosols, and upper respiratory secretions.⁵⁻⁶ The incubation period is estimated to be 4.6–5.8 days with a median of ~5 days.⁷ The symptoms of COVID-19 are non-specific, ranging from asymptomatic to severe pneumonia and death.⁸ Fever and cough are the most common clinical symptoms but also include shortness of breath, fatigue, muscle aches, headache, new loss of smell or taste, sore throat, congestion or runny nose, diarrhea, and vomiting, which typically appear between 2–14 days following exposure to the virus.⁸⁻¹⁰ Roughly 20% of those infected with SARS-CoV-2 will experience severe symptoms, including Acute Respiratory Distress Syndrome (ARDS) that often requires mechanical ventilation.¹¹

The standard medical practice for definitive diagnosis of active SARS-CoV-2 infection relies on the molecular detection of viral RNA.¹²⁻¹⁴ Typically, IgA and IgM production occur simultaneously and shortly after viral infection with IgM ebbing rapidly in convalescence.¹⁵⁻¹⁶ IgG antibody responses occur after IgM and IgA seroconversion and are longer sustained.¹⁷ The QIAreach[™] Anti-SARS-CoV-2 Total Test detects total antibodies to SARS-CoV-2 and is an indirect test for indicating recent or prior infection in populations of interest. The clinical and public health applications of serologic assays may include support to clinical assessment in persons suspected of having a post-infectious syndrome and understanding transmission dynamics in populations.¹⁸

Principles of the assay

The QIAreach[™] Anti-SARS-CoV-2 Total Test is a rapid, qualitative serological test that detects total antibody responses to expressed SARS-CoV-2 viral antigens, in serum or plasma (sodium heparin, lithium heparin, dipotassium EDTA and tripotassium EDTA).

Antibodies are detected on a single-use, lateral flow, digital detection cartridge (eStick) via nanoparticle fluorescence. The eStick contains state-of-the-art optoelectronic technology and a microprocessor that converts a fluorescent signal into a qualitative readout for the presence of SARS-CoV-2 specific antibodies in patient test samples.

The QIAreach[™] Anti-SARS-CoV-2 Total Test is performed by inserting the eStick into an QIAreach[™] eHub (sold separately). The QIAreach[™] eHub (referred to as eHub hereafter) is a connection hub that provides power to perform multiple QIAreach[™] Anti-SARS-CoV-2 Total Tests simultaneously. The eHub acts as a power source and features a rechargeable lithium battery to allow QIAreach Anti-SARS-CoV-2 Total Tests to be performed when a continuous power supply is not available.

To perform the test, QIAreach Diluent Buffer is first added to the QIAreach Processing Tube and reconstitutes a SARS-CoV-2 viral S1 protein-fluorescent nanoparticle conjugate that is spray dried on an immobilized accretion pad within the tube. Patient serum or plasma is then added to the Processing Tube and mixed with the resuspended conjugate using a pipette. If anti-SARS-CoV-2 antibodies are present in the sample, they will bind to the SARS-CoV-2 S1 antigen-nanoparticle conjugate. The sample is then transferred from the Processing Tube to the eStick sample port.

Once in the eStick, the test sample migrates on a nitrocellulose membrane and across the test line. The migrating anti-SARS-CoV-2 antibodies will bind to immobilized SARS-CoV-2 viral S1 protein at the test line, where they will bridge the two SARS-CoV-2 S1 viral proteins on the test line and in the conjugate. A photosensor will detect light emitted from the fluorescent nanoparticles in the presence of excitation light filtered onto the test line. Signal is interpreted on the eStick firmware and transmitted to the eHub, which then communicates a positive or negative test result to the user by means of a visual display.

QIAreach[™] Anti-SARS-CoV-2 Total test results are determined as Positive or Negative according to the assay result algorithm on the eStick firmware.

Software is available to backup test results, generate test reports, and support online data transfer.

Time required for performing the assay

The time required to perform the QIAreach Anti-SARS-CoV-2 Total Test is estimated below. The time of testing multiple samples when batched is also indicated.

Digital detection:	Approx. 10 minutes for one test
	(1 individual)
<20 minutes labor in total	

Add up to 3 minutes for each extra eStick

Pipet use

This assay requires use of an adjustable volume pipet. Users should familiarize themselves with pipet use prior to performing the QIAreach[™] Anti-SARS-CoV-2 Total Test.

Kit Contents

QIAreach™ Anti-SARS-CoV-2 TotalTest		
Catalog number		645033
Number of tests/pack		60
QIAreach Anti-SARS-CoV-2Total Test Detection System Components*		
QIAreach eStick	Packaged together with Processing Tube in foil wrapper	
	Contains recombinant SARS-CoV-2 S1 protein and human serum albumin	
QIAreach Processing Tube	Packaged together with eStick in foil wrapper	
	Contains an accretion pad that contains spray dried fluorescent nanoparticles conjugated to SARS-CoV-2 S1 viral protein and bovine serum albumin, along with maltose.	
QIAreach Diluent Buffer	Contains bovine serum albumin and ProClin® 300	3 x 10 ml

*See Warnings and Precautions for precautions and hazard statements.

Materials Required but not Provided

Collection tubes for patient serum and plasma (lithium heparin, sodium heparin, dipotassium EDTA, tripotassium EDTA) QIAreach™ eHub (including USB adapter and cable)*- Catalogue# 9003063 QIAreach® Anti-SARS-CoV-2 Controls (Catalogue # 647030)

Equipment required but not provided

Calibrated pipets* for delivery of 50 µl, 150 µl, and 300 µl with disposable tips Optional: Centrifuge for isolating patient serum or plasma Optional: QIAreach Software (Catalogue# 1118894)

Storage and Handling

Kit reagents

Store kit reagents at 2-30°C for up to 4 months.

Stability

The test must be initiated within 60 minutes of opening the foil-wrapped eStick and Processing Tube.

The QIAreach[™] Anti-SARS-CoV-2 Total Test should be performed in a test environment with ≤ 65% relative humidity.

Refer to the expiration date printed on the device labeling for component shelf life.

* See Warnings and Precautions for precautions and hazard statements.

QIAreach Diluent Buffer should be used within 3 months after opening the bottle.

Warnings and Precautions

Warnings

For prescription use only. For in vitro diagnostic use only. For Emergency Use Authorization Only.

This test has not been FDA cleared or approved but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. 263a, that meet the requirements to perform moderate or high complexity tests. This test has been authorized only for detecting the presence of total antibodies to SARS-CoV-2, not for any other viruses or pathogens.

The emergency use of this test 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.

Do not use kits or reagents after the expiration dates shown on the labels.

Ensure test is run away from direct sunlight.

Do not use damaged test kit. Do not reuse the test kit.

Human serum and plasma samples should be considered as potentially infectious. Operators should wear protective clothing, masks, gloves and take other appropriate safety precautions to avoid or reduce the risk of infection.

Precautions

When working with chemicals, always wear a suitable lab coat, disposable gloves, and eye protection goggles. For more information, please consult the appropriate safety data sheets (SDSs).



Handle all human blood serum, and plasma as if potentially (C1) infectious. Observe relevant blood and blood product handling guidelines. Dispose of samples and materials in contact with blood or blood products in accordance with federal, state, and local regulations.

The following hazards and precautionary statements apply to components of the QIAreach[™] Anti-SARS-CoV-2 Total Test kit.

Hazard statements

WARNING	QIAreach Diluent Buffer	(W1)
	Contains: Mixture of 2-methyl-1,2-thiazol-3(2H)-one, and 5- chloro-2-methyl-1,2-thiazol-3(2H)-one. Warning! May cause an allergic skin reaction. Harmful to aquatic life with long lasting effects. Wear protective gloves/ protective clothing/ eye protection/face protection.	

WARNING	QIAreach™ eHub	(W2)
	Do not open the eHub. No serviceable parts inside. Opening of the eHub device could lead to electric shock or damage of the device.	

CAUTION	QIAreacheStick	(C2)
	Do not open the eStick. No serviceable parts inside. Opening of the eStick could lead to user exposure of infectious patient body fluids. Opening the eStick could also damage the eStick device.	

Further information

Deviations from the *QlAreach™ Anti-SARS-CoV-2 Total Test Instructions for Use* may yield erroneous results. Please read the instructions carefully before use. Important: Inspect materials prior to use. Do not use kit if the Diluent Buffer, Processing Tube, or eStick show signs of damage or leakage, or if the seals have been compromised prior to use. Do not handle broken eSticks. Discard used or unused materials and biological samples in accordance with local and government regulations. Do not use the QlAreach™ Anti-SARS-CoV-2 Total Test kit after the expiration date.

Do not mix consumables and reagents from multiple lots.

Procedures

Preparing samples

Note: The QIAreach[™] Anti-SARS-CoV-2 Total Test requires 50 µl of serum or plasma for an individual test.

Follow all instructions provided by the manufacturer of the sample collection device when collecting test specimens. Collect all specimens using standard procedures. Refer to the following guidelines¹⁹ for handling of samples prior to performing the QIAreach[™] Anti-SARS-CoV-2 Total Test.

Samples that require fractionation from red blood cells should be isolated by appropriate means (e.g. centrifugation using a gel separator) prior to analysis. Samples collected in serum tubes should be allowed to clot prior to serum separation.

Samples should optimally be tested as soon as possible following collection and may be held at room temperature (17–27°C) for up to 8 hours prior to testing.

Samples may be stored for up to 2 days at 2-8°C prior to testing.

Samples that require long term storage prior to testing may be stored at ≤ -20°C. Specimens may be frozen and thawed once.

Detection assay

Materials required

QIAreach Processing Tube (packaged together with eStick in foil wrapper) QIAreach eStick (packaged together with Processing Tube in foil wrapper) QIAreach Diluent Buffer

QIAreach™ eHub (with associated power cable and adapter), sold separately

Important points before starting

All samples and reagents (if stored in the refrigerator), must be brought to room temperature (17–27°C) before use. Allow at least 60 minutes for sample equilibration to room temperature. Any clots visible in the samples may be removed by centrifugation.

The eStick and Processing Tube are packaged together in a foil wrapper. The packaging should only be opened before performing the assay. Important: The QIAreach™ Anti-SARS-CoV-2 Total Test must be started within 60

minutes of removing the components from the packaging.

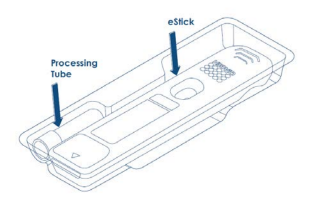


Figure 1. Contents of foil wrapper packaging – QIAreach Processing Tube and eStick.

The eStick is a single use device. It is recommended to label the eStick with test information using a permanent marker or by applying a label directly on the eStick. If a label is applied to the eStick, ensure that the label is not placed over the sample port or the sloped front end (with arrow) of the eStick as this could affect the connection between the eStick and eHub.

There is a small white pad contained within the Processing Tube that is critical component of the QIAreach[™] Anti-SARS-CoV-2 Total Test. DO NOT remove the pad from the Processing Tube. This pad will not be dislodged or come loose during pipetting.

If not connected to a power source, the eHub should have sufficient battery power to complete the test. A fully charged eHub should maintain internal battery power for 8 hours. The QIAreachTM Anti-SARS-CoV-2 Total Test should not be performed if the eHub battery power is less than 10% and is not connected to a power source. The battery LED indicator will display the battery status. The battery level can also be checked by connecting the eHub to a laptop via the provided USB cable and launching the software. The software displays the level of battery charge in the bottom right hand corner of the screen. Refer to the $Q/Areach^{TM}$ eHub User Manual and software quide (provided with the optional QIAreach software) for details.

The eHub comes with a cover to protect the internal ports from dust buildup and contamination. The cover should be placed over the front panel of the eHub when the eHub is not in use.

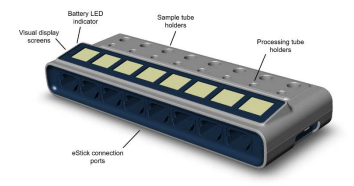


Figure 2. eHub layout. Note: The cover should be in place when the eHub is not in use.

Note: It is recommended to fully charge the eHub in a switched off state overnight (when not in use) or to charge for 4 hours before use. To charge the unit, connect the eHub to a power outlet using the provided USB power adapter and USB cable. We also recommend that the eHub is connected to a USB power source (either a USB adapter or PC) during operation.

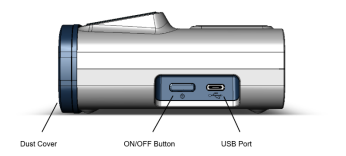
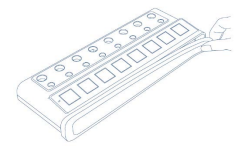


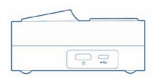
Figure 3. Side panel view of eHub with dust cover, ON/OFF switch and USB connection port.

Procedure

1. Remove the dust cover from the front panel of the eHub and set aside.



2. Press the ON/OFF switch on the right side of the eHub to turn it on.



3. Remove the eStick from the packaging, label with patient identifier, and insert into the eHub.

Note: The test sample must be added to the eStick sample port within 60 minutes of removal from the foil packaging.



4. Remove the Processing Tube from the packaging and insert into the empty tube slot directly in line with the eStick.



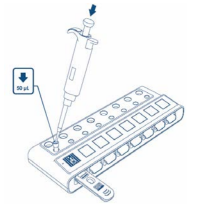
5. Add 300 μl of QIAreach Diluent Buffer to the Processing Tube using a pipet.



6. Carefully remove 50 μl of the patient sample (serum or plasma) from the sample tube using a pipet.



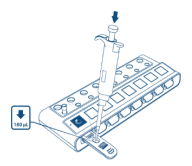
7. Add the patient sample to the Processing Tube containing the QIAreach Diluent Buffer.



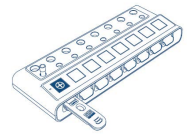
8. With the pipette set to 150 μl, mix the contents of the Processing Tube by pipetting up and down at least 4 times. Take care to not introduce foam while pipetting.



9. Remove 150 µl of the mixed sample from the Processing Tube and dispense into the sample port of the inserted eStick.



10. The test will start automatically within approximately 30 seconds following sample addition, signaled by a countdown timer on the display. Do not remove the inserted eStick until the test is complete and a result is displayed.



11. After completion of the test, the result will be displayed on the eHub.



Positive



Negative

Results Analysis and Test Interpretation

The standard time from sample addition to the eStick to the test result is 10 minutes. The time to result will be less than 10 minutes for positive samples containing elevated anti-SARS-CoV-2 antibody levels. The time to test result will be displayed on the eHub following a positive result.

QIAreach Anti-SARS-CoV-2 Total Test raw data is analyzed on the eStick firmware, which then interprets a positive or negative QIAreach[™] Anti-SARS-CoV-2 Total Test result based on an internal algorithm. The result is transmitted to the eHub, which displays the result. If the optional QIAreach software (Catalogue# 1118894) is used, the eHub will transfer the test result to a computer for data backup and report printing. Please refer to the QIAreach software user manual for detailed instructions about how to use the QIAreach software with the QIAreach Anti-SARS-CoV-2 Total Test.

Quality control of test

All eSticks have built-in controls to ensure reliable performance of the eStick optoelectronics and lateral flow strip and also monitor procedural steps after sample addition to confirm suitability. A failure alert will be communicated to the user in the form of a test error if any fault conditions are detected on the eStick firmware.

Mechanical performance controls are in place to confirm that the eStick components are functioning correctly and are not compromised due to improper handling or transport. Once the sample is added to the eStick, the eStick will continually monitor progress, including the proper flow rate of sample across the strip as well as the correct range of detector particles in the sample. The eStick has at least 48 unique controls built into the firmware to alert the user if the test has not been successfully completed or if the test strip has been compromised, providing an additional level of control over standard lateral flow tests that rely on a single control line. External positive and negative controls are required but not supplied with this kit; and are available for purchase separately from QIAGEN (QIAreach® Anti-SARS-CoV-2 Controls, Catalog # 647030). External controls should be run as outlined in the procedure for testing the samples. Positive and negative controls are required to be tested each time when a new lot is used, when a new operator performs the test, or when the test is run in a new room/laboratory, etc. as a good laboratory practice to confirm the test procedure and to verify proper test performance.

If the test is invalid, an error code will be displayed on the eHub. The test should be repeated if there is \geq 50 µl of patient sample remaining. See Table 7 in the Technical Information section for the list of QIAreach Anti-SARS-CoV-2 Total test error codes.

Interpretation of results

Assessment of QIAreach Anti-SARS-CoV-2 Total Test results is to be performed after the positive and negative controls have been examined, as recommended in Quality Control section above, and determined to be valid and acceptable. If the controls are not valid, the results cannot be interpreted.

QIAreach Anti-SARS-CoV-2 Total Test results are interpreted using the following criteria:

Table 1. Interpretation of QIAreach™ Anti-SARS-CoV-2 Total Test results

QIAreach Anti-SARS-CoV-2Total Test result	Report/Interpretation
Positive (+)	Positive for antibodies for SARS-CoV-2
Negative (-)	Negative for antibodies for SARS-CoV-2

QIAreach Anti-SARS-CoV-2 Total Test results should not be used to diagnose or exclude acute infection. Results are not intended to be used as the sole basis for patient management decisions. Test results should be interpreted in conjunction with clinical observations, patient history, epidemiological information, and other laboratory findings. Important: Laboratories within the United States and its territories are required to report all results to the appropriate public health authorities.

If acute infection is suspected, direct testing for SARS-CoV-2 is necessary.

Limitations

- This test is only to be used in CLIA certified laboratories that meet requirements to perform moderate or high complexity testing and not in point-of-care or athome testing settings.
- This test can only be used for the analysis of serum and plasma (sodium heparin, lithium heparin, dipotassium EDTA and tripotassium EDTA) samples. Do not use with venous or fingerstick (capillary) whole blood samples.

- The test is limited to the qualitative detection of total antibodies specific for the SARS-CoV-2 virus.
- Results from antibody testing should not be used to diagnose acute SARS-CoV-2 infection or to inform infection status. An assay that directly detects the virus should be used to evaluate symptomatic patients for acute COVID-19.
- It is unknown at this time if the presence of antibodies to SARS-CoV-2 confers immunity to infection.
- False negative results may occur for immune-compromised individuals or individuals who receive immunosuppressive therapy.
- Antibodies to SARS-CoV-2 are generally detectable in blood several days after initial infection, although the duration of time antibodies are present post-infection is not well characterized. Individuals may have detectable virus present for several weeks following seroconversion.
- Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for patient management decisions. Antibodies may not be detected in the first few days of infection; The sensitivity of QIAreach[™] Anti-SARS-CoV-2 Total Test early after infection is unknown. Negative results do not preclude acute SARS-CoV-2 infection. If acute infection is suspected, direct testing for SARS-CoV-2 is necessary.
- False positive results may occur due to cross-reactivity from pre-existing antibodies or other possible cause. Positive results may be due to past or present infection with non-SARS-CoV-2 coronavirus strains, such as coronavirus HKU1, NL63, OC43, or 229E.
- A negative result can occur if the quantity of the anti-SARS-CoV-2 antibodies present in the specimen is below the detection limits of the assay, or the antibodies that are detected are not present during the stage of disease in which a sample is collected.
- A positive result may not indicate previous SARS-CoV-2 infection. Consider other information, including clinical history and local disease prevalence, in assessing the need for a second, different serology test to confirm an adaptive immune response.
- Significantly hemolyzed (reddish brown) samples can potentially interfere with the QIAreach[™] Anti-SARS-CoV-2 Total Test optical measurement system. The eStick firmware features built-in controls to determine unacceptably high levels

of hemolysate (> 5mg/ml) and will return an invalid result in the form of an error code if interference is present. Refer to the troubleshooting section if observing elevated hemolysate in samples.

- Not to be used for the screening of donated blood.
- Note: Unreliable results may occur due to deviations from the procedure described in this handbook.
- The performance of this test has not been established in individuals that have received a COVID-19 vaccine. The clinical significance of a positive or negative antibody result following COVID-19 vaccination has not been established, and the result from this test should not be interpreted as an indication or degree of protection from infection after vaccination.
- The performance of this test was established based on the evaluation of a limited number of clinical specimens. The specimens for the negative agreement studies were collected in Pennsylvania, USA between July 2019 and September 2019 and in Florida, USA between August 2018 and October 2019. The specimens for the positive agreement studies were collected in Florida, Louisiana, Texas, and the state of Washington, USA between April 2020 and June 2020 and in Lima, Peru in May 2020. The clinical performance has not been established in all circulating variants but is anticipated to be reflective of the prevalent variants in circulation at the time and location of the clinical evaluation. Performance 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.

Conditions of Authorization for the Laboratory

The QIAreach[™] Anti-SARS-CoV-2 Total Test Letter of Authorization, along with the authorized Fact Sheet for Healthcare Providers, the authorized Fact Sheet for Recipients, and authorized labeling are available on the FDA website: <u>https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/vitro-diagnostics-euas</u>. Authorized laboratories using the QIAreach[™] Anti-SARS-CoV-2 Total Test (referred to as "your product" in the conditions below), must adhere to the Conditions of Authorization indicated in the Letter of Authorization as listed below:

- Authorized laboratories using your product must include with test result reports, all authorized Fact Sheets. Under exigent circumstances, other appropriate methods for disseminating these Fact Sheets may be used, which may include mass media.
- Authorized laboratories must use your product as outlined in the authorized labeling. Deviations from the authorized procedures, including the authorized instruments, authorized clinical specimen types, authorized control materials, authorized other ancillary reagents and authorized materials required to use your product are not permitted.
- Authorized laboratories that receive your product must notify the relevant public health authorities of their intent to run your product prior to initiating testing.
- Authorized laboratories using your product must have a process in place for reporting test results to healthcare providers and relevant public health authorities, as appropriate.
- Authorized laboratories must collect information on the performance of your product and report to DMD/OHT7-OIR/OPEQ/CDRH (via email: <u>CDRH-EUA-</u> <u>Reporting@fda.hhs.gov</u>) and to you (<u>techservice-na@qiagen.com</u>) any suspected occurrence of false positive or false negative results and significant deviations from the established performance characteristics of your product of which they become aware.
- All laboratory personnel using your product must be appropriately trained in immunoassay techniques and use appropriate laboratory and personal protective equipment when handling this kit and use your product in accordance with the authorized labeling. All laboratory personnel using the assay must also be trained in and be familiar with the interpretation of results of the product.

 Authorized distributors, and authorized laboratories using your product will ensure that any records associated with this EUA are maintained until otherwise notified by FDA. Such records will be made available to FDA for inspection upon request.

Performance Characteristics

Clinical performance

Positive Percent Agreement (PPA)/Sensitivity

PPA/Sensitivity was estimated by evaluating serum and plasma samples from study subjects with PCR-confirmed symptomatic SARS-CoV-2 infection. A total of 65 previously collected samples from 65 subjects, collected from April 2020 to July 2020 in the United States (43 samples) and Peru (22 samples), were tested using the QIAreach Anti-SARS-CoV-2 Total Test.

The following table describes the positive percent agreement by time of sampling after onset of symptoms.

Number of days after symptom onset	Number of samples tested	Number of QIAreach™ Anti- SARS-CoV-2 Total Test positive results	Positive percent agreement	95% confidence interval
0 – 7 days	3	2	66.67 %	9.43 - 99.16 %
8 – 14 days	13	12	92.31%	63.97 - 99.81 %
≥ 15 days	49	47	95.92 %	86.02 - 99.50 %
All	65	61	93.85 %	84.99 - 98.30 %

Table 2. Clinical sensitivity by days post-symptom onset

Negative Percent Agreement/Specificity

NPA/Specificity was estimated by evaluating samples collected before the start of the SARS-CoV-2 pandemic (before December 2019). A total of 230 previously collected serum and plasma samples from 230 subjects in the United States were tested using the QIAreach Anti-SARS-CoV-2 Total Test. The following table shows the negative percent agreement.

Table 3. Negative Percent Agreement

Number of samples tested	Number of QlAreach™ Anti-SARS-CoV-2 Total Test negative results	Negative percent agreement	95% confidence interval
230	225	97.83 %	95.00 – 99.29 %

Independent evaluation of clinical performance

The QIAreach Anti-SARS-CoV-2 Total Test was tested on December 10, 2020 at the Frederick National Laboratory for Cancer Research (FNLCR) sponsored by the National Cancer Institute (NCI). Tests were from lot number 86652. The test was evaluated against "Panel 3" which is a panel of previously frozen samples consisting of 30 SARS-CoV-2 antibody-positive serum samples and 80 antibody-negative serum and plasma samples. Each of the 30 antibody-positive samples were confirmed with a nucleic acid amplification test (NAAT) and both SARS-CoV-2 IgM and IgG antibodies were confirmed to be present in all 30 samples. The presence of antibodies in the samples was confirmed by several orthogonal methods prior to testing with the QIAreach Anti-SARS-CoV-2 Total Test. The presence of SARS-CoV-2 antibodies specifically was confirmed by one or more comparator methods. Antibody-positive samples were selected at different antibody titers.

All antibody-negative samples were collected prior to 2020 and include i) seventy (70) samples selected without regard to clinical status, "Negatives" and ii) Ten (10) samples selected from banked serum from HIV+ patients, "HIV+". Testing was performed by one operator using 1 lot of QIAreach Anti-SARS-CoV-2 Total Test. Confidence intervals for sensitivity and specificity were calculated per a score method described in CLSI EP12-A2 (2008).

For the evaluation of cross-reactivity with HIV+, it was determined whether an increased false positive rate among antibody-negative samples with HIV was statistically higher than the false positive rate among antibody-negative samples without HIV (for this, a confidence interval for the difference in false positive rates was calculated per a score method described by Altman). The results and data analysis are shown in the following tables:

Table 4. Summary Results

	Comparator Method			Collected pre-2020		Total
	Antibody Positive		Antibody Negative			
QIAreach Anti- SARS-CoV-2 Total Test	IgM+, IgG+	IgM+, IgG-	IgM-, IgG+	Negative	HIV+	Total
Panlg+	30			2		32
Panlg-				68	10	78
Total	30			70	10	110

Table 5. Summary Statistics

Measure	Estimate	Confidence Interval
Pan Ig Sensitivity	100% (30/30)	(88.7%; 100%)
Pan Ig Specificity	97.5% (78/80)	(91.3%; 99.3%)
Combined Sensitivity	100% (30/30)	(88.7%; 100%)
Combined Specificity	97.5% (78/80)	(91.3%; 99.3%)
Combined PPV for prevalence = 5.0%	67.8%	(35.0%; 88.4%)
Combined NPV for prevalence = 5.0%	100%	(99.4%; 100%)
Combined NPV for prevalence = 5.0%	100%	(99.4%, 100%)
Cross-reactivity with HIV+	0.0% (0/10), not detected	

Important limitations:

- 1. Samples were not randomly selected, and sensitivity and specificity estimates may not be indicative of the real-world performance of the device
- These results are based on serum and ACD plasma samples only and may not be indicative of performance with other sample types, such as whole blood, including finger stick blood.
- 3. The number of samples in the panel is a minimally viable sample size that still provides reasonable estimates and confidence intervals for test performance, and the samples used may not be representative of the antibody profile observed in patient populations.

Cross-reactivity

QIAreach[™] Anti-SARS-CoV-2 Total Test was evaluated for potential cross-reactivity with antibodies to the pathogens listed below. All samples were collected from SARS-CoV-2 negative individuals prior to November 2019 (pre-pandemic). A total of 167 individual specimens were tested. No cross-reactivity was observed.

Pathogen	Sample type	Ν	Number of cross- reactive	Number of non- reactive
Adenovirus	Plasma	28	0	28
Anti-nuclear Antibody (ANA)	Serum	5	0	5
Bordetella pertussis	Plasma	31	0	31
Chlamydia pneumoniae	Plasma	42	0	42
CoV 229E	Serum	17	0	17
CoV HKU1	Serum	5	0	5
CoVNL63	Serum	5	0	5
CoVOC43	Serum	14	0	14
Dengue	Serum	10	0	10
Enterovirus	Plasma	20	0	20
Haemophilus influenzae	Plasma	5	0	5
Hepatitis B- HBc	Serum	5	0	5
Hepatitis B- HBs	Serum	5	0	5
Hepatitis C	Serum	5	0	5
Influenza A	Plasma	63	0	63
Influenza A	Serum	5	0	5
Influenza B	Plasma	72	0	72
Legionella pneumophila	Plasma	12	0	12
Mycoplasma pneumoniae	Plasma	81	0	81
Parainfluenza	Plasma	103	0	103
Respiratory Syncytial Virus (RSV)	Plasma	76	0	76
Respiratory Syncytial Virus (RSV)	Serum	5	0	5

Table 6. Cross-reactivity summary for QIAreach™ Anti-SARS-CoV-2Total test*

* This table evaluates cross-reactivity separately for each pathogen category. With several specimens, containing antibodies to multiple pathogens, some specimens are listed more than once, but in separate pathogen categories.

Interference

The effect of potentially interfering substances on QIAreach[™] Anti-SARS-CoV-2 Total Test was evaluated by spiking endogenous and exogenous interferents at CLSI recommended high level concentrations into SARS-CoV-2 negative plasma and negative plasma spiked with low titer SARS-CoV-2 antibody. No significant interference was observed at the following concentrations:

Bilirubin, conjugated	0.4 mg/ml
Bilirubin, unconjugated	0.4 mg/ml
Hemoglobin*	5 mg/ml
Prednisolone	0.12 mg/ml
Triglycerides	15 mg/ml

* Hemoglobin levels above 5 mg/ml (reddish brown colored samples) can potentially interfere with the QIAreach[™] Anti-SARS-CoV-2 Total Test optical measurement system. The eStick firmware features built-in controls to determine unacceptably high levels of hemolysate and will return an invalid result in the form of an error code if interference is present. See Troubleshooting Guide for more information.

Matrix equivalency

QIAreach Anti-SARS-CoV-2 Total Test performance was evaluated in serum, lithium heparin plasma, sodium heparin plasma, dipotassium EDTA plasma, and tripotassium EDTA plasma by obtaining each sample matrix in a single blood collection from each of five negative donors and one positive donor. Low and Medium level positive samples were contrived by diluting the positive subject sample into each of the negative subject samples, with each sample matrix diluted into the same matrix that was being assessed in order to maintain matrix integrity. The contrived Low and Medium positive samples were tested in parallel with uncontrived negative samples, with 2 replicates tested at each concentration level. QIAreach Anti-SARS-CoV-2 Total Test results were evaluated against the expected

Negative result for uncontrived samples and the expected Positive result for Low and Medium level contrived positive samples. Result agreement is shown in Table 7.

Sample matrix	Negative agreement (n=10)	Low positive agreement (n=10)	Med positive agreement (n=10)	Overall agreement (n=40)
Serum	10/10 (100%)	10/10 (100%)	10/10 (100%)	30/30 (100%)
Lithium Heparin	10/10 (100%)	10/10 (100%)	10/10 (100%)	30/30 (100%)
Sodium Heparin	10/10 (100%)	10/10 (100%)	10/10 (100%)	30/30 (100%)
Dipotassium EDTA	10/10 (100%)	10/10 (100%)	10/10 (100%)	30/30 (100%)
Tripotassium EDTA	10/10 (100%)	10/10 (100%)	10/10 (100%)	30/30 (100%)

Table 7. Sample matrix equivalency test agreement to expected result

Technical Information

Clotted plasma samples

Should fibrin clots occur with long-term storage of plasma samples at or below -20° C, the samples may be centrifuged to sediment clotted material and to facilitate pipetting of plasma.

eHub display icons

Table 8. eHub display icons

lcon	ID	Description
	Please Insert	The eHub port is available for eStick use.
3 ¹¹	Self-test	The eStick has been inserted and a self-test is being performed.
•	Add sample	The eStick is ready for sample addition to the detection port. The sample must be added within 60 minutes of removing the eStick from the foil packaging.
C0/3.46	Processing	The eStick has detected sample and is processing the test. A test countdown timer is displayed. Do not remove the eStick until a result is displayed. The test will take up to 10 minutes.
Anti-CoV2	Positive	Anti-SARS-CoV-2 antibody present
Anti-CoV2	Negative	Anti-SARS-CoV-2 antibody NOT present
X-123	Error	The eStick has encountered an error. The letter below the symbol denotes the type and the numbers are code for the error. Refer to the error code section for more information.

Error codes

The following table lists the error codes in the QIAreach Anti-SARS-CoV-2 test:

Table 0 OlAreach™	' Anti-SARS-CoV-2 Total test e	urror codes categories -	apparal description
	Anti-JANJ-COV-2 Total test e		general description

Error type	Error code format	Description
Self-Test	A-[Error code]	eStick electronic failure
Algorithm	B-[Error code]	Run error or user workflow error
Communication/ Other	C-[Error code]	Invalid data or missed communication betweeneStick and eHub

Table 10. "A" error codes

Error code	Description	Recommended action
A-1	Used eStick	Discard and use new eStick
A-2	Metadata error	Discard and use new eStick
A-4	Metadata error	Discard and use new eStick
A-8	Voltage Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-16	Voltage Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-32	Voltage Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-64	Voltage Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-128	Frequency Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.

Table continued on next page

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Table 10. "A" error codes (cont'd)

Error code	Description	Recommended action
A-256	Frequency Failure	Remove and re-insert the eStick. If error persists, discard and use new eStick.
A-512	Frequency Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-1024	Frequency Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-2048	LED Current Failure	Remove and re-insert the eStick. If error persists, discard and use new eStick.
A-4096	LED Current Failure	Remove and re-insert the eStick. If error persists, discard and use new eStick.
A-8192	LED Current Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-16384	LED Current Failure	Remove and re-insert the eStick If error persists, discard and use new eStick.
A-32768	Dark Frequency Failure	Remove and re-insert the eStick. If error persists, discard and use new eStick.
A-65535	Unknown value	Remove and re-insert the eStick If error persists, discard and use new eStick.

Table 11. "B" error codes

Error code	Description	Recommended action
B-0	No result	Discard and use new eStick
B-8	Conjugate Wave Too Early	Ensure eStick is inserted prior to adding sample. Discard and use new eStick.
B-9	Conjugate Wave Too Early	Check color of sample*. Discard and use new eStick.
B-10	High Dark Frequency	Ensure test is run away from direct sunlight. Discard and use new eStick.
B-12	No Frequency	Discard and use new eStick.
B-13	No Frequency	Discard and use new eStick.
B-14	No Conjugate Wave (Timeout)	Run test within 60 minutes of removing eStick from foil. Check color of sample. Discard and use new eStick.
B-15	Frequency Out of Range	Discard and use new eStick.
B-16	Low Frequency	Ensure sample is mixed well in processing tube prior to adding test sample to eStick. Discard and use new eStick.
B-17	High Frequency	Discard and use new eStick.
B-18	Frequency Out of Range	Discard and use new eStick.
B-19	Low Frequency	Ensure sample is mixed well in processing tube prior to adding test sample to eStick. Discard and use new eStick.
B-21	Peak Data Failure	Check color of sample*. Discard and use new eStick.

Table continued on next page

Table continued from previous page Table 11. "B" error codes (cont'd)

Error code	Description	Recommended action
B-22	Result Timeout	Discard and use new eStick.
B-23	Baseline Issue	Discard and use new eStick
B-24	Baseline Issue	Discard and use new eStick.
B-25	Signal Noise	Discard and use new eStick
B-255	Test Removed Early	Wait for test completion before removing eStick. Discard and use new eStick.

* See Troubleshooting Guide section of applicable kit Instructions for Use regarding optical interference.

Table 12. "C" error codes

Error code	Description	Recommended action
C-0	Connection Error	Remove and re-insert the eStick If error persists, discard and use new eStick.
C-1	Expired eStick	Test is past expiry date. Use an eStick within expiration.
C-2	Sample Not Detected	Run test within 60 minutes of removing eStick from foil. Discard and use new eStick.
C-3	Start Not Acknowledged	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.
C-4	Self Test Failure	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.
C-5	Metadata Failure	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.
C-6	Measurement Data Failure	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.
C-9	Algorithm Failure	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.

Table continued on next page

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Table 12. "C" error codes (cont'd)

Error code	Description	Recommended action
C-10	Unexpected Result Time	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.
C-11	eStick Timeout	Run test within 60 minutes of removing eStick from foil. Discard and use new eStick.
C-12	Test Removed Too Early	Wait for test completion before removing eStick Discard and use new eStick.
C-13	Connection Error	Remove and re-insert the eStick If error persists, discard and use new eStick. If error persists with new eStick, discontinue use of eHub port.
C-14	eHub Low Battery	Charge eHub or connect to main power prior to repeating test. Remove and re-insert the eStick If error persists, discard and use new eStick.
C-15	eHub Internal Error	The eHub can no longer be used. Contact QIAGEN Customer Support.
C-16	eHub RTC Failure	The eHub can no longer be used. Contact QIAGEN Customer Support.

* The eHub does not have to be fully charged before running a test, but we recommend keeping the eHub plugged in to a power source and charging at all times, if possible.

Troubleshooting Guide

This troubleshooting guide may be helpful in solving any problems that may arise. For more information, see also the Frequently Asked Questions page at our Technical Support Center: <u>www.giagen.com/FAQ/FAQList.aspx</u>. The scientists in QIAGEN Technical Services are always happy to answer any questions you may have about either the information or protocols in this handbook (for contact information, see back cover or visit www.giagen.com).

QIAreach Anti-SARS-CoV-2 Total Test troubleshooting

See Table 9-12 in Technical Information section for the list of QIAreach Anti-SARS-CoV-2 Total Test error codes.

Significantly hemolyzed (reddish brown) samples can potentially interfere with the QIAreach[™] Anti-SARS-CoV-2 Total Test optical measurement system. The eStick firmware features built-in controls to determine unacceptably high levels of hemolysate and will return an invalid result in the form of an error code if interference is present. If a reddish-brown test sample results in a "B" error code or if the sample is added to the eStick and the test will not start within 1 minute, then the sample may contain elevated levels of hemoglobin that interfere with the test. Causes of in vitro hemolysis may include improper sample collection from the patient and improper storage / handling of the sample prior to analysis. See Preparing samples for guidelines and for pre-analytical specimen stability and follow the sample collection device manufacturer's recommendations for sample collection and centrifugation.

Additional user warnings

When cleaning, avoid any deliberate water ingress deep into the test ports. The eHub can be cleaned using mild detergent, 10% bleach or 70% EtOH.

Only use the eHub with the USB cable and USB adapter supplied with the device.

Contact Information

For technical assistance and more information, please see our Technical Support Center at <u>www.qiagen.com/support</u>, call 800-344-3631, email <u>techservice-na@qiagen.com</u>, or contact one of the QIAGEN Technical Service Departments or local distributors (see back cover or visit www.qiagen.com).

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Symbols

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The following symbols may appear on the packaging and labelling:

Symbol	Symbol definition
Σ	Use by date
\	Temperature limitation
REF	Catalog number
MAT	Material number
	Manufacturer
業	Protect from light
	Consult instructions for use
	Caution
	Do not open electrical unit
2	Do not reuse

Ordering Information

Product	Contents	Cat. no.
QIAreach™ Anti-SARS-CoV-2 Total Test Kit	60 QIAreach eSticks / Processing Tubes 3 x 10 ml QIAreach Diluent Buffer	645033
Related Products		
QIAreach™ eHub	QIAreach™ eHub, power adapter and USB connector cable	9003063
QIAreach™ Software	n/a	1118894
QIAreach® Anti-SARS-CoV-2 Controls	QIAreach Anti-SARS-CoV-2 Positive Control and the QIAreach Anti-SARS- CoV-2 Negative Control	647030

For up-to-date licensing and product-specific disclaimers, see the respective QIAGEN kit handbook or user manual. QIAGEN kit handbooks and user manuals are available at www.qiagen.com or can be requested from QIAGEN Technical Services or your local distributor.

Document Revision History

Date	Changes
R1, August 2020	Initial release
R2, December 2020	Rebranding from Access to QIAreach™ Updated clinical agreement section for sensitivity, specificity and predictive value Updated available sample matrices study Updated Error Codes Table
R3, May 2021	Updated to include QIAreach® Anti-SARS-CoV-2 Controls. Corrected eHub catalog number

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May 2021

QIAreach[™] eHub User Manual

For *in vitro* diagnostic use For Emergency Use Authorization Only Rx Only





Sample to Insight

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1 Introduction

This manual describes how to operate the QIAreach[™] eHub (also referred to as eHub herein). Before using the QIAreach[™] eHub, it is essential that you read this user manual carefully and pay particular attention to the safety information. The instructions and safety information in the user manual must be followed to ensure safe operation of the instrument and to maintain the instrument in a safe condition.

1.1 About this user manual

This user manual provides information about QIAreach™ eHub in the following sections:

- Introduction
- Safety Information
- General Description
- Installation Procedures
- Operation
- System Functions
- Maintenance
- Troubleshooting
- Technical Specifications

The Appendix section contains the following information:

- Declaration of Conformity
- Waste Electrical and Electronic Equipment (WEEE)
- Disclaimer of warranties

1.2 General Information

- This test has not been FDA cleared or approved but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C 263a, that meet requirements to perform moderate or high complexity tests.
- This test has been authorized only for detecting the presence of total antibodies to SARS-CoV-2, not for any other viruses or pathogens.
- The emergency use of this test 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.

1.2.1 Technical assistance

At QIAGEN®, we pride ourselves on the quality and availability of our technical support. Our Technical Services Departments are staffed by experienced scientists with extensive practical and theoretical expertise in molecular biology and the use of QIAGEN products. If you have any questions or experience any difficulties regarding the QIAreach™ eHub or QIAGEN products in general, do not hesitate to contact us.

For technical assistance and more information, please see our Technical Support Center at www.qiagen.com/support/technical-support or call one of the QIAGEN Technical Service Departments or local distributors (see back cover or visit www.qiagen.com).

When contacting QIAGEN Technical Services about errors, please have the following information ready:

- QIAreach™ eHub serial number
- Test type and test kit lot number
- Error code (if applicable)
- Timestamp when the error occurred for the first time
- Frequency of error occurrence (i.e., intermittent or persistent error)
- Photo of error, if possible

1.2.2 Policy statement

It is the policy of QIAGEN to improve products as new techniques and components become available. QIAGEN reserves the right to change specifications at any time.

In an effort to produce useful and appropriate documentation, we appreciate your comments on this user manual. Please contact QIAGEN Technical Services.

1.3 Intended use of the QIAreach[™] eHub

The QIAreach[™] eHub is intended for use in conjunction with QIAreach[™] Anti-SARS-CoV-2 Total Test. QIAreach[™] technology on the digital detection eStick provides diagnostic results that are displayed to the user on the QIAreach[™] eHub visual display.

The QIAreach[™] eHub and QIAreach[™] Anti-SARS-CoV-2 Total test are intended for professional use only and not intended for self-testing. The QIAreach[™] Anti-SARS-CoV-2 Total test is for in vitro diagnostic use, for Emergency Use Authorization Only, and for prescription use only.

1.3.1 Limitations of use

- The QIAreach[™] eHub can only be used with eSticks according to the instructions contained in this user manual and in the Instructions for Use for the QIAreach[™] Anti-SARS-CoV-2 Total Test.
- When powering the QIAreach[™] eHub or connecting to a computer, use only the USB cable supplied with this product.
- When charging the QIAreach[™] eHub, use only the USB charger and USB cable supplied with this product.
- The QIAreach[™] eHub should only be operated on a flat, horizontal surface with no angles or tilts.
- Do not re-run an eStick if it has already been used successfully or if it has been associated with an error or an incomplete run.
- Ensure that the QIAreach[™] eHub is positioned away from any air conditioning outlets, heaters, or sources of intense light.
- Do not move the QIAreach[™] eHub while a test is running.
- Do not remove an eStick from the QIAreach™ eHub before the run has completed.

1.4 Symbols on the QIAreach[™] eHub

The following symbols may appear on the packaging or labelling.

Symbol	Location	Description
CE	Type plate and outer box label of the QIAreach™ eHub	CE mark
F©	Type plate on the bottom of the QIAreach™ eHub	FCC Mark
X	Type plate and outer box label of the QIAreach™ eHub	WEEE Mark for Europe
	Type plate and outer box label of the QIAreach™ eHub	Legal Manufacturer
REF	Type plate and outer box label of the QIAreach™ eHub	Catalog Number
SN	Type plate and outer box label of the QIAreach™ eHub	Serial Number
GTIN	Type plate and outer box label of the QIAreach™eHub	Global Trade Item Number
Ţ	Outer box label of the QIAreach™ eHub Fragile	
	Type plate and outer box label of the QIAreach™ eHub	Electrical Safety Regulatory Compliance Mark (Australia and New Zealand)
Ĩ	Type plate and outer box label of the QIAreach™ eHub	Consult Instructions for Use

	Type plate and outer box label of the Attention	
<u> </u>	QIAreach™ eHub	

2 Safety Information

Before using the QIAreach[™] eHub, it is essential that you read this user manual carefully and pay particular attention to the safety information. The instructions and safety information in the user manual must be followed to ensure safe operation of the QIAreach[™] eHub and to maintain a safe working condition.

Possible hazards that could harm the user or result in damage to the instrument are clearly stated at the appropriate places throughout this user manual.

If the QIAreach[™] eHub is used in a manner not specified by the manufacturer, the protection provided by the equipment may be impaired.

The following types of safety information appear throughout the *QlAreach™ eHub User Manual*.

The term WARNING is used to inform you about situations that could result in personal injury to you or others.
Details about these circumstances are given in a box like this one.

The term CAUTION is used to inform you about situations that could result in damage to the QIAreach™ eHub or to other equipment.
Details about these circumstances are given in a box like this one.

Important	The term Important is used to highlight information that is critical for the completion of a task or optimal performance of the system.
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	The term Note is used for information that explains or clarifies a specific case or task.
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The guidance provided in this manual is intended to supplement, not supersede, the normal safety requirements prevailing in the user's country.

2.1 Proper use

- Use the QIAreach[™] eHub according to this user manual. We recommend you carefully read and become acquainted with the Instructions for Use before running a QIAreach[™] Anti-SARS-CoV-2 Total Test.
- Improper use of the QIAreach[™] eHub may cause personal injuries or damage to the QIAreach[™] eHub.
- The QIAreach[™] eHub must only be operated by qualified and appropriately trained QIAGEN personnel.

2.2 Electrical safety

Observe all general safety precautions that apply to electrical instruments. This device has been tested for compliance with electrical safety requirements as per IEC 61010-1: Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 1: General requirements.



Do not open the QIAreach[™] eHub. No user-serviceable parts inside. (W1) Opening of the QIAreach[™] eHub device could lead to user injury or damage of the device.

2.3 Chemical safety

Safety Data Sheets (SDSs) for QIAreach[™] Anti-SARS-CoV-2 Total Test kit materials are available and can be requested from QIAGEN. Used eSticks should be disposed of in accordance with all national, state, and local health and safety regulations and laws.

2.4 Biological safety

Samples tested on the QIAreach[™] eHub may contain infectious agents. Users should be aware of the health hazard presented by such agents and should use, store and dispose of such samples according to the required safety regulations. Wear personal protective equipment when handling reagents or samples, and wash hands thoroughly thereafter.

Always observe safety precautions as outlined in relevant guidelines. Avoid contamination of the QIAreach[™] eHub and workspace by handling samples and eSticks with care. In the event of contamination, clean and decontaminate the affected area of the QIAreach[™] eHub.

For instructions on cleaning and decontaminating the QIAreach[™] eHub, see Maintenance.

2.5 Waste disposal

For disposal of waste electrical and electronic equipment (WEEE), see Waste Electrical and Electronic Equipment (WEEE).

3 General Description

3.1 System description

The QIAreach[™] eHub, in combination with eSticks, provides a simplified workflow for multiple in vitro diagnostic tests, using state-of-the-art nanoparticle fluorescence detection technology. The eSticks are single-use tests that include a lateral flow strip and optoelectronics that perform test measurements and interpret results. The QIAreach[™] eHub provides power to the eStick to run the test when the eStick is connected to any one of the eHub-eStick ports. The QIAreach[™] eHub visually communicates test progress and results to the user via a display screen specific to each QIAreach[™] eHub port.

Optional QIAreach software is not provided with the QIAreach[™] eHub and can be purchased separately from Qiagen (Catalog # 1118894). The QIAreach[™] eHub will transmit test information and results when connected to a computer running QIAreach[™] software.

3.2 QIAreach[™] eHub description

The QIAreach[™] eHub is a connection hub that provides power to perform multiple QIAreach[™] Anti-SARS-CoV-2 Total Tests simultaneously. The QIAreach[™] eHub is connected to a power source using the provided connection cable and features a rechargeable lithium battery to allow QIAreach[™] Anti-SARS-CoV-2 Total Test to be performed when a continuous power supply is not available. QIAreach[™] Anti-SARS-CoV-2 Total Test results are interpreted on the eStick firmware, and results are transmitted to the QIAreach[™] eHub which then communicates to the user by means of a visual display.

The QIAreach[™] eHub USB charger and USB cable allow the QIAreach[™] eHub to be powered from either an electrical outlet or from a computer equipped with USB ports. Use of the optional QIAreach[™] software (Catalogue# 1118894) requires the QIAreach[™] eHub to be connected to a computer. For instructions on how to use the software, refer to the QIAreach[™] software user guide (available separately),

The QIAreach[™] eHub includes the following elements:

- eStick connection ports for up to eight (8) separate tests
- Visual display screen centered above each individual eStick connection port
- QIAreach processing tube holder slot positioned directly behind each visual display screen

- Sample tube holder slot positioned directly behind each processing tube holder slot, to be used for QIAreach[™] Anti-SARS-CoV-2 Total Test
- USB-C port for connection to a USB charger (supplied) or computer
- Battery LED indicator

Figures 1 and 2 show the location of various QIAreach™ eHub features.

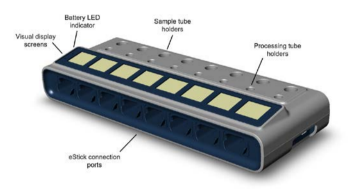


Figure 1. Front view of QIAreach™ eHub.

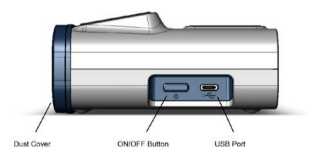


Figure 2. Side view of QIAreach™ eHub.

4 Installation Procedures

4.1 Site requirements

Select a flat, dry, and clean workbench space for the QIAreach[™] eHub. Make sure that the space is free of excessive moisture and dust, protected from direct sunlight, large temperature fluctuations, heat sources, vibration and electrical interference. Refer to Section 9 for the weight and dimensions of the QIAreach[™] eHub and the correct operating conditions (temperature and humidity). There should be sufficient clearance on the workbench to allow unimpeded access to the eStick connection ports, USB port, and ON/OFF button.

Note: Before installing and using the QIAreach[™] eHub, see Operating the QIAreach[™] eHub to become familiar with the QIAreach[™] eHub operating conditions.

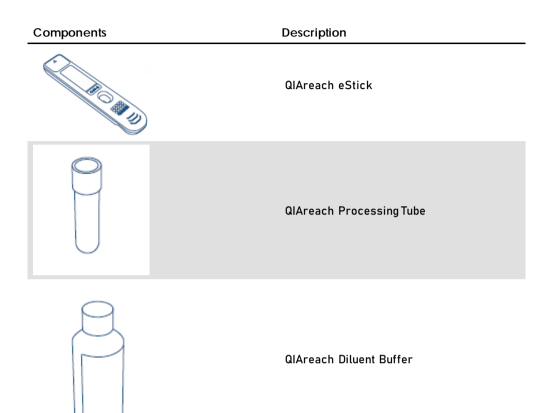
4.2 QIAreach[™] eHub delivery and components

The QIAreach[™] eHub is delivered in a single box and includes all the necessary components for setting up and connecting the QIAreach[™] eHub. The contents of the box are described below:

Note: QIAreach[™] Anti-SARS-CoV-2 Total Test kits (sold separately) are required to perform testing on QIAreach[™] eHub devices.

Components	Description
ALL DE LE CONTRACTOR	1x QIAreach™ eHub
	1x Dust cover
	1x USB drive containing QIAreach™ software
	1x USB-C – USB-A Cable, 1.5m length
	1x USB Charger Power adapter with region specific plugs

The following components are required for testing but are provided separately in the QlAreach[™] Anti-SARS-CoV-2 Total Test kit (Cat#645033).



4.3 Unpacking and installing the QIAreach[™] eHub

The QIAreach[™] eHub is delivered ready for use and does not require any hands-on assembly procedures.

Remove the QIAreach[™] eHub from its delivery box and place on a flat, level surface.

The QIAreach[™] eHub should be charged prior to use. To charge the QIAreach[™] eHub, connect the USB cable from the QIAreach[™] eHub USB port to either the supplied power adapter or to a computer.

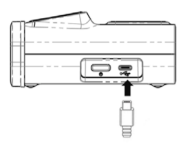


Figure 3. Connecting the USB cable to the QIAreach™ eHub USB port.

Note: The QIAreach[™] eHub will charge more quickly when charged through the supplied USB charging adapter, compared to when it is charged through a computer USB port.

The QIAreach[™] eHub comes with a dust cover to protect the internal ports from dust buildup and contamination. The cover should always be placed over the front panel of the QIAreach[™] eHub when not in use. When ready to operate the QIAreach[™] eHub, the dust cover can be removed from the front panel and set aside.

5 Operating the QIAreach[™] eHub

5.1 Setting up the QIAreach[™] eHub for use

1. Remove the dust cover from the front panel of the QIAreach™ eHub and set aside.

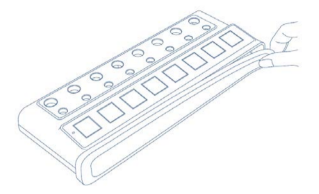


Figure 4. Removing the dust cover from the front of QIAreach™ eHub prior to use.

2. Ensure that the provided USB cable is connected to the QIAreach[™] eHub and an adequate power source (power outlet or computer).

Note: We recommend to fully charge the QIAreach[™] eHub in a switched off state overnight (when not in use). We also recommend that you connect the QIAreach[™] eHub to a USB power source (either a USB adapter or computer) during operation. If testing is performed without the use of associated QIAreach[™] software, then we recommend connecting the QIAreach[™] eHub to a power outlet (if available) through the provided USB power adapter and USB cable.

3. To turn on the QIAreach[™] eHub, press ON/OFF on the side of the unit until the visual display screen lights up.

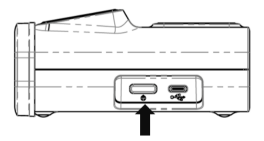


Figure 5. Press the ON/OFF switch to turn on the QIA reach[™] eHub.

4. Check battery LED indicator to ensure that the QIAreach[™] eHub has sufficient charge for the test operation. For different battery LED indicator states, see Battery LED indicator.

Once the QIAreach[™] eHub is ready for use, the visual display screen above each available eStick connection port will display the "Insert eStick" icon shown below.



Figure 6. Insert eStick icon. This signifies that an QIAreach™ eHub port is available for use.

5.2 Running a test on the QIAreach[™] eHub

Important: The steps described in this section are general test workflow cues provided on the QIAreach[™] eHub visual display screen. Refer to the QIAreach[™] Anti-SARS-CoV-2 Total Test Instructions for Use for instructions on performing a test with the QIAreach[™] eHub.

Note: Each of the connection ports on the QIAreach[™] eHub operates separately. Up to eight (8) QIAreach[™] Anti-SARS-CoV-2 Total tests can be run simultaneously.

1. When an eStick has been inserted in an eHub-eStick connection port, the self-test icon will be displayed while the eStick performs a self-test.



Figure 7. Self-test screen display.

2. Once the eStick self-test has successfully completed, the "Add sample" icon will be displayed, signifying the eStick is ready for sample addition.



Figure 8. Add sample screen display.

3. After the test sample has been added to the eStick, the "Processing" icon will be displayed along with a test countdown timer.

Important: The eStick must not be removed from the QIAreach[™] eHub until the test has been completed.



Figure 9. Processing screen display and countdown timer.

4. Once the QIAreach[™] Anti-SARS-CoV-2 Total Test has completed, the test result will be displayed on the screen and the eStick can be safely removed.



Figure 10. Test result screen.

5. If an error occurs during the test, the error icon will appear along with a specific error code. Refer to the Troubleshooting section of this manual for more information.



Figure 11. Error icon and associated error code.

5.3 Shutting down the QIAreach[™] eHub

After use, the QIAreach[™] eHub should be turned off by pressing the ON/OFF button. The QIAreach[™] eHub battery will continue to charge when turned off if connected to a power source.

After each use, clean the QIAreach[™] eHub according to the instructions in Cleaning the QIAreach[™] eHub after use.

Replace the dust cover on the front panel of the QIAreach[™] eHub to protect the eStick connection ports from moisture and dust.

6 QIAreach[™] eHub Functions

6.1 Display screen icons

Table 1. Display screen icons

Icon	ID	Description
	Please Insert	The QIAreach™ eHub port is available for eStick use.
7. No. 10. 10. 10. 10. 10. 10. 10. 10. 10. 10	Self-test	The eStick has been inserted and a self-test is being performed.
•	Add sample	The eStick is ready for sample addition to the detection port. The sample must be added within 60 minutes of removing the eStick from the foil packaging.
1959 1959	Processing	The eStick has detected sample and is processing the test. A test countdown timer is displayed. Do not remove the eStick until a result is displayed. Test times may vary across Access products.
12:34	Positive (varies with test)	The test has returned a positive result.
Θ	Negative (varies with test)	The test has returned a negative result.
X-123	Error	The test has encountered an error. The letter denotes the type and the numbers are code for the error. Refer to the Troubleshooting section for more information.

6.2 Battery LED indicator

If not connected to a power source, the QIAreach[™] eHub should have sufficient battery power to complete all in-progress QIAreach[™] Anti-SARS-CoV-2 Total Tests. A fully charged QIAreach[™] eHub should maintain internal battery power for at least 8 hours. QIAreach[™] Anti-SARS-CoV-2 Total tests should not be performed if the battery power is less than 10% and is not connected to a power source. The battery level can be checked by connecting the QIAreach[™] eHub to a computer through the provided USB cable and launching the software. The software displays the level of battery charge in the bottom right hand corner of the screen. The battery level is also indicated by the various battery LED states listed below.

Table 2. Battey levels

Display	LED state	Meaning
None	off	The QIAreach™ eHub is off
*	Flashing green	Battery charging in progress
	Solid green	The QIAreach™ eHub is turned on, battery charge > 50%
•	Solid amber	The QIAreach™ eHub is turned on, battery charge 10 - 50%
	Solid red	The QIAreach™ eHub is turned on, battery charge < 10%
*	Flashing red	The QIAreach™ eHub is turned on, battery fault

7 Maintenance

The QIAreach™ eHub does not require any service maintenance or calibration.

7.1 Cleaning the QIAreach[™] eHub after use



Risk of personal injury and material damage(W2)Disconnect the QIAreach™ eHub from all power sources before
cleaning.(C3)Ensure the QIAreach™ eHub is turned off before cleaning.

CAUTION	Risk of damage to the QIAreach™ eHub	(C4)
	When cleaning, avoid any deliberate water ingress into the eStick connection ports.	

CAUTION	Risk of damage to the QIAreach™ eHub	(C5)
	Avoid the use of excessive volumes of liquid that could enter the interior of the unit when cleaning the QIAreach™ eHub.	

Only use the following materials to clean the QIAreach™ eHub exterior surfaces:

- Mild detergent
- Water

When cleaning the QIAreach[™] eHub surface:

- Wear laboratory gloves, coat, and protective glasses.
- Wet a paper towel in mild detergent and wipe down the QIAreach[™] eHub surface and the surrounding workbench area. Take care not to intentionally wet the eStick connection ports or ON/OFF button and USB port.
- Dry the QIAreach™ eHub surface with a fresh paper towel.

8 Troubleshooting

8.1 General information

This section provides information on some issues that may occur with the QIAreach[™] eHub along with possible causes and solutions. Specific information may vary with QIAreach[™] Anti-SARS-CoV-2 Total Tests. For troubleshooting relevant to QIAreach[™] Anti-SARS-CoV-2 Total Test, see the kit instructions for use.

8.2 Contacting QIAGEN Technical Services

When contacting QIAGEN Technical Services about an error with the QIAreach[™] eHub, note the steps leading up to the error. This information will help QIAGEN Technical Services solve the problem.

When contacting QIAGEN Technical Services about errors, please have the following information ready:

- QIAreach™ eHub serial number
- Test type and test kit lot number
- Error code (if applicable)
- Timepoint when the error occurred for the first time
- Frequency of error occurrence (i.e., intermittent or persistent error)
- Photo of error, if possible

8.3 QIAreach[™] eHub error codes

If the QIAreach[™] eHub displays an error code, refer to the table below specific error descriptions and solutions.

Table 2. QIAreach™ eHub error codes

Error type	Format	Description
Self-Test	A-[Error code]	eStick electronic failure
Algorithm	B-[Error code]	Run error or user workflow error
Communication/ Other	C-[Error code]	Invalid data or missed communication betweeneStick and eHub
A-[all error codes]	eStick electronic failure	Discard and use new eStick.
		Repeat test.
B-8	Sample added too early	Only add sample after "Add Sample" icon is displayed.
		Repeat test.
B-14	No sample detected	Add sample within 60 min of insertion.
		Repeat test.
B-16	Baseline signal at reference line too low	Ensure solution in processing tube is well mixed.
		Repeat test.
B-19	Baseline signal at test line too low	Ensure solution in processing tube is well mixed.
		Repeat test.
B-255	eStick removed too early from eHub	Remove eStick only after a valid test result or error code is displayed.
B-[all other error codes]	Error code specific	Repeat test.
C-1	Out of date, i.e. expired test	Discard and use another eStick from a kit within expiry.
C-14	eHub battery charge too low	Connect eHub to a power outlet or computer via USB cable*. Remove and reinsert eStick.
C-[all other error codes]	Electrical contact or timing issue	Remove and reinsert eStick. If issue persists, discard and use new eStick.

The QIAreach™ eHub does not have to be fully charged before running a test, but we recommend to keep the QIAreach™ eHub plugged in to a power source and charging at all times, if possible.

9 Technical Specifications

Dimensions and weight

Dimensions	Width: 304 x 112 x 51 mm (12 x 4.4 x 2 in)
Weight	1000 g (2.2 lb)

Power requirements

USB Charger Power Adapter (supplied)			
Voltage	90-264 VAC		
Power	0.4A max		
Frequency	50-60 Hz		
Line regulation	± 1%		
Load regulation	± 5%		
Line Frequency Variation	\pm 3 Hz		
QIAreach™ eHub device:			
Voltage	5V DC		
Power	1.0A		
Internal Li-Ion battery (non u	ser-seviceable):		
 Voltage: 	3.7V nominal		

Capacity:	3350 mAh nominal
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Operating conditions

Air Temperature	15–30°C (59–86°F)
Relative Humidity	30-65% (non-condensing)
Place of Operation	For Indoor use only

Transport conditions

Air Temperature	5–50°C (41–122°F)
Relative Humidity	Maximum 70% relative humidity, non-condensing

Storage conditions

Air Temperature	15–30°C (59–86°F)
Relative Humidity	30–65% (non-condensing)

RoHS (Responsibility of Health and Safety) Compliance:

• Compliant with RoHS 3 EU Directive 2015/863: Restriction of the Use of certain Hazardous Substances in electrical and electronic equipment

9.1 Electromagnetic compatibility (EMC)

- Compliant with IEC 61326-1: Electrical equipment for measurement, control and laboratory use EMC requirements Part 1: General requirements
- Meets the requirements of CISPR 11:2015: ISM Equipment Radiated RF Emissions as a Group 1, Class A device
- Meets the requirements of FCC Title 47 CFR Part 15 Subpart B- unintentional radiators as a Class A device

9.2 Electrical Safety

• Compliant with IEC 61010-1: Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 1: General requirements

9.3 Cybersecurity

- Negligible or acceptable failure modes or hazards identified after a failure mode and effects analysis (FMEA) of the cybersecurity risks to the test system, including the optional QIAreach software.
- Standard Windows Filesharing provides authenticated and encrypted communication.

9.4 Software Validation

 Complaint with ISO 62304: Software verification of the QIAreach software has been done in accordance with the Ellume Quality System Design Control Procedures and ISO 62304 on both the 32-bit and 64-bit version.

10 Appendix A – Technical Data

10.1 FCC Compliance: Supplier's Declaration of Conformity

47 CFR § 2.1077 Compliance Information

Unique Identifier: 9002969 - QIAGEN QIAreach™ eHub

Responsible Party - U.S. Contact Information

QIAGEN Inc. - USA

19300 Germantown Road

Germantown, MD 20874

Telephone: (800-362-7737)

Email: customercare-us@qiagen.com

FCC Compliance Statement

This product has been tested and found to comply with the limits for a Class A digital device pursuant to Part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference when the equipment is operated in a commercial environment. This product generates, uses, and can radiate radio frequency energy and, if not installed and used in accordance with the manufacturer's instruction manual, may cause harmful interference with radio communications. Operation of this product in a residential area is likely to cause harmful interference, in which case you will be required to correct the interference at your own expense.

This device complies with Part 15 of the FCC Rules. Operation is subject to the following two conditions:

1) This device may not cause harmful interference.

2) This device must accept any interference received, including interference that may cause undesired operation.

Notice: The FCC regulations provide that changes or modifications not expressly approved by QIAGEN, Inc. could void your authority to operate this equipment.

These limits are designed to provide reasonable protection against harmful interference in a non-residential installation. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference with radio or television reception, which can be determined by turning the equipment off and on, you are encouraged to try to correct the interference by one or more of the following measures:

• Reorient or relocate the antenna of the radio/television receiver.

• Increase the separation between this equipment and the radio/television receiver.

• Plug the equipment into a different outlet so that the equipment and the radio/television receiver are on different power mains branch circuits.

Consult a representative of QIAGEN or an experienced radio/television technician for additional suggestions.

In order to maintain compliance with FCC regulations, this equipment must be used with the supplied USB cable and USB Charger. Operation with non-approved equipment or alternate cables may result in interference to radio and TV reception.

10.2 Waste Electrical and Electronic Equipment (WEEE)

This section provides information about disposal of waste electrical and electronic equipment by users.

The crossed-out wheeled bin symbol (see below) indicates that this product must not be disposed of with other waste; it must be taken to an approved treatment facility or to a designated collection point for recycling, according to local laws and regulations.

The separate collection and recycling of waste electronic equipment at the time of disposal helps to conserve natural resources and ensures that the product is recycled in a manner that protects human health and the environment.



Recycling can be provided by QIAGEN upon request at additional cost. In the European Union, in accordance with the specific WEEE recycling requirements and where a replacement product is being supplied by QIAGEN, free recycling of its WEEE-marked electronic equipment is provided.

To recycle electronic equipment, contact your local QIAGEN sales office for the required return form. Once the form is submitted, you will be contacted by QIAGEN either to request follow-up information for scheduling collection of the electronic waste or to provide you with an individual quote.

10.3 Disclaimer of warranties

EXCEPT AS PROVIDED IN QIAGEN TERMS AND CONDITIONS OF SALE FOR THE QIAREACH[™] EHUB, QIAGEN ASSUMES NO LIABILITY WHATSOEVER AND DISCLAIMS ANY EXPRESS OR IMPLIED WARRANTY RELATING TO THE USE OF THE QIAREACH[™] EHUB INCLUDING LIABILITY OR WARRANTIES RELATING TO MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE, OR INFRINGEMENT OF ANY PATENT, COPYRIGHT, OR OTHER INTELLECTUAL PROPERTY RIGHT ANYWHERE IN THE WORLD.

11 Ordering Information

Product	Contents	Cat. no.
QIAreach™ eHub	Includes 1 QIAreach™ eHub; 1 USB charging cable; 1 power adapter; 1 QIAreach™ eHub dust cover	9003063
Optional QIAreach Software	N/A	1118894

For up-to-date licensing information and product-specific disclaimers, see the respective QIAGEN kit handbook or user manual. QIAGEN kit handbooks and user manuals are available at www.qiagen.com or can be requested from QIAGEN Technical Services or your local distributor.

12 Document Revision History

Date	Changes
R1, July 2020	Initial release
R2, May 2021	Change to catalog Number

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Limited License Agreement for OlAreach™ eHub

Use of this product signifies the agreement of any purchaser or user of the product to the following terms:

- The product may be used solely in accordance with the protocols provided with the product and this handbook and for use with components contained in the kit only. 1 The product may be used solely in accordance with the protocols provided with the product and this handbook and for use with components contained in the kit own QIAGEN grants no license under any of its intellectual property to use or incorporate the enclosed components of this kit with any components not included within this kit except as described in the protocols provided with the product, this handbook, and additional protocols available at www.qiagencom. Some of these additional protocols have been provided by QIAGEN users for QIAGEN users. These protocols have not been thoroughly tested or optimized by QIAGEN. QIAGEN neither guarantees them nor warrants that they do not infringe the rights of third-parties.
- 2. Other than expressly stated licenses, QIAGEN makes no warranty that this kit and/or its use(s) do not infringe the rights of third-parties.
- 3. This kit and its components are licensed for one-time use and may not be reused, refurbished, or resold.
- QIAGEN specifically disclaims any other licenses, expressed or implied other than those expressly stated. 4.
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QIAreach[™] Anti-SARS-CoV-2 **Total Test**



Emergency Use Authorization (EUA) only

Please be advised:

- This test has not been FDA cleared or approved, but has been 0 authorized for emergency use by FDA under an EUA for use by authorized laboratories:
- This test has been authorized only for detecting the presence of 0 total antibodies to SARS-CoV-2, not for any other viruses or pathogens; and
- The emergency use of this test is only authorized for the duration 0 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.

Please contact QIAGEN Technical Services (1-800-426-8157) if you require a printed copy free of charge.

Please find the "QIAreach Anti-SARS-CoV-2 Total Test Instructions for Use (Handbook)", and the "QIAreach Anti-SARS-CoV-2 Total Test Quick Reference Guide" instructions for use at the following web address:

www.giagen.com/us/products/diagnostics-and-clinicalresearch/infectious-disease/giareach-solutions/giareach-antisars-cov-2-total-us/#orderinginformation

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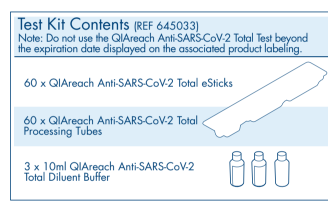
QlAreach® Anti-SARS-CoV-2 Total Test Quick Reference Guide Pre-analytical Steps

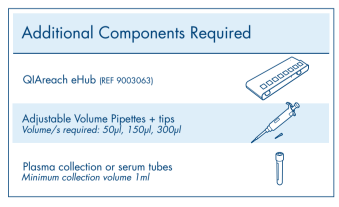
The QIAreach® Anti-SARS-CoV-2 Total Test has not been FDA cleared or approved, but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories. Testing of serum laboratories certified under CLIA that meet the requirements to perform moderate- or high-complexity tests.

The QIAreach™ Anti-SARS-CoV-2 Total Test has been authorized only for detecting the presence of total antibodies to SARS-CoV-2, not for any other viruses or pathogens.

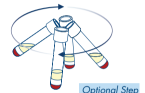
The emergency use of this test is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the declaration is terminated or authorization is revoked sooner. Due to the risk of false-positive results, confirmation of positive results should be considered using a second, different total antibody assay. Please study the *QlAreach™ Anti-SARS-CoV-2 Total Test Instructions for Use (Handbook)* thoroughly before referring to this Quick Reference Guide. This Quick Reference Guide is not intended as an exhaustive instructional document.











Draw patient sample using a plasma collection tube or standard serum tube following phlebotomy best practice.

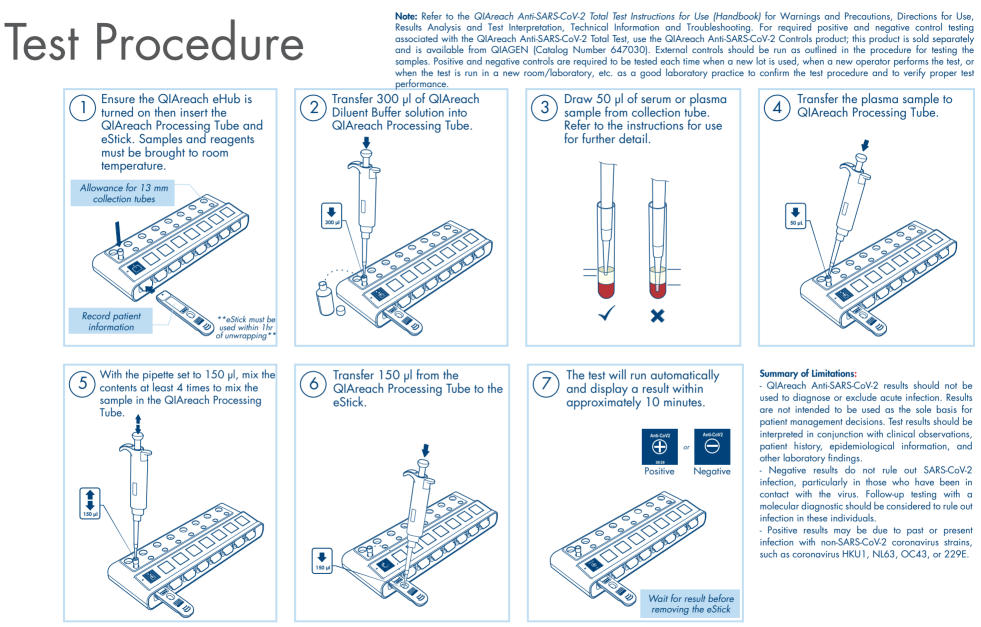
Centrifuge the sample per the collection tube manufacturer's specifications.

QIAreach eHub Setup



Remove QIAreach eHub from its packaging. Remove the dust cover from the front of the QIAreach eHub. Connect the eHub to power via USB (wall plug or PC) and turn on by pushing the power button.

NB: Refer to QIAreach *eHub User Manual* for a complete guide to device operation and troubleshooting.



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Sample to Insight

QIAreach® Anti-SARS-CoV-2 Controls Package Insert

For use under Emergency Use Authorization (EUA) only For In Vitro Diagnostic Use Rx Only

Intended use

For use with the QIAreach® Anti-SARS-CoV-2 Total Test (Cat# 645033)

The QIAreach Anti-SARS-CoV-2 Controls are intended for use as positive and negative external quality controls to assess the performance of the QIAreach Anti-SARS-CoV-2 Total Test (Cat# 645033). The QIAreach Anti-SARS-CoV-2 Controls are provided to assist laboratories with training new operators and qualifying individual lots of the QIAreach Anti-SARS-CoV-2 Total Test.

The QIAreach Anti-SARS-CoV-2 Controls are designed for quality assurance purposes, and should be used when:

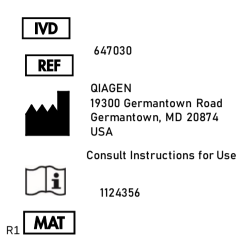
- Training new operators
- Qualifying new lots or shipments of the QIAreach Anti-SARS-CoV-2 Total Test.
- Identifying systematic issues at a particular lab or testing site.
- External controls are deemed necessary by organizational quality control procedures, and/or in accordance with Local, State and Federal regulations or accreditation requirements.

Summary and explanation

The QIAreach Anti-SARS-CoV-2 Controls test kit consists of the QIAreach Anti-SARS-CoV-2 Positive Control and the QIAreach Anti-SARS-CoV-2 Negative Control. The Negative Control contains SARS-CoV-2 seronegative human serum and the Positive Control contains SARS-CoV-2 antibodies in a human serum matrix diluted to a low reactive level.

Principles of the procedure

The QIAreach Anti-SARS-CoV-2 Controls are provided frozen in single-use vials and are substituted with the patient sample in the QIAreach Anti-SARS-CoV-2 Total Test workflow. For instructions on how to perform the test, refer to the *QIAreach Anti-SARS-CoV-2 Total Test Instructions For Use (Handbook)*.





Kit contents

QIAreach Anti-SARS-CoV-2 Controls			
Catalog no.		647030	
QIAreach Anti-SARS-CoV-2 Positive Control (100 μl)	Contains anti-SARS-CoV-2 antibodies in human serum	4 ea	
QIAreach Anti-SARS-CoV-2 Negative Control (100 μl)	Contains SARS-CoV-2 seronegative human serum	4 ea	

Store QIAreach Anti-SARS-CoV-2 Controls at -30 to -15°C. Do not use after the expiration date printed on the labelling. Each control vial is single use only and should be used within 4 hours of thawing to room temperature.

Warnings and Precautions

When working with chemicals, always wear a suitable lab coat, disposable gloves, and eye protection goggles. For more information, please consult the appropriate safety data sheets (SDSs). Refer to the *QlAreach Anti-SARS-CoV-2 Total Test Instructions For Use (Handbook)* for further safety information.

Important: Inspect vials prior to use. Do not use control vials that show signs of damage or if the vial cap has been compromised. Do not handle broken vials. Take appropriate safety precautions to dispose of vials safely.

- 1. For Emergency Use Authorization only.
- 2. For *in vitro* diagnostic use only.
- 3. For professional use only.
- This test has not been FDA cleared or approved, but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C §263a, that meet requirements to perform moderate or high complexity tests.
- 5. This test has been authorized only for detecting the presence of total antibodies to SARS-CoV-2, not for any other viruses or pathogens.
- 6. The emergency use of this test 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. § 360bbb3(b)(1), unless the declaration is terminated or authorization is revoked sooner.
- 7. All specimens of human origin should be considered potentially infectious and handled with care.

Procedure

- 1. Equilibrate all controls to room temperature (15-30°C) prior to use. Controls should be used within 4 hours of thawing. Do not refreeze vials after thawing.
- Controls are substituted with the patient sample in the QIAreach Anti-SARS-CoV-2 Total Test workflow and 50 μl of control sample should be used in the assay workflow.
- 3. Discard opened vials after use in accordance with the local and government regulations.

Expected results

The QIAreach Anti-SARS-CoV-2 Positive Control should return a Positive result and the QIAreach Anti-SARS-CoV-2 Negative Control should return a Negative result in the QIAreach Anti-SARS-CoV-2 Total Test. Control test results that do not match with the expected results may identify a potential issue with operator training, laboratory handling procedures or with a specific QIAreach Anti-SARS-CoV-2 Total Test lot or shipment. If the control results do not match the expected results should not be reported.

For up-to-date licensing information and product-specific disclaimers, see the respective QIAGEN kit handbook or user manual. QIAGEN kit handbooks and user manuals are available at www.qiagen.com or can be requested from QIAGEN Technical Services or your local distributor.

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