OPTI SARS-CoV-2 RT-PCR Test

English Version

Used for real-time PCR identification of SARS-CoV-2 RNA extracted from upper respiratory specimens (such as nasal, nasopharyngeal, oropharyngeal sw abs, and nasopharyngeal w ash/aspirate or nasal aspirate) and bronchoalveolar lavage.



IVD (E R

For *in vitro* diagnostic use only For Emergency Use Authorization Only For Prescription Use only

REF 99-57003 and 99-57004





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English version

OPTI SARS-CoV-2 RT-PCR Test

Intended Use

The OPTI SARS-CoV-2 RT-PCRTest is a real-time fluorescent reverse transcription polymerase chain reaction testfor the qualitative detection of nucleic acid from the SARS-CoV-2 in upper respiratory specimens (such as nasal swabs, nasopharyngeal swabs, or or pharyngeal swabs, and nasopharyngeal wash/aspirate or nasal as pirate) and bronchoalveolar lavage from patients suspected of COVID-19 by their healthcare provider. 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 high complexity tests.

Results are for the identification of SARS-CoV-2 RNA. The SARS-CoV-2 RNA is generally detectable in respiratory samples during the acute phase of infection. Positive results are indicative of the presence of SARS-CoV-2 RNA; clinical correlation with patient history and other diagnostic information is necessary to determine patient infection status. Positive results do not rule out bacterial infection or co-infection with other viruses. The agent detected may not be the definite cause of disease. Laboratories within the United States and its territories are required to report all test results to the appropriate public health authorities.

Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for patient management decisions. Negative results must be combined with clinical observations, patient history, and epidemiological information.

The OPTI SARS-CoV-2 RT-PCR Test is intended to be used by qualified clinical laboratory personnel specifically instructed and trained in the techniques of real-time PCR and *in vitro* diagnostic procedures.

The OPTI SARS-CoV-2 RT-PCR Test is only for use under the Food and Drug Administration's Emergency Use Authorization.

Product Description

The OPTI SARS-CoV-2 RT-PCR Test is a real-time reverse transcription polymerase chain reaction (rRT-PCR) test that uses the N1 and N2 primer and probe sequences which are described by the CDC design¹. The OPTI SARS-CoV-2 RNA Mix (SARS-CoV-2 Mix) includes primers and probes for the detection of SARS-CoV-2 RNA when amplified with the OPTI RNA Master Mix (RNA MMx). SARS-CoV-2 RNA targets (N1 and N2) are both detected in the FAM channel. The internal control for the test is RNase P (RP), which is detected in the HEX channel. The internal control for the test is based on the detection of a conserved nucleic acid sequence present in human samples. This host target is referred to as the internal sample control (ISC). Detection of endogenous nucleic acid in the test sample controls for sample addition, extraction, and amplification. Primers and probe for detection of the internal sample control are included in the SARS-CoV-2 Mix.

During the real-time reverse transcription polymerase chain reaction, viral RNA is reverse transcribed into cDNA and subsequently amplified in a real-time PCR cycling protocol. During the process, the probe anneals to a specific target sequence located between the forward and reverse primers. During the extension phase of the PCR cycle, the 5' nuclease activity of Taq polymerase degrades the probe, causing

the reporter dye to separate from the quencher dye, generating a fluorescent signal. With each cycle, additional reporter dye molecules are cleaved from their respective probes, increasing the fluorescence intensity exponentially. Fluorescence intensity is monitored at each PCR cycle by one of the PCR thermal cycler instruments listed in Section "Materials Required but Not Provided".

In addition, the OPTI SARS-CoV-2 RT-PCR Test utilizes the OPTI Positive Control (PC) and OPTI PCR Grade Water (Negative Control). The OPTI Positive Control (PC) contains SARS-CoV-2 and ISC synthetic material and works as a positive control for the reaction. OPTI PCR Grade Water is used as the PCR negative control, as well as to reconstitute the dried SARS-CoV-2 Mix and the PC.

"Coronavirus Disease 2019 (COVID-19) Real-Time rRT-PCR Panel Primers and Probes." Centers for Disease Control
and Prevention, Centers for Disease Control and Prevention, 6 Mar. 2020, www.cdc.gov/coronavirus/2019-ncov/lab
/rt-pcr-panel-primer-probes.html.

Materials and Storage

		Quantity		tity	Storage			
	Identification/ General Information	Cap color	100 tests 99-57003	500 tests 99-57004	At receipt	After reconstitution	Freeze/ Thaw cycles	
	OPTI SARS-CoV-2 Mix (SARS-CoV-2 Mix), dried	Red	1 x 1.0 mL	5 x 1.0 mL	–25 to 8°C	−25 to −15°C	≤6	

61-56616-00

Contains N1, N2 and ISC primers and probes. Reconstitute to 1 mL in PCR Grade Water. Store the SARS-CoV-2 Mixin the dark. The expiration date on the vial is valid for either the dry or reconstituted form.

OPTI RNA Master Mix (RNA MMx)	Black	1 x 1.0 mL	5 x 1.0 mL	−25 to −15°C (Long-term)	N/A	≤6
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61-56618-00

Concentrated master mix that includes reverse transcriptase and hot-start polymerase. The RNA MMx is more viscous than most master mixes— see the Test Procedure section for handling recommendations. A reference dye (ROX) has been added for normalizing volume inaccuracies. Protect the RNA MMx from light.

OPTI Positive Control, dried (PC)	Blue	1 x 200 μL	1 x 200 μL	–25 to 8°C	−25 to −15°C	≤6
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The PCcontains the targets for SARS-CoV-2 (N1 target region) and the internal control (RNase P). Reconstitute to 200 µL in PCR Grade Water. The expiration date on the vial is valid for either the dry or reconstituted form.

OPTI PCR Grade Water	Clear	2 x 1.0 mL	7 x 1.0 mL	−25 to 8°C	N/A
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61-56619-00

44-56617-00

PCR Grade Water has been qualified for reverse transcription-PCR (RT-PCR) use. It is used for the reconstitution of the SARS-CoV-2 Mix and PC. It is also used as the negative control for PCR. Do not transport PCR Grade Water vials between PCR work areas. Separate vials of water are needed for each area to avoid contamination risk.

Note: See table at the end of the insert for a description of symbols used on the insert and labels.

Materials Required but Not Provided

Real-Time PCR Instrument and consumables	Source and part number
Thermo Scientific	
Applied Biosystems® 7500 FAST	7500 instrument (4351106) and 7500software v2.0.6
Applied Biosystems® QuantStudio 5	QS5 instrument (A28138) and QuantStudio Design
96 well PCR plate	and Analysis Desktop software (v1.5.1)
Optical plate cover	plate: 4346906 cover: 4311971
Agilent	
Agilent Mx3005P™	3005P instrument (401449) and MxPro qPCR software v4.10
96 well PCR plate	plate: 401334
Optical cap strips	caps: 401425
IDEXX Laboratories	
Bio Molecular Systems Mic qPCR Tubes and caps	Instrument (98-0012758-00) and micPCR software v2.8.10 tubes + caps: 98-0012759-01
Roche	
Roche LightCycler® 480 96 well PCR plate + cover	Instrument (05015278001) and LightCycler 480 SW v1.5.1 plate + cover: 04 729 692 081
Extraction Equipment and Consumables	Source and part number
RealPCR DNA/RN A Magnetic Bead Kit	IDEXX 99-56102 (384 samples) / 99-56106 (96 samples)
Nucleo Mag VET Magnetic Extraction Kit	Macherey Nagel 744200.4
OPTI RNA/DNA Magnetic Bead Kit PurePrep Pathogens Extraction Kit	OPTI Medical Systems 99-58015 MolGen BV OE00290096 (n=96) and OE00290960 (n=960)
Turcing Turnogens Extraction int	Worden By October (11-30) and October 300 (11-300)
Thermo Scientific	[[]
Thermo Scientific™ KingFisher™ Flex 96 deep well plate	Flex instrument (5400630) and software v1.0.1.0 Deep well plate: 95040460
96 well elution plate	Elution plate: 97002540
96 tip comb for deep well magnet	Tip Comb: 97002534
Thermo Scientific™ KingFisher™ Duo Prime 96 deep well plate 12-tip elution strip for deep well plate 12-tip comb for deep well plate	Duo instrument (5400110) and software v1.02.27. RT18 Deep well plate: 95040460 Elution strip: 97003520 Tip comb: 97003500
Manual Magnetic Extraction (OPTI)	Manual Extraction (OPTI RNA/DNA Magnetic Bead Kit)
Magnetic separator for 96 well plates	MLS
Plate shaker/heater	MLS
Multichannel pipette	MLS
Extraction control containing human specimen (HSC) material	See Quality Controls section

Equipment and Lab Consumables	Source and part number
Molecular grade water (used as extraction negative control)	MLS
Micro-centrifuge for 2 mL microtubes capable of 1500–3000 x g	MLS
Vortex mixer	MLS
1.5 mL microcentrifuge tubes (DNase/ RNase free)	MLS
Pipettes and multi-channel pipettes (5–1000 μL); dedicated pipettes for preparation of PCR Mix	MLS
Nuclease-free, aerosol resistant pipette tips	MLS
Personal protective equipment consistent with current guidelines for handling infectious samples	MLS
Optional: Centrifuge with rotor and adapters for multi-well plates	MLS

MLS = Major Laboratory Supplier, such as vwr.com or fisherscientific.com

Warnings and Precautions

General

- The assay is for in vitro diagnostic (IVD) use under the FDA Emergency Use Authorization Only.
- For prescription use only.
- This product has not been FDA cleared or approved, but has been authorized for emergency
 use by FDA under an Emergency Use Authorization (EUA) for use by authorized laboratories.
- This product has been authorized only for the detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens.
- The emergency use of this product is only authorized for the duration of the declaration that
 circumstances exist justifying the authorization of emergency use of in vitro diagnostics tests
 for detection and/ordiagnosis 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.
- Handle all specimens as of infectious using safe laboratory procedures. Refer to Interim Laboratory Biosafety Guidelines for Handling and Processing Specimens Associated with SARS-CoV-2: https://www.cdc.gov/coronavirus/2019-nCoV/lab-biosafetyguidelines.html.
- Use personal protective equipment (PPE) consistent with current guidelines for the handling of potentially infectious samples.
- Do not eat, drink, smoke, apply cosmetics or handle contact lenses in areas where reagents and human specimens are handled.
- Modifications to assay reagents, assay protocol, or instrumentation are not permitted, and are in violation of the product Emergency Use Authorization.
- Dispose of waste in compliance with the local, state, and federal regulations.
- Laboratories within the United States and its territories are required to report all test results to the appropriate public health authorities.

PCR

- Reagents must be stored and handled as specified in these instructions for use. Do not use reagents
 past expiration date.
- The entire procedure must be performed under nuclease-free conditions.
- Wear powder-free gloves when working with the reagents and nucleic acids.
- Always use pipette tips with aerosol barriers. Tips that are used must be sterile and free from DNases and RNases.
- Keep reagents and PCR Mixtubes capped or covered as much as possible.
- To avoid cross-contamination, use nuclease-free, aerosol-resistant pipette tips for all pipetting, and physically separate the workplaces for nucleic acid extraction/handling, PCRs etup and PCR.
- Work surfaces, pipettes, and centrifuges should be cleaned and decontaminated with cleaning products such as 10%bleach, "DNAZap™" or "RNaseAWAY" to minimize riskofnucleicacid contamination. Residual bleach should be removed using 70% ethanol.
- The internal control for the test detects human nucleic acid; it is important to avoid environmental sources of human nucleic acid contamination.

Specimen Collection

- The Sample collection device is not a part of the test kit. The OPTI SARS-CoV-2 RT PCR Test is compatible with FDA recommended swabs and transport media. Refer to Interim Guidelines for Collecting, Handling, and Testing Clinical Specimens from Patients Under Investigation (PUIs) for 2019 Novel Coronavirus (2019-nCoV):
 - https://www.cdc.gov/coronavirus/2019-nCoV/guidelines-clinical-specimens.html
- Follow specimen collection manufacturer instructions for proper collection methods.
- Swab specimens should be collected using only swabs with a synthetic tip, such as nylon or Dacron® and an aluminum or plastic shaft. Calcium alginate swabs should not be used and cotton swabs with wooden shafts are not recommended. Place swabs immediately into sterile tubes containing 2–3 mL of viral transport media.

Transporting Specimens

Specimens must be packaged, shipped, and transported according to the current edition of the International Air Transport Association (IATA) Dangerous Goods Regulation. Follow shipping regulations for UN 3373 Biological Substance, Category B when sending potential 2019-nCoV specimens. Store specimens at 2–8°C and ship on ice packs

Storing Specimens

- Specimens can be stored at 2-8°C for up to 72 hours after collection.
- If a delay in extraction is expected, store specimens at -70°C or lower.

Reconstitution of Dried Components

Reconstitute the SARS-CoV-2 Mix and Positive Control by pipetting PCR Grade Water to the volume indicated on the component label. Allow to sit at 18 to 26° C for at least 10 minutes; mix and microcentrifuge briefly prior to use. Once the SARS-CoV-2 Mix and the Positive Control are reconstituted, aliquot as appropriate and store the solutions frozen. When handling frozen components, thaw at 18 to 26° C for approximately 15 to 30 minutes, mix gently and then microcentrifuge briefly ($^{\sim}1,500-3,000\times g$).

Extraction

Magnetic Bead Extraction kits (for automated use on Thermo Scientific™ KingFisher™ Flex and Thermo Scientific™ KingFisher™ Duo Prime)

- RealPCR DNA/RNA Magnetic Bead Kit (IDEXX, Part#99-56102/ 99-56106) for use on Thermo Scientific™ KingFisher™ Flex
 - Sample input volume: 200 uL; elution volume 100 uL
- Nucleo Mag
 VET Magnetic Extraction Kit (Macherey Nagel, Part #744200.4) for use on Thermo Scientific™ KingFisher™ Duo Prime
 - Sample input volume: 200 uL; elution volume 100 uL
- OPTI DNA/RNA Magnetic Bead Kit (OPTI Medical Systems 99-58015) for use on Thermo Scientific™ KingFisher™ Flex and KingFisher™ Duo Prime instruments or Manual protocol
 - Sample input volume: 200 uL; elution volume 100 uL
- PurePrep Pathogens Extraction Kit (MolGen BV, Part # OE00290996, 96 samples and OE00290960, 960 samples) for use on Thermo Scientific™ KingFisher™ Duo Prime instrument
 - Sample input volume: 200 uL; elution volume 100 uL

Store the purified RNA at <-15°C if testing is not performed immediately after RNA extraction.

Quality Controls

Control(s) that are provided with the OPTI SARS-CoV-2 RT-PCR Test are listed below:

- PCR Negative Control (OPTI PCR Grade Water): A "no template" (negative) control is needed to confirm
 the PCR plate is valid. PCR Grade water is used and should be included for each PCR run. The negative
 control should test negative for the SARS CoV-2 target and internal control. The no template control is
 not included during extraction.
- Positive Control (OPTI Positive Control): A positive template control is needed to confirm the PCR plate is valid. Synthetic nucleic acid for the N1 target region is used at 20 copies per µL. The positive control should be included on each PCR run and should test positive for both the SARS CoV-2 target and internal control channels. The positive control is not included during extraction.
- The internal control for the test is a human endogenous nucleic acid sequence (RNase P) and controls for sample addition, extraction and PCR. The internal control is expected to test positive for each sample tested.

Control(s) that are required but not provided with the OPTI SARS-CoV-2 RT-PCR Test are listed below:

- Extraction Negative Control: A "no sample" (negative) control is needed to confirm that the extraction
 process is valid. Molecular grade water should be included as a sample for each extraction run. The
 purpose of an Extraction Negative Control is to monitor for cross-contamination of samples with
 amplifiable materials from SARS-CoV-2 and/or human sourced material (RNase P) within the extraction
 run. The Extraction Negative Control should always test negative for the SARS CoV-2 target.
- Extraction Sample control: An extraction control containing human specimen control (HSC) material should be extracted and tested with each set of patient samples. The extraction control is used to demonstrate successful recovery of RNA during the extraction process and should test negative for the SARS CoV-2 target and positive for the RNase P internal control. Laboratories may use confirmed negative human specimen material (e.g. a negative respiratory specimen). This material should be prepared in enough volume to be used across multiple runs. Material should be tested prior to use as the extraction control to ensure it generates the expected results.

Test Procedure

- 1 Preparation of the PCR Mix.
 - Mix the thawed RNA MMx by inversion or gentle vortex.
 - The RNA MMx is a viscous solution; always pipette it slowly.
 - To prepare the PCR Mix add 10 μL SARS-CoV-2 Mix and 10 μL RNA MMx for each reaction.
 - When preparing the PCR Mix, first pipette SARS-CoV-2 Mix into the tube and then add the RNA MMx.
 Pipette up and down a few times to rinse the MMx pipette tip.
 - Gently vortex the solution to ensure the components are mixed well.
 - · Pipette the PCR Mix slowly into the PCR plate.

Load the PCR plate within 20 minutes or store at 2 to 8°C for up to 4 hrs. The PCR Mix can be stored at -25 to -15°C for up to 2 weeks. Protect from light.

- 2 Pipette 20 µL of the PCR Mix into the required wells of the multiwell plate.
- 3 Add 5 μ L of sample RNA to each well. The final reaction volume is 25 μ L.
- Include 5 μ L each of the Positive Control , PCR Negative Control , Extraction Negative Control and Extraction Sample Control (5 μ L) for each test run.
- 5 Cover the plate and briefly spin the plate, if necessary, to settle contents and remove air bubbles.
- 6 Load the plate into the PCR instrument. Set up thermal cycler with Cycling Program below. Start the run.

Settings for Reporter and Quencher

<u>Targe</u> t	Reporter	Quencher
SARS-CoV-2	FAM™	BHQ [®] (none)
Internal Control (RNase P)	HEX [™] (VIC)	BHQ (none)
Passive Reference	ROX™	N/A

Cycling Program (used for all instruments)

<u>Step</u>	Temperature	<u>Time</u>	Cycles
Reverse transcription (RT)	50°C	15 min.	1
Denaturation	95°C	1 min.	1
Amplification**	95°C 60°C	15 sec. 30 sec.	45

^{**}Ensure the instrument is set to record fluorescence following the 60°C amplification step.

7 Examination and Interpretation of Quality Control Results

All test controls should be examined prior to interpretation of patient results. If the controls are not valid, the patient results cannot be interpreted.

Using the PCR instrument software, assign a unique identifier for the SARS-CoV-2 and internal control targets on the plate. To obtain appropriate Ct values, analysis for both the SARS CoV-2 target and internal control target should be performed by manually setting the threshold. Each target threshold should be set separately. The threshold should be adjusted to the inflection point for the exponential phase of the curve and above background signal. This is best done while viewing all amplification curves, for each respective target for a given run, on a logarithmic scale. It is important to follow the same procedure run to run when setting the manual threshold.

Refer to specific instrument's user manual for guidance on how to analyze data.

Plate Validity Criteria

The following control results must be obtained for each PCR run in order for the run to be deemed valid. If the plate controls are not valid, the patient results cannot be interpreted, are not valid, and the plate must be repeated.

<u>Contro</u> l	SARS-CoV-2 FAM Ct Value	SARS-CoV-2 FAM Result	HEX Ct Value	Internal Control HEX Ct Result
Positive Control	<40	Positive	<36	Positive
PCR Negative Control	No Signal	Negative	No signal*	Negative
Extraction Negative Control	No Signal	Negative	No signal*	Negative
Extraction Sample Control	No Signal	Negative	<36	Positive

^{*}The negative controls are expected to test negative for both the SARS-CoV-2 and Internal Control targets. If the laboratory observes nucleic acid contamination (e.g. HEX Ct values > 36), please review and evaluate your established laboratory procedures intended to prevent environmental sources of human nucleic acid contamination. The internal control target is human RNase P nucleic acid and trace amounts may be present in the laboratory environment.

Sample Validity: The validity for each sample is determined by the internal control result for the respective sample. The table below details the results interpretation of the SARS-CoV-2 and internal control target for each sample.

8 Examination and Interpretation of Patient Specimen Results

Assessment of clinical specimen test results should be performed after the positive and negative controls have been examined and determined to be valid and acceptable. If the controls are not valid, the patient results cannot be interpreted.

<u>Sample Resul</u> t	SARS-CoV-2 FAM Ct Value	Internal Control HEX Ct Value	Other Characteristics
SARS-COV-2 RNA POSITIVE	≤40	Any Ct value	A characteristic amplification curve in comparison to the PCR negative control. An internal control amplification curve in the HEX (VIC) channel is expected. A strong positive SARS CoV-2 sample may result in a negative internal control result.
SARS-CoV-2 RNA NEGATIVE	No Ct value	≤36	Amplification curve in the HEX (VIC) internal control channel
Invalid Sample**	No Ct value	>36	Absence of an amplification curve in the FAM and HEX (VIC) channels indicates an invalid result for the sample.

^{**}An invalid sample can be an indication of failed sample addition, extraction and/or PCR. It is recommended that the RNA be diluted five-fold into PCR grade water and retested; include the undiluted RNA as a sample. If the test is still not valid a new extraction is recommended.

Conditions of Authorization

The OPTI SARS-CoV-2 RT-PCR Test's Letter of Authorization, User Manual, and Labeling are available on FDA website: https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/vitro-diagnostics-euas.

To assist clinical laboratories using the OPTI SARS-CoV-2 RT-PCR Test, the relevant Conditions of Authorization are listed below.

- a) Authorized laboratories using the OPTI SARS-CoV-2 RT-PCR Test will 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.
- b) Authorized laboratories using the OPTI SARS-CoV-2 RT-PCR Test will use your test as outlined in the Instructions for Use only. Deviation from the authorized procedures, such as the authorized instruments, authorized extraction methods, authorized clinical specimen types, authorized control materials, authorized other ancillary reagents and authorized materials required to use your product are not permitted.
- c) Authorized laboratories that receive the OPTI SARS-CoV-2 RT-PCR Test will notify the relevant public health authorities of their intent to run your test prior to initiating testing.
- d) Authorized laboratories using the OPTI SARS-CoV-2 RT-PCR Test will have a process in place for reporting test results to healthcare providers and relevant public health authorities.
- e) Authorized laboratories will collect information on the performance of the OPTI SARS-CoV-2 RT-PCR Test and report to DMD/OHT7-OIR/OPEQ/CDRH (via email: CDRH-EUA-Reporting@fda.hhs.gov) and OPTI Medical Systems, Inc. (via email: COVID19@optimedical.com) if they become aware of any suspected occurrence of false positive or false negative results and significant deviations from the established performance characteristics of the OPTI SARS-CoV-2 RT-PCR Test.
- f) All laboratory personnel using the OPTI SARS-CoV-2 RT-PCR Test must be appropriately trained in RT-PCR techniques and use appropriate laboratory and personal protective equipment when handling this kit and use your test in accordance with the authorized labeling.
- g) OPTI Medical Systems, Inc., 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.
- 1. The letter of authorization refers to, "Laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet requirements to perform high complexity tests" as "authorized laboratories."

Limitations

- The use of this assay as an In vitro diagnostic under the FDA Emergency Use Authorization (EUA) is limited to laboratories that are certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. § 263a, to perform high complexity tests.
- The OPTI SARS-CoV-2 RT-PCR Test can be used with the specimens listed in the Intended Use statement. Other specimen types should not be tested with this assay. Negative results do not preclude infection with SARS-CoV-2 virus and should not be the sole basis of a patient management decision.
- Laboratories are required to report all test results to the appropriate public health authorities.
- Samples must be collected, transported, and stored using appropriate procedures and conditions. Improper collection, transport, or storage of specimens may affect the test performance.
- A false negative result may occur if a specimen is improperly collected, transported or handled. False negative
 results may also occur if amplification inhibitors are present in the specimen or if inadequate numbers of
 organisms are present in the specimen.
- Extraction and amplification of nucleic acid from clinical samples must be performed according to the specified methods listed in this procedure. Other extraction approaches and processing systems have not been validated.
- If the virus mutates in the test target region, SARS-CoV-2 RNA may not be detected or may be detected less predictably. Inhibitors or other types of interference may produce a false negative result.
- False-positive results may arise from cross contamination during specimen handling, preparation, nucleic acid extraction, PCR assay set-up or product handling.
- This test cannot rule out diseases caused by other bacterial or viral pathogens.
- The impacts of vaccines, antiviral therapeutics, antibiotics, chemotherapeutic, immunosuppressant drugs or cold medications have not been evaluated.

Assay Performance

Limit of Detection (LoD)

Limit of detection (LoD) is defined as the lowest concentration of SARS-CoV-2 RNA at which greater than or equal to 95% of all replicates test positive. LoD for the OPTI SARS-CoV-2 RT-PCR Test was determined using serial dilutions of synthetic SARS-CoV-2 RNA (Twist Bioscience, San Francisco, CA, part # MT007544.1) prepared in nasopharyngeal (NP) swab (upper respiratory sample type) sample pools. Samples were collected prior to 2020 and were considered negative for SARS-CoV-2.

The initial LoD was determined with 3-fold serial dilutions tested in triplicate. Each replicate was extracted using the RealPCR DNA/RNA Magnetic Bead Kit on Thermo Scientific™ KingFisher™ Flex following the standard protocol. Extracted RNA was tested on the Applied Biosystems® 7500 PCR instrument (software v2.0.6). To confirm the LoD, 20 replicates of each sample matrix spiked with SARS-CoV-2 RNA were extracted with the RealPCR DNA/RNA Magnetic Bead Kit on the Thermo Scientific™ KingFisher™ Flex and the NucleoMag VET Magnetic Extraction Kit on the Thermo Scientific™ KingFisher™ Duo Prime. Extracted RNA was tested on the Applied Biosystems® 7500 FAST PCR instrument (software v2.0.6). The LoD was confirmed to be 0.9 copies/ µL in NP swab samples (19/20) with both extraction methods. Results are shown in Tables 1 and 2 below.

Additional studies showed comparable LoD results (within 3X of the original confirmed LoD) when using the OPTI DNA/RNA Magnetic Bead Kit (OPTI Medical Systems 99-58015) and PurePrep Pathogens Extraction Kit (MolGen BV OE00290096) on the Thermo Scientific™ KingFisher™ Duo Prime with extracted RNA tested on the Applied Biosystems® 7500 FAST PCR instrument. The Manual protocol for the OPTI DNA/RNA Magnetic Bead Kit was also shown to have comparable LoD results (within 3X of the original confirmed LoD) with the RealPCR DNA/RNA Magnetic Bead Kit, NucleoMag VET Magnetic Extraction Kit and OPTI DNA/RNA Magnetic Bead Kit on the Thermo Scientific™ KingFisher™ Duo Prime or Flex with extracted RNA tested on the Applied Biosystems® 7500 FAST PCR instrument.

Table 1: LoD Initial Determination

NP Swab						
RNA Mean Ct Detection % copies/μL (N1/N2) Rate Detectio						
6.7	NT	NT	NT			
2.2	36.9	3/3	100%			
0.7	38.9	2/3	67%			
0.2	40.0	1/3	33%			

Table 2. LoD Confirmation- Applied Biosystems® 7500 FAST PCR Instrument

		NP Sw	ab	
	RNA copies/μL	Mean Ct (N1/N2)	Detection Rate	LoD copies/µL
RealPCR DNA/RNA	0.9	37.6	19/20	
Magnetic Bead 100uL elution (KingFisher Flex)	0.3	38.5	12/20	0.9
Macherey Nagel NucleoMag VET	0.9	37.3	19/20	0.0
100uL elution (KingFisher DUO Prime)	0.3	38.3	10/20	0.9

Alternate Instrument Testing

An additional study was conducted to determine the LoD for the OPTI SARS-CoV-2 RT-PCR Test using additional PCR instruments. Applied Biosystems QuantStudio 5, Agilent Mx3005P, Roche LightCycler® 480, and Bio Molecular Systems Mic PCR instruments were included in this study. See section "Materials Required but Not Provided" for software versions.

The LoD with was evaluated by testing 20 replicates of pooled nasopharyngeal (NP) swab matrix spiked with SARS-CoV-2 synthetic RNA sourced from Twist Bioscience (part # MT007544.1). Samples were spiked at the LoD concentration which had been previously confirmed for the Applied Biosystems® 7500 FAST PCR instrument. Twenty replicates at the LoD were tested on each instrument. The lowest concentration at which 95% of the replicates were detected was considered the LoD for the instrument. The LoD was 0.9 copies/ul in NP swabs on each of the instruments tested. These results are shown in table 3, below.

Table 3. PCR Instrument LoD Determination

	NP Swab			
PCR Instrument	RNA copies/μL	Mean Ct (N1/N2)	Detection Rate	LoD copies/µL
ABI 7500	0.9	37.6	19/20	0.9
FAST	0.3	38.5	12/20	0.9
ABI QuantStudio 5	0.9	36.5	20/20	0.9
	0.3	37.9	14/20	0.9
Agilent MX3005P	0.9	33.7	19/20	0.9
	0.3	35.2	15/20	0.9
Roche LC480	0.9	36.2	20/20	0.9
	0.3	36.4	13/20	0.9
Bio Molecular Systems MIC	0.9	33.7	20/20	0.9
	0.3	35.0	17/20	0.9

Inclusivity (analytical reactivity)

To assess the in silico inclusivity of the OPTI SARS-CoV-2 RT-PCR Test, a multiple sequence alignment (MSA) was generated from the GISAID CoV database sequences submitted between the dates of December 23, 2019 and October 23, 2020 and compared for identity with the test primers and probes. Only full length, high coverage sequences were used, resulting in over 115,200 strains with sequences in the design regions. 99.7% of sequences match the N1 forward primer, 98.1% match the N1 probe and 99.7% match the N1 reverse primer. Likewise, 99.1% of sequences match the N2 forward primer, 99.7% match the N2 probe and 98.6% match the N2 reverse primer. In total, the combined N1 and N2 designs match 115,249 of 115,277 (99.98%) strains with known sequence in either design region. Generally, failure to detect a positive sample is mitigated through the dual target design of the test. Most sequences only had single base pair mismatches that were not expected to impact detection of the strains due to the location of the mismatch. There were a total of 43 strains that had sequence mismatches in both N1 and N2. Of these, 40 strains were identified in the entire cohort that had a single mismatch in both the N1 and N2 target regions, however all mismatches were located in regions not expected to impact test performance. Two strains contained two mismatches in N1(forward and reverse primers) and one mismatch in N2 (reverse primer). The two N1 mismatches were not in the same primer/probe binding region and are therefore not expected to impact detectability of the strain. One strain contained two mismatches in both N1 (forward and reverse primers) and N2 (reverse primer and probe). Again, the two sets of mismatches were not in the same primer/probe binding region. Therefore, the OPTI SARS-CoV-2 RT-PCR Test would be predicted to amplify and detect all analyzed sequences.

Specificity (Cross-Reactivity)

To access the *in silico* exclusivity of the OPTI SARS-CoV-2 RT-PCR Test, an MSA was generated from several high priority pathogens from the same genetic family as SARS-CoV-2 as well as other high-profile pathogens likely in the same biological niche as SARS-CoV-2. This alignment was then compared for identity to the test primers and probes.

The N1 and N2 design regions were aligned with SARS coronavirus (NC_004718), MERS coronavirus (NC_019843), and human coronaviruses NL63 (NC_005831), OC43 (KX344031), 229E (NC_002645), and HKU1 (NC_006577). No single primer or probe sequence contained greater than 80% identity to the design region. For the organisms listed in Table 4 below, there was insufficient identity to align any of the additional organisms listed. No single organisms contained greater than 80% identity to the design region.

It can reasonably be concluded that the N1 and N2 primers and probes will not amplify and detect any of the virus, bacterial or yeast sequences analyzed.

Table 4: List of organisms analyzed in silico

Organism	Strain	Accession or WGS number
Human Adenovirus	А	NC_001460
Human Metapneumovirus (hMPV)	00-1	NC_039199
Parainfluenza virus 1	NM001	KX639498
Parainfluenza virus 2	VIROAF10	KM190939
Parainfluenza virus 3	CFI1849/2012	KJ672618
Parainfluenza virus 4	SC3019/2015	KY986647
Influenza A	8/1934(H1N1)	NC_002016 to NC_002023
Influenza B	2/2012 BX-51C	MT056021 to MT056028
Enterovirus	D68	MN389735
Respiratory syncytial virus	B/WI/629-Q0190/10	JN032120
Human Rhinovirus	14	NC_001490
Chlamydia pneumoniae	neumoniae CWL029 AE001363	
Haemophilus influenzae	NCTC8143	LN831035
Legionella pneumophila	Phil. 1	CP015928
Mycobacterium tuberculosis	HN-506	AP018036
Streptococcus pneumoniae	NCTC7465	LN831051
Streptococcus pyogenes	NCTC8198	LN831034
Bordetella pertussis	18323	HE965805
Mycoplasma pneumoniae	FH	CP010546
Pneumocystis jirovecii	E2178	NJFV01000001 to NJFV01000219
Candida albicans	dida albicans SC5314 CP017623 to CP017630	
Pseudomonas aeruginosa	PAO1	AE004091
Staphylococcus epidermis	ATCC 12228	NC_004461
Staphylococcus (Streptococcus) salivarius	NCTC8618	NZ_LR134274

Clinical Evaluation

A clinical evaluation study was conducted using real NP swab specimens from patients suspected of COVID-19 by their health care provider that were collected from a CLIA laboratory in the United States. A total of 34 positive (including 7 low positives or 20.6%) and 32 negative NP swab samples were tested with the OPTI SARS-CoV-2 RT PCR Test and compared to results obtained with an FDA EUA authorized RT-PCR test. Samples were extracted with either the OPTI DNA/RNA Magnetic Bead Kit (automated protocol) or RealPCR DNA/RNA Magnetic Bead Kit and RT-PCR was performed using the Applied Biosystems® 7500 FAST PCR instrument. Table 5 and Table 6 summarizes the results including the positive and negative percent agreement with 95% confidence limits.

Table 5. Clinical Samples- OPTI DNA/RNA Magnetic Bead Kit

		FDA EUA RT-PCR Test		
		Positive Patient Specimen	Negative Patient Specimen	Total
OPTI SARS-CoV- 2 RT PCR Test	Positive Patient Specimen	34	0	34
	Negative Patient Specimen	0	32	32
	Total	34	32	

The positive and negative percent agreements between the OPTI SARS-CoV-2 RT PCR Test and FDA EUA test with the OPTI DNA/RNA Magnetic Bead Kit is:

PPA = 100% * 34/34 = 100%. (95% C.I. = 89.85% - 100%)

NPA = 100% * 32/32 = 100%. (95% C.I. = 89.28% - 100%)

Table 6. Clinical samples - RealPCR DNA/RNA Magnetic Bead Kit

	FDA E		RT-PCR Test	
		Positive Patient Specimen	Negative Patient Specimen	Total
OPTI SARS-CoV-2	Positive Patient Specimen	34	0	34
RT PCR Test	Negative Patient Specimen	0	32	32
	Total	34	32	

The positive and negative percent agreements between the OPTI SARS-CoV-2 RT PCR Test and FDA EUA test with the RealPCR DNA/RNA Magnetic Bead Kit is:

PPA = 100% * 34/34 = 100%. (95% C.I. = 89.85% - 100%)

NPA = 100% * 32/32 = 100%. (95% C.I. = 89.28% - 100%)

FDA SARS-CoV-2 Reference Panel Testing

The evaluation of sensitivity and MERS-CoV cross-reactivity was performed using reference material (T1), blinded samples and a standard protocol provided by the FDA. The study included a range finding study and a confirmatory study for LoD. Blinded sample testing was used to establish specificity and to confirm the LoD. The extraction method and instrument used were the OPTI DNA/RNA Magnetic Bead Kit (automated protocol) and the Applied Biosystems® 7500 FAST PCR instrument. The results are summarized in Table 7.

Table 7: Summary of LoD Confirmation Result using the FDA SARS-CoV-2 Reference Panel

Reference Materials Provided by FDA	Specimen Type	Product LoD	Cross-Reactivity
SARS-CoV-2	ND surph	1800 NDU/mL	N/A
MERS-CoV	NP swab	N/A	ND

NDU/mL = RNA NAAT detectable units/mL

N/A: Not applicable ND: Not detected

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Patent information: idexx.com/patents

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Symbol Descriptions

LOT	Batch Code (Lot)
SN	Serial Number
REF	Catalog Number
ECREP	Authorized Representative in the European Community
2	Use by date
~~	Date of manufacture
•••	Manufacturer
	Temperature limitation
	Consult instructions for use
	Major change in the user instructions
IVD	In vitro diagnostics

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