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TaqPath[™] COVID-19 Combo Kit and TaqPath[™] COVID-19 Combo Kit Advanced*

Multiplex real-time RT-PCR test intended for the qualitative detection of nucleic acid from SARS-CoV-2

*TaqPath™ COVID-19 Combo Kit Advanced reagent volumes have been optimized for workflows that use 14.0 µL or 17.5 µL of purified sample RNA

Catalog Numbers A47813 and A47814

Publication Number MAN0019181

Revision J.0







Lif e Technologies Corporation | 6055 Sunol Blv d | Pleasanton, California 94566 USA For descriptions of symbols on product labels or product documents, go to thermofisher.com/symbols-definition.

The customer is responsible for compliance with regulatory requirements that pertain to their procedures and uses of the instrument.

The information in this guide is subject to change without notice.

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Revision history: Pub. No. MAN0019181

Revision	Date	Description	
J.0	22 February 2021	Provided new options for RT-PCR plates.	
H.01	10 December 2020	Internal document control update only, with no changes to Instructions for Use rev. H.0 released on DCO1199965 and posted to the thermofisher.com website. Update to align the 20 Nov 2020 FDA-released Instructions for Use with removal of the licensing statement covering Limited Use Label Licenses on page 2.	
H.0	10 Nov ember 2020	 The instructions for using the TaqPath[™] COVID-19 Combo Kit Advanced were updated to indicate that the kit is now compatible with all instruments. The compatible COVID-19 Interpretive Software versions were updated to v1.5 and 2.5, which include amplification curves. Instructions were added on what to do in the event of bead carry over from RNA extraction. The licensing statement covering Limited Use Label Licenses was removed from page 2. 	
G.0	9 October 2020	 Guidance and protocols were added for samples collected using the Everly well[™] COVID-19 Test Home Collection Kit, including use of the new TaqMan[™] SARS-CoV-2 RNase P Assay Kit. Information and protocols were added for the TaqPath[™] COVID-19 Combo Kit Advanced (Cat. No. A47813), which supports preparation of RT-PCR reactions from 17.5 µL and 14.0 µL of purified sample RNA. "Intended Use" on page 9 and Chapter 12, "Conditions of authorization for labs" were updated with information for the Everly well[™] COVID-19 Test Home Collection Kit and the TaqPath[™] COVID-19 Combo Kit Advanced. COVID-19 Interpretive Software v 1.4 and v .2.4 were added to support use of the RNase P assay, and instructions for obtaining the software were updated. Minor corrections were made to the kit configuration information for the TaqPath[™] COVID-19 Combo Kit. General laboratory recommendations were added (see page 18). The name of the Wash Buffer was changed to Wash Solution in the RNA extraction procedures. A note was added that the Wash Solution can develop particulates but this does not affect performance. "Interpretation of the results" on page 106 was updated to include results for the RNase P assay and guidance on reporting results to the appropriate public health authorities. Minor updates were made to the Intended Use. 	
F.0	15 July 2020	 Added Applied Biosy stems[™] COVID-19 Interpretive Software v1.3 and Applied Biosy stems[™] COVID-19 Interpretive Software v2.3. Removed Applied Biosy stems[™] COVID-19 Interpretive Software v1.2, Applied Biosy stems[™] COVID-19 Interpretive Software v2.0, and Applied Biosy stems[™] COVID-19 Interpretive Software v2.2. Added Ct cutoff information (Appendix B, "Ct cutoff values for assay targets"). Removed instructions to mix by pipetting up and down 10 times when preparing RT-PCR plates. Added instructions to vortex the plates to ensure proper mixing. Updated centrifuge and sealing instructions when preparing the RT-PCR plates. Updated instructions to create a unique name for each well in the physical plate, not just the wells with a patient sample. 	

Revision	Date	Description
F.0 (continued)	15 July 2020	Updated reactivity (inclusivity) (page 115) and the warnings and precautions (page 16).
(continued)		Specified that retesting must be performed by re-extracting the original sample and repeating the RT-PCR ("Interpretation of the results" on page 106).
E.0	12 May 2020	Added the following products as an alternative to the KingFisher [™] 96 KF microplate for the tip comb plate:
		Tip Comb Presenting Plate for KF 96
		 Nunc[™] MicroWell[™] 96-Well Microplate, barcoded
		 Nunc[™] MicroWell[™] 96-Well Microplate, Flat Bottom
		Nunc [™] F96 MicroWell [™] Black Polysty rene Plate
		 KingFisher[™] Deep-Well 96 Plate
D.0	11 May 2020	Updated the list of acceptable sample types and added specimen storage conditions.
		Removed catalog numbers for combo kit components.
		Updated materials listed in "Required materials not supplied" on page 11.
		 Updated the automated RNA extraction procedure with new plastics, new instrument programs, and revised wash and plate handling steps.
		 Updated the manual RNA extraction procedure with an optional ethanol volume for 200-µL sample input volumes, and an option to use the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit for 400-µL sample input volumes.
		Updated the real-time RT-PCR preparation procedure with new instructions for 384-well plates, clarified guidelines for negative controls, and optional plates without barcodes.
		 Added procedures for the QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well block) and the QuantStudio[™] 5 Real-Time PCR Instrument (384-well block).
		 Added instrument firmware requirements and removed calibration plates for the QuantStudio[™] 5 Real-Time PCR Instrument.
		 Added Applied Biosystems[™] COVID-19 Interpretive Software v 2.2.
		Updated control requirements in interpretation of results, based on addition of 384-well plates.
C.0	20 April 2020	Removed 100-reaction kit.
		 Added a catalog number for the KingFisher[™] Deep-Well 96 Plate.
		Updated the catalog number for the Compact Digital Microplate Shaker.
		 Added catalog number for the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit and removed catalog numbers for individual components of the kit.
		 Added the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit.
		• Added an option to extract RNA with 200 µL of sample.
		 Added Applied Biosystems[™] 7500 Real-Time PCR Instrument and Applied Biosystems[™] QuantStudio[™] 5 Real-Time PCR Instruments.
		 Added Applied Biosystems[™] COVID-19 Interpretive Software v 1.2 and Applied Biosystems[™] COVID-19 Interpretive Software v 2.0.
		 Removed Applied Biosystems[™] COVID-19 Interpretive Software v 1.0 and Applied Biosystems[™] and COVID-19 Interpretive Software v 1.1.
		• Added specific instructions to vortex and centrifuge the reaction plate for RT-PCR ("Prepare the RT-PCR reactions (400-µL sample input, 96-well reaction plate, COVID-19 assay only)" on page 40 and "Prepare the RT-PCR reactions (200-µL sample input, 96-well reaction plate, COVID-19 assay only)" on page 36).
		 Specified that retesting must be done with the original sample ("Interpretation of the results" on page 106).
		Reorganized the content to perform RT-PCR based on the real-time PCR instrument.
		Added "Interfering substances" on page 115.
		Added information to customer and technical support (page 124).

Revision	Date	Description
B.0	24 March 2020	 Added MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit components for 1,000 preparations to Required Materials.
		 Added manual RNA extraction protocol and required materials for the manual RNA extraction protocol.
		 Added Applied Biosystems[™] 7500 Fast Real-Time PCR Instrument.
		Removed storage options for RNA after extraction.
		 Updated guidelines for RT-PCR to run the plate immediately after preparation and to keep the plate on ice until it is loaded into the real-time PCR instrument.
		 When setting up the RT-PCR reaction, added instructions to mix by pipetting up and down 10 times and seal and centrifuge the reaction plate.
		 Added COVID-19 Interpretive Software v 1.1 (compatible with Applied Biosystems[™] 7500 Fast Real-Time PCR Instrument and Applied Biosystems[™] 7500 Fast Dx Real-Time PCR Instrument).
		 Added that the run file must be opened, analyzed, and saved in the instrument software before it is opened in COVID-19 Interpretive Software.
		 For TaqPath[™] COVID-19 RT-PCR Kit, 1,000 reactions (Cat. No. A47817), changed MS2 Phage Control from 20 tubes x 500 µL to 10 tubes x 1,000 µL.
		Updated instructions to obtain the COVID-19 Interpretive Software.
		Changed Limit of Detection and Clinical Evaluation data to 1 decimal place.
A.0	15 March 2020	New document.

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TaqPath[™] COVID-19 Combo Kit and TaqPath[™] COVID-19 Combo Kit Advanced product information

Intended Use

The TaqPath™ COVID-19 Combo Kit, which can be labeled as the TaqPath™ COVID-19 Combo Kit Advanced, contains the assays and controls for a real-time reverse transcription polymerase chain reaction (RT-PCR) test intended for the qualitative detection of nucleic acid from SARS-CoV-2 in upper respiratory specimens (such as nasopharyngeal, oropharyngeal, nasal, and mid-turbinate swabs, and nasopharyngeal aspirate) and bronchoalveolar lavage (BAL) specimens from individuals 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.

This test is also authorized for use with the Everlywell[™] COVID-19 Test Home Collection Kit for individuals to self-collect nasal swab specimens when determined by a healthcare provider to be appropriate based on results of a COVID-19 questionnaire.

Results are for the identification of SARS-CoV-2 RNA. The SARS-CoV-2 RNA is generally detectable in upper respiratory and bronchoalveolar lavage (BAL) specimens 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.

Testing with the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced are intended for use by qualified and trained clinical laboratory personnel specifically instructed and trained in the techniques of real-time PCR and *in vitro* diagnostic procedures. The TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced are only for use under the Food and Drug Administration's Emergency Use Authorization.

Product description

The TaqPath[™] COVID-19 Combo Kit and TaqPath[™] COVID-19 Combo Kit Advanced include the assays and controls for a multiplex real-time RT-PCR test for the qualitative detection of RNA from SARS-CoV-2 in upper respiratory specimens (such as nasopharyngeal, oropharyngeal, nasal, and mid-turbinate swabs, and nasopharyngeal aspirate) and bronchoalveolar lavage (BAL) specimens from individuals suspected of COVID-19 by their healthcare provider.

Each kit includes the following components:

- Multiplexed assays that contain three primer/probe sets specific to different SARS-CoV-2 genomic regions and primers/probes for bacteriophage MS2
- MS2 Phage Control as an internal process control for nucleic acid extraction
- TaqPath[™] COVID-19 Control as a positive RNA control that contains targets specific to the SARS-CoV-2 genomic regions targeted by the assays

Contents and storage

Table 1 TaqPath[™] COVID-19 Combo Kit, 1,000 reactions (Cat. No. A47814)

Вох	Components	Amount	Storage
TaqPath™ COVID-19 RT-PCR Kit	COVID-19 Real Time PCR Assay Multiplex (ORF1ab, N gene, S gene, MS2)	1,500 μL	–30°C to −10°C
	MS2 Phage Control	10 × 1,000 μL	-30°C to −10°C
TaqPath [™] COVID-19 Control (1 x 10 ⁴ copies/μL)		2 x 10 μL per box; 5 boxes per kit	≤ –70°C
TaqPath [™] COVID-19 Control Dilution Buffer		2 x 250 µL per box; 5 boxes per kit	-30°C to -10°C

Table 2 TaqPath™ COVID-19 Combo Kit Advanced, 200 reactions (Cat. No. A47813)

Вох	Components	Amount	Storage
TaqPath [™] COVID-19 RT-PCR Kit Advanced	COVID-19 Assay Multiplex Advanced (ORF1ab, N gene, S gene, MS2)	300 μL	−30°C to −10°C
	MS2 Phage Control	2 × 1,000 µL	–30°C to −10°C
TaqPath [™] COVID-19 Control (1 x 10 ⁴ copies/μL)		2 × 10 µL	≤ -70°C
TaqPath™ COVID-19 Control Dilution Buffer		2 × 250 µL	-30°C to -10°C

Required materials not supplied

Unless otherwise indicated, all materials are available through **thermofisher.com**. "MLS" indicates that the material is available from **fisherscientific.com** or another major laboratory supplier.

Catalog numbers that appear as links open the web pages for those products.

Item	Source		
Real-time PCR instrument			
Applied Biosystems [™] 7500 Fast DxReal-Time PCR Instrument (used with SDS Software v1.4.1)	4406984 (with laptop computer) 4406985 (with tower computer)		
Applied Biosystems [™] 7500 Fast Real-Time PCR Instrument (used with SDS Software v1.5.1 or 7500 Software v2.3)	4351106 (with laptop computer) 4351107 (with desktop computer)		
Applied Biosystems [™] 7500 Real-Time PCR Instrument (used with 7500 Software v2.3)	4351104 (with laptop computer) 4351105 (with desktop computer)		
Applied Biosystems [™] QuantStudio [™] 5 Real-Time PCR Instrument, 96-well, 0.2-mL block (used with QuantStudio [™] Design and Analysis Desktop Software v1.5.1)	A28569 (with laptop computer) A28574 (with desktop computer) A28139 (instrument only)		
Applied Biosystems [™] QuantStudio [™] 5 Real-Time PCR Instrument, 96-well, 0.1-mL block (used with QuantStudio [™] Design and Analysis Desktop Software v1.5.1)	A28568 (with laptop computer) A28573 (with desktop computer) A28138 (instrument only)		
Applied Biosystems [™] QuantStudio [™] 5 Real-Time PCR Instrument, 384-well block (used with QuantStudio [™] Design and Analysis Desktop Software v1.5.1)	A28570 (with laptop computer) A28575 (with desktop computer) A28140 (instrument only)		
Applied Biosystems [™] QuantStudio [™] 7 Flex Real-Time PCR Instrument, 384–well block (used with QuantStudio [™] Real-Time PCR Software v1.3)	4485695 (with laptop computer) 4485701 (with desktop computer)		
Equipment			
Laboratoryfreezers • -30°C to -10°C • ≤ -70°C	MLS		
Centrifuge, with a rotor that accommodates standard and deepwell microplates	MLS		
Microcentrifuge	MLS		
Laboratorymixer, vortex or equivalent	MLS		



(continued)

ltem	Source
Single and multichannel adjustable pipettors (1.00 μ L to 1,000.0 μ L)	MLS
Cold block (96-well or 384-well) or ice	MLS
Automated nucleic acid extraction system and materials	
KingFisher [™] FlexMagnetic Particle Processor with 96 Deep- Well Head	5400630
KingFisher [™] Flex96 Deep-Well Heating Block	24075430
KingFisher [™] Deep-Well 96 Plate	95040450, A48305, A48424, 95040455
96-well plate for the tip comb, one of the following: • KingFisher™96 KF microplate • Tip Comb Presenting Plate for KF 96 • Nunc™ MicroWell™ 96-Well Microplate, Flat Bottom • Nunc™ MicroWell™ 96-Well Microplate, barcoded • ABgene™ 96-Well Polypropylene Storage Microplate • ABgene™ 96-Well 1.2-mL Polypropylene Deepwell Storage Plate • Nunc™ F96 MicroWell™ Black Polystyrene Plate • Nunc™ F96 MicroWell™ White Polystyrene Plate • KingFisher™ Deep-Well 96 Plate	 97002540 267600 167008 269787 AB0796 AB1127 137101 136101 95040450, A48305, A48424, 95040455
KingFisher [™] 96 tip comb for DW magnets	97002534, A48438, A48414
Manual nucleic acid extraction system and materials	
Magnetic Stand-96 Compact Digital Microplate Shaker	AM10027 AM10050 88882005
Incubator capable of reaching 65°C with slatted shelves	MLS
KingFisher [™] Deep-Well 96 Plate	95040450, A48305, A48424, 95040455
 Standard 96-well plate for the eluate, one of the following: KingFisher[™] 96 KF microplate MicroAmp[™] Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL MicroAmp[™] Fast Optical 96-Well Reaction Plate, 0.1 mL MicroAmp[™] Optical 96-Well Reaction Plate with Barcode, 0.2 mL MicroAmp[™] Optical 96-Well Reaction Plate, 0.2 mL 	 97002540 4346906, 4366932 4346907 4306737, 4326659 N8010560, 4316813

(continued)

Item	Source	
MicroAmp [™] Clear Adhesive Film	4306311	
Kits and reagents		
Required if specimens were collected using the Everlywell™ COVID-19 Test Home Collection Kit: TaqMan™ SARS-CoV-2 RNase P Assay Kit	A49564	
MagMAX [™] Viral/Pathogen Nucleic Acid Isolation Kit (up to 200 preparations, when 200 µL of sample is used)	A42352	
MagMAX [™] Viral/Pathogen Nucleic Acid Isolation Kit (up to 2,000 preparations, when 200 μL of sample is used)	A48310	
MagMAX [™] Viral/Pathogen II Nucleic Acid Isolation Kit (up to 2,000 preparations, when 200 μL of sample is used)	A48383	
TaqPath [™] 1-Step MultiplexMaster Mix (No ROX [™])	A28521, A28522, A28523	
Fisher BioReagents [™] Ethanol, Absolute, Molecular Biology Grade ^[1] , or equivalent	BP2818100,BP2818500,BP28184	
Nuclease-free Water (not DEPC-Treated)	MLS	
Calibration plates (7500 Real-Time PCR Instrument series)		
ABY [™] Dye Spectral Calibration Plate for MultiplexqPCR, Fast 96-well (0.1-mL)	A24734	
JUN [™] Dye Spectral Calibration Plate for Multiplex qPCR, Fast 96-well (0.1-mL)	A24735	
ABY [™] Dye Spectral Calibration Plate for Multiplex qPCR, 96-well (0.2-mL)	A24738	
JUN [™] Dye Spectral Calibration Plate for Multiplex qPCR, 96-well (0.2-mL)	A24737	
Calibration plates (QuantStudio [™] 7 Flex Real-Time PCR Instru	ment)	
ABY [™] Dye Spectral Calibration Plate for Multiplex qPCR, 384-well	A24736	
JUN [™] Dye Spectral Calibration Plate for Multiplex qPCR, 384-well	A24733	
Tubes, plates, and other consumables		
MicroAmp [™] Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL	4346906, 4366932	
MicroAmp [™] Fast Optical 96-Well Reaction Plate, 0.1 mL	4346907	



(continued)

ltem	Source
MicroAmp [™] EnduraPlate [™] Optical 96-Well Fast Clear Reaction Plates with Barcode ^[2]	4483485, 4483494
MicroAmp [™] EnduraPlate [™] Optical 96-Well Fast Clear Reaction Plates ^[2]	A36930
MicroAmp [™] Optical 96-Well Reaction Plate with Barcode, 0.2 mL	4306737,4326659
MicroAmp [™] Optical 96-Well Reaction Plate, 0.2 mL	N8010560, 4316813
MicroAmp [™] EnduraPlate [™] Optical 96-Well Clear Reaction Plates with Barcode ^[2]	4483354, 4483352
MicroAmp [™] EnduraPlate [™] Optical 96-Well Clear Reaction Plates ^[2]	A36924
MicroAmp [™] Optical 384-Well Reaction Plate with Barcode	4309849,4326270,4343814
MicroAmp [™] Optical 384-Well Reaction Plate	4343370
MicroAmp [™] EnduraPlate [™] Optical 384-Well Clear Reaction Plates with Barcode ^[2]	4483285, 4483273
MicroAmp [™] EnduraPlate [™] Optical 384-Well Clear Reaction Plates ^[2]	A36931
MicroAmp™ Clear Adhesive Film	4306311
MicroAmp [™] Optical Adhesive Film	4311971,4360954
MicroAmp [™] Adhesive Film Applicator	4333183
Nonstick, RNase-free microcentrifuge tubes (1.5 mL and 2.0 mL)	thermofisher.com/plastics
Sterile aerosol barrier (filtered) pipette tips	thermofisher.com/pipettetips

 $^{^{[1]}}$ Available at fisherscientific.com.

^[2] Not compatible with the Everlywell[™] COVID-19 Test Home Collection Kit procedure or the TaqPath[™] COVID-19 RT-PCR Kit Advanced procedure.



Instrument, assay, and software compatibility

The following table lists the version of the Applied Biosystems[™] COVID-19 Interpretive Software that is compatible with your instrument, its associated analysis software, and whether the TaqMan[™] SARS-CoV-2 RNase P Assay is used in the test procedure.

Note: The TaqMan[™] SARS-CoV-2 RNase P Assay is required for specimens collected using the Everlywell[™] COVID-19 Test Home Collection Kit.

For information on how to obtain the Applied Biosystems[™] COVID-19 Interpretive Software, see "Obtain the Applied Biosystems[™] COVID-19 Interpretive Software" on page 105.

To obtain the analysis software or firmware for your real-time PCR instrument, go to **thermofisher.com/ qpcrsoftware**, then select your instrument in the **Real-Time PCR** section.

Instrument	Analysis software used with the instrument	Minimum compatible COVID-19 Interpretive Software version
7500 Fast Dx Real-Time PCR Instrument	SDS Software v1.4.1	v1.5
7500 Fast Real-Time PCR Instrument	SDS Software v1.5.1 or 7500 Software v2.3	v1.5
7500 Real-Time PCR Instrument	7500 Software v2.3	v1.5
QuantStudio [™] 5 Real-Time PCR Instrument with instrument firmware v1.3.3 • 96-well, 0.2-mL block • 96-well, 0.1-mL block • 384-well block	QuantStudio [™] Design and Analysis Desktop Software v1.5.1	v2.5
QuantStudio [™] 7 Flex Real-Time PCR Instrument with instrument firm ware v1.0.4 • 384-well block	QuantStudio [™] Real-Time PCR Software v1.3	v2.5

Warnings and precautions

The TaqPath[™] COVID-19 RT-PCR Kit and the TaqPath[™] COVID-19 RT-PCR Kit Advanced workflows should be performed by qualified and trained staff to avoid the risk of erroneous results. Use separate areas for the preparation of patient samples and controls to prevent false positive results. Samples and reagents must be handled in a biological safety cabinet.

- The assay is for in vitro diagnostic use under the FDA Emergency Use Authorization Only.
- This test has not been FDA cleared or approved.
- This test has been authorized only for the detection of nucleic acid from SARS-CoV-2, not for any other viruses or pathogens.
- 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 Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization
 is terminated or revoked sooner.
- Samples and controls should always be treated as if infectious and/or biohazardous in accordance with safe laboratory procedures.
- Follow necessary precautions when handling specimens. Use personal protective equipment (PPE) consistent with current guidelines for the handling of potentially infectious samples.
- Always use pipette tips with aerosol barriers. Tips that are used must be sterile and free from DNases and RNases.
- Do not eat, drink, smoke, or apply cosmetic products in the work areas.
- Modifications to assay reagents, assay protocol, or instrumentation are not permitted, and are in violation of the product Emergency Use Authorization.
- Reagents must be stored and handled as specified in "Contents and storage" on page 10.
- Do not use the kits after the indicated expiry date.
- Dispose of waste in compliance with local, state, and federal regulations.
- Safety Data Sheets are available upon request.
- Laboratories within the United States and its territories are required to report all test results to the appropriate public health authorities.
- Positive results are indicative of the presence of SARS-CoV-2 RNA.



Assay 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, that meet requirements to perform high complexity tests.
- The TaqPath™ COVID-19 RT-PCR Kit and the TaqPath™ COVID-19 RT-PCR Kit Advanced performance was established using nasopharyngeal and oropharyngeal swab, nasopharyngeal aspirate, and bronchoalveolar lavage samples only. Nasal swabs and mid-turbinate swabs are considered acceptable specimen types for use with the TaqPath™ COVID-19 RT-PCR Kit and the TaqPath™ COVID-19 RT-PCR Kit Advanced, but performance with these specimen types has not been established. Refer to FDA's FAQs on Diagnostic Testing for SARS-CoV-2 for additional information. Specimen types other than nasopharyngeal, oropharyngeal, nasal and mid-turbinate nasal swabs, nasopharyngeal aspirate, and bronchoalveolar lavage should not be tested with this assay.
- Samples must be collected, transported, and stored using appropriate procedures and conditions.
 Improper collection, transport, or storage of specimens may hinder the ability of the assay to detect the target sequences.
- Extraction and amplification of nucleic acid from clinical samples must be performed according the specified methods listed in this procedure. Other extraction approaches and processing systems have not been evaluated.
- Specimens submitted using the Everlywell[™] COVID-19 Test Home Collection Kit must be tested using the TaqMan[™] SARS-CoV-2 RNase P Assay Kit and the TaqPath[™] COVID-19 RT-PCR Kit.
- False-negative results may arise from:
 - Improper sample collection
 - Degradation of the SARS-CoV-2 RNA during shipping/storage
 - Specimen collection after SARS-CoV-2 RNA can no longer be found in the specimen matrix
 - Using unauthorized extraction or assay reagents
 - The presence of RT-PCR inhibitors
 - Mutation in the SARS-CoV-2 virus
 - Failure to follow instructions for use
- False-positive results may arise from:
 - Cross contamination during specimen handling or preparation
 - Cross contamination between patient samples
 - Specimen mix-up
 - RNA contamination during product handling
- The impacts of vaccines, antiviral therapeutics, antibiotics, chemotherapeutic or immunosuppressant drugs have not been evaluated. The TaqPath[™] COVID-19 RT-PCR Kit and the TaqPath[™] COVID-19 RT-PCR Kit Advanced cannot rule out diseases caused by other bacterial or viral pathogens.
- 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.

- Detection of RNase P indicates that human nucleic acid is present and implies that human biological material was collected and successfully extracted and amplified. It does not necessarily indicate that the specimen is of appropriate quality to enable detection of SARS-CoV-2.
- MicroAmp[™] EnduraPlate[™] products are not compatible with the Everlywell[™] COVID-19 Test Home Collection Kit procedure or the TaqPath[™] COVID-19 RT-PCR Kit Advanced procedure.

General laboratory recommendations

- Implement standard operating procedures in your laboratory to prevent contamination, such as the following:
 - Frequent glove changes
 - Frequent decontamination of surfaces, equipment, and pipettes with 10% bleach or decontamination solution, followed by 70% ethanol
 - Use of ultraviolet light during biosafety cabinet decontamination (when available)
- To prevent degradation, keep eluted sample RNA, master mixes, assays, and controls on ice or in cold blocks while in use.
- Limit freeze-thaw cycles.
- Aliquot reagents to prevent stock contamination and reduce the number of freeze-thaw cycles.
- After each run, review the amplification curves in the interpretive software for signs of inadequate vortexing or centrifugation. Contact your Applications Support team for additional information or training on data QC in your instrument software.

Samples and controls

Patient samples must be collected according to appropriate laboratory guidelines. Positive and negative test controls must be included to accurately interpret patient test results.

Store patient samples according to CDC guidelines. See the CDC website: https://www.cdc.gov/coronavirus/2019-ncov/lab/guidelines-clinical-specimens.html.

Include the following controls:

Control	Used to monitor	Assays
Positive Control (TaqPath [™] COVID-19 Control Kit)	RT-PCR reaction setup and reagent integrity	All three SARS-CoV-2 assays
MS2 Phage Control	RNA extraction	MS2 assay
Negative Control	Cross-contamination during RNA	All three SARS-CoV-2 assays
	extraction and reaction setup	MS2 assay
TaqMan [™] SARS-CoV-2 RNase P Assay control ^[1]	Sample adequacy	RNase Passay

^[1] Used with the Everlywell[™] COVID-19 Test Home Collection Kit workflow.



Sample collection, transport, and storage

Note: Handle all samples and controls as if they are capable of transmitting infectious agents.

Workflow

Extract RNA from patient sample



Perform RT-PCR



Analyze data using the Applied Biosystems[™] COVID-19 Interpretive Software



Review run control results



Review results interpretation for patient samples

The workflow begins with nucleic acid extraction from upper respiratory specimens (such as nasopharyngeal, oropharyngeal, nasal, and mid-turbinate swabs, and nasopharyngeal aspirate) and bronchoalveolar lavage (BAL) specimens that arrive in the testing site in transport media. Nucleic acids are isolated and purified from the specimens using the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit. Nucleic acid isolation can be performed manually or via an automated process using the KingFisher[™] Flex Purification System (KingFisher). For more information about using the kit, see "Related documentation" on page 124.

The RT-PCR reactions are prepared. For specimens collected using the Everlywell[™] COVID-19 Test Home Collection Kit, the TaqMan[™] SARS-CoV-2 RNase P Assay Kit is included as an additional control.

The nucleic acid is reverse transcribed into cDNA and amplified using the TaqPath[™] COVID-19 RT-PCR Kit or the TaqPath[™] COVID-19 RT-PCR Kit Advanced and one of the following real-time PCR instruments:

- Applied Biosystems[™] 7500 Fast Dx Real-Time PCR instrument
- Applied Biosystems[™] 7500 Fast Real-Time PCR Instrument
- Applied Biosystems[™] 7500 Real-Time PCR Instrument
- Applied Biosystems[™] QuantStudio[™] 5 Real-Time PCR Instrument, 96-well, 0.2–mL block
- Applied Biosystems[™] QuantStudio[™] 5 Real-Time PCR Instrument, 96-well, 0.1–mL block
- Applied Biosystems[™] QuantStudio[™] 5 Real-Time PCR Instrument, 384-well block
- Applied Biosystems[™] QuantStudio[™] 7 Flex Real-Time PCR Instrument, 384–well block



In the process, the probes anneal to three (3) specific SARS-CoV-2 target sequences located between three (3) unique forward and reverse primers for the following genes:

- ORF1ab
- N Gene
- S Gene

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. Fluorescence intensity is monitored at each PCR cycle by the real-time PCR instrument.

The data are analyzed, then interpreted by the Applied Biosystems[™] COVID-19 Interpretive Software.

The workflow options have been updated to reduce the number of consumables. Previous versions of the workflow in previous revisions of this document are still validated to run the TaqPath[™] COVID-19 RT-PCR Kit.



Extract RNA (automated method)

Before you begin	2′
Extract RNA—Automated method (200-µL sample input volume)	22
Extract RNA—Automated method (400-µL sample input volume)	25

Automated RNA extraction is performed using the KingFisher[™] Flex Magnetic Particle Processor with 96 Deep-Well Head and the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit or MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit with a sample input volume of 200 µL or 400 µL.

Before you begin

Note: During the wash steps, the Wash Solution may develop inert white or brown particulates that float in solution. This is not a cause for concern and does not negatively affect performance.

- Determine the number of required reactions based on the number of patient samples to be processed, plus one Negative Control per plate.
- Prepare fresh 80% Ethanol using Ethanol, Absolute, Molecular Biology Grade and Nuclease-free Water (not DEPC-Treated) for the required number of reactions, sufficient for 1 mL per reaction, plus 10% overage.
- Label the short side of each KingFisher[™] Deep-Well 96 Plate (4):

Label	Number of plates
Sample plate	1
Wash 1	1
Wash 2	1
Elution plate	1

Label the short side of the KingFisher[™] 96 KF microplate (1):

Label	Number of plates
Tip comb	1

Note: The following items can be used to hold the tip comb instead of the KingFisher[™] 96 KF microplate:

- Tip Comb Presenting Plate for KF 96
- Nunc[™] MicroWell[™] 96-Well Microplate, Flat Bottom
- Nunc[™] MicroWell[™] 96-Well Microplate, barcoded
- ABgene[™] 96–Well Polypropylene Storage Microplate

- ABgene[™] 96–Well 1.2–mL Polypropylene Deepwell Storage Plate
- Nunc[™] F96 MicroWell[™] Black Polystyrene Plate
- Nunc[™] F96 MicroWell[™] White Polystyrene Plate
- KingFisher[™] Deep-Well 96 Plate
- Mark the Negative Control well on the plate.

Extract RNA—Automated method (200-µL sample input volume)

The following procedure uses components from the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit or the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit.

Set up the instrument (200-µL sample input volume)

- 1. Ensure that the KingFisher[™] Flex Magnetic Particle Processor with 96 Deep-Well Head is set up with the KingFisher[™] Flex 96 Deep-Well Heating Block.
 - IMPORTANT! Failure to use the proper magnetic head and heat block results in lower yields and potential harm to the instrument.
- 2. Ensure that the MVP_2Wash_200_Flex program has been downloaded from the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit product page at www.thermofisher.com and loaded onto the instrument.

Prepare the processing plates (200-µL sample input volume)

Prepare the processing plates according to the following table. Cover the plates with a temporary seal (such as MicroAmp[™] Clear Adhesive Film), then store at room temperature for up to 1 hour while you set up the sample plate.

Plate ID	Plate position	Plate type	Reagent	Volume per well
Wash 1 Plate	2		Wash Solution	500 μL
Wash 2 Plate	3	KingFisher [™] Deep-Well 96 Plate	80% Ethanol	1,000 µL
Elution Plate	4		Elution Solution	50 μL
Tip Comb Plate	5	Place a KingFisher [™] 96 tip comb f KF mi	or DW magnets in a Kir croplate	ngFisher [™] 96

Note: The following items can be used to hold the tip comb instead of the KingFisher[™] 96 KF microplate:

- Tip Comb Presenting Plate for KF 96
- Nunc[™] MicroWell[™] 96-Well Microplate, Flat Bottom
- Nunc[™] MicroWell[™] 96-Well Microplate, barcoded
- ABgene[™] 96–Well Polypropylene Storage Microplate
- ABgene[™] 96–Well 1.2–mL Polypropylene Deepwell Storage Plate
- Nunc[™] F96 MicroWell[™] Black Polystyrene Plate
- Nunc[™] F96 MicroWell[™] White Polystyrene Plate
- KingFisher[™] Deep-Well 96 Plate

Prepare Binding Bead Mix (200-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Total Nucleic Acid Magnetic Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well ^[1]
Binding Solution	265 μL
Total Nucleic Acid Magnetic Beads	10 µL
Total volume per well	275 μL

^[1] Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

3. Mix well by inversion, then store at room temperature.

Prepare sample plate (200-µL sample input volume)

This section provides volumes for the sample plate. Your plate layout will depend on the number of samples you run.

IMPORTANT! Samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit have special plate layout considerations. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)".

- Add 5 µL of Proteinase K to each well in the KingFisher[™] Deep-Well 96 Plate labeled "Sample Plate".
- 2. Add 200 µL of sample to each sample well.
 - For Everlywell[™] COVID-19 Test Home Collection Kit samples, not all sample wells are used. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)".
- 3. Add 200 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.

 For Everlywell™ COVID-19 Test Home Collection Kit samples, prepare 2 Negative Control wells on the extraction plate if you are splitting a single extraction plate into two 96-well RT-PCR plates. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)".
- 4. Invert the Binding Bead Mix 5 times gently to mix, then add 275 μL to each sample well and the Negative Control well in the Sample Plate.

Note: Remix Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

5. Add 5 µL of MS2 Phage Control to each sample well and Negative Control well.

Process the samples (200-µL sample input volume)

- Select the MVP_2Wash_200_Flex on the KingFisher[™] Flex Magnetic Particle Processor with 96 Deep-Well Head.
- 2. Start the run, then load the prepared plates into position when prompted by the instrument.
- 3. After the run is complete (~22 minutes after start), immediately remove the Elution Plate from the instrument, then cover the plate with MicroAmp[™] Clear Adhesive Film.

IMPORTANT! To prevent evaporation, seal the plate containing the eluate immediately.

The samples are eluted in 50 µL of Elution Solution (see "Prepare the processing plates (200-µL sample input volume)" on page 23).

Note:

- Significant bead carry over may adversely impact RT-PCR performance. If bead carry over is observed, repeat the test by re-extracting a new aliquot of the sample.
- To ensure reliable performance of the KingFisher[™] Flex Magnetic Particle Processor, perform preventive maintenance as instructed by the manufacturer.

Place the Elution Plate on ice for immediate use in real-time RT-PCR.

Extract RNA—Automated method (400-µL sample input volume)

The following procedure uses components from the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit or the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit.

Set up the instrument (400-µL sample input volume)

- Ensure that the KingFisher[™] Flex Magnetic Particle Processor with 96 Deep-Well Head is set up with the KingFisher[™] Flex 96 Deep-Well Heating Block.
 - IMPORTANT! Failure to use the proper magnetic head and heat block results in lower yields and potential harm to the instrument.
- Ensure that the MVP_2Wash_400_Flex program has been downloaded from the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit product page at www.thermofisher.com and loaded onto the instrument.

Prepare the processing plates (400-µL sample input volume)

Prepare the processing plates according to the following table. Cover the plates with a temporary seal (such as MicroAmp[™] Clear Adhesive Film), then store at room temperature for up to 1 hour while you set up the sample plate.

Plate ID	Plate position	Plate type	Reagent	Volume per well
Wash 1 Plate	2		Wash Solution	1,000 µL
Wash 2 Plate	3	KingFisher [™] Deep-Well 96 Plate	80% Ethanol	1,000 µL
Elution Plate	4		Elution Solution	50 µL
Tip Comb Plate	5	Place a KingFisher [™] 96 tip comb for DW magnets in a KingFisher [™] 96 KF microplate		

Note: The following items can be used to hold the tip comb instead of the KingFisher[™] 96 KF microplate:

- · Tip Comb Presenting Plate for KF 96
- Nunc[™] MicroWell[™] 96-Well Microplate, Flat Bottom
- Nunc[™] MicroWell[™] 96-Well Microplate, barcoded
- ABgene[™] 96–Well Polypropylene Storage Microplate
- ABgene[™] 96–Well 1.2–mL Polypropylene Deepwell Storage Plate
- Nunc[™] F96 MicroWell[™] Black Polystyrene Plate
- Nunc[™] F96 MicroWell[™] White Polystyrene Plate
- KingFisher[™] Deep-Well 96 Plate

Prepare Binding Bead Mix (400-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Total Nucleic Acid Magnetic Beads to ensure that the bead mixture is homogeneous.
- For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well ^[1]
Binding Solution	530 µL
Total Nucleic Acid Magnetic Beads	20 μL
Total volume per well	550 µL

^[1] Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

3. Mix well by inversion, then store at room temperature.

Prepare sample plate (400-µL sample input volume)

This section provides volumes for the sample plate. Your plate layout will depend on the number of samples you run.

IMPORTANT! Samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit have special plate layout considerations. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)".

- Add 10 µL of Proteinase K to each well in the KingFisher[™] Deep-Well 96 Plate labeled "Sample Plate".
- 2. Add 400 µL of sample to each sample well.
 - For Everlywell[™] COVID-19 Test Home Collection Kit samples, not all sample wells may be used. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)".
- 3. Add 400 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.

 For Everlywell™ COVID-19 Test Home Collection Kit samples, prepare 2 Negative Control wells on the extraction plate if you are splitting a single extraction plate into two 96-well RT-PCR plates. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)".
- 4. Invert the Binding Bead Mix 5 times gently to mix, then add 550 μL to each sample well and the Negative Control well in the Sample Plate.

Note: Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

5. Add 10 µL of MS2 Phage Control to each sample well and to the Negative Control well.

Process the samples (400-µL sample input volume)

- Select the MVP_2Wash_400_Flex on the KingFisher[™] Flex Magnetic Particle Processor with 96 Deep-Well Head.
- 2. Start the run, then load the prepared plates into position when prompted by the instrument.
- 3. After the run is complete (~24 minutes after start), immediately remove the Elution Plate from the instrument, then cover the plate with MicroAmp[™] Clear Adhesive Film.

IMPORTANT! To prevent evaporation, seal the plate containing the eluate immediately.

The samples are eluted in 50 μ L of Elution Solution (see "Prepare the processing plates (400- μ L sample input volume)" on page 26).

Chapter 2 Extract RNA (automated method) Extract RNA—Automated method (400-µL sample input volume)



- Significant bead carry over may adversely impact RT-PCR performance. If bead carry over is observed, repeat the test by re-extracting a new aliquot of the sample.
- To ensure reliable performance of the KingFisher[™] Flex Magnetic Particle Processor, perform preventive maintenance as instructed by the manufacturer.

Place the Elution Plate on ice for immediate use in real-time RT-PCR.



Extract RNA (manual method)

Before you begin	29
Extract RNA—Manual method (200-µL sample input volume)	29
Extract RNA—Manual method (400-µL sample input volume)	32

Manual RNA extraction can be performed from a sample input volume of 200 µL or 400 µL using either the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit or the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit.

Before you begin

Note: During the wash steps, the Wash Solution may develop inert white or brown particulates that float in solution. This is not a cause for concern and does not negatively affect performance.

- Determine the number of required reactions based on the number of patient samples to be processed, plus one Negative Control per plate.
- Prepare fresh 80% Ethanol using Ethanol, Absolute, Molecular Biology Grade and Nuclease-free Water (not DEPC-Treated) for the required number of reactions, plus 10% overage.

Sample input volume	Volume of 80% Ethanol per reaction
200 μL	0.75 mL
400 μL	1.5 mL

Mark the Negative Control well on the plate.

Extract RNA—Manual method (200-µL sample input volume)

The following procedure uses components from the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit or the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit.

Prepare Binding Bead Mix (200-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Total Nucleic Acid Magnetic Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well ^[1]
Binding Solution	265 μL
Total Nucleic Acid Magnetic Beads	10 µL
Total volume per well	275 μL

^[1] Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

3. Mix well by inversion, then store at room temperature.

Digest with Proteinase K (200-µL sample input volume)

This section provides volumes for the sample plate. Your plate layout will depend on the number of samples you run.

IMPORTANT! Samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit have special plate layout considerations. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)".

- 1. Add 5 µL of Proteinase K to each well of a KingFisher[™] Deep-Well 96 Plate.
- 2. Add 200 µL of sample to each sample well.
 For Everlywell™ COVID-19 Test Home Collection Kit samples, not all sample wells may be used. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)".
- 3. Add 200 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well. For Everlywell™ COVID-19 Test Home Collection Kit samples, prepare 2 Negative Control wells on the extraction plate if you are splitting a single extraction plate into two 96-well RT-PCR plates. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)".
- Invert the Binding Bead Mix 5 times gently to mix, then add 275 μL to each sample well and Negative Control well.

Note: Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

5. Add 5 μ L of MS2 Phage Control to each sample well and to the Negative Control well.

- 6. Seal the plate with MicroAmp[™] Clear Adhesive Film, then shake the sealed plate at 1,050 rpm for 2 minutes.
- 7. Incubate the sealed plate at 65°C for 5 minutes (ensure the bottom of the plate is uncovered), then shake the plate at 1,050 rpm for 5 minutes.
- 8. Place the sealed plate on the magnetic stand for 10 minutes or until all of the beads have collected.

Wash the beads (200-µL sample input volume)

1. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

IMPORTANT! Avoid disturbing the beads.

- 2. Remove the plate from the magnetic stand, then add 500 µL of Wash Solution to each sample.
- 3. Reseal the plate, then shake at 1,050 rpm for 1 minute.
- 4. Place the plate back on the magnetic stand for 2 minutes, or until all the beads have collected.
- 5. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

IMPORTANT! Avoid disturbing the beads.

- 6. Repeat step 2 to step 5 using 500 µL of 80% Ethanol.
- 7. Repeat step 2 to step 5 using 250 µL of 80% Ethanol.
- 8. Dry the beads by shaking the plate (uncovered) at 1,050 rpm for 2 minutes.

Elute the nucleic acid (200-µL sample input volume)

- Add 50 µL of Elution Solution to each sample, then seal the plate with MicroAmp[™] Clear Adhesive Film.
- 2. Shake the sealed plate at 1,050 rpm for 5 minutes.
- 3. Place the plate in an incubator at 65°C for 10 minutes.
- 4. Remove the plate from the incubator, then shake the plate at 1,050 rpm for 5 minutes.
- 5. Place the sealed plate on the magnetic stand for 3 minutes or until clear to collect the beads against the magnets.
- 6. Keeping the plate on the magnet, carefully remove the seal, transfer the eluates to a fresh standard (not deep-well) 96-well plate, then seal the plate with MicroAmp[™] Clear Adhesive Film.
 - IMPORTANT! To prevent evaporation, seal the plate containing the eluate immediately after the transfers are complete.

Note: Significant bead carry over may adversely impact RT-PCR performance. If bead carry over is observed, repeat the test by re-extracting a new aliquot of the sample.

Place the plate on ice for immediate use in real-time RT-PCR.

Extract RNA—Manual method (400-µL sample input volume)

The following procedure uses components from the MagMAX[™] Viral/Pathogen Nucleic Acid Isolation Kit or the MagMAX[™] Viral/Pathogen II Nucleic Acid Isolation Kit.

Prepare Binding Bead Mix (400-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Total Nucleic Acid Magnetic Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well ^[1]
Binding Solution	530 µL
Total Nucleic Acid Magnetic Beads	20 µL
Total volume per well	550 μL

^[1] Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

3. Mix well by inversion, then store at room temperature.

Digest with Proteinase K (400-µL sample input volume)

This section provides volumes for the sample plate. Your plate layout will depend on the number of samples you run.

IMPORTANT! Samples collected using the Everlywell™ COVID-19 Test Home Collection Kit have special plate layout considerations. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)".

- 1. Add 10 µL of Proteinase K to each well of a KingFisher[™] Deep-Well 96 Plate.
- 2. Add 400 µL of sample to each sample well.

For Everlywell[™] COVID-19 Test Home Collection Kit samples, not all sample wells may be used. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)".

- 3. Add 400 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.

 For Everlywell™ COVID-19 Test Home Collection Kit samples, prepare 2 Negative Control wells on the extraction plate if you are splitting a single extraction plate into two 96-well RT-PCR plates. See Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19
- 4. Invert the Binding Bead Mix 5 times gently to mix, then add 550 μL to each sample well and Negative Control well.

Note: Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

- 5. Add 10 µL of MS2 Phage Control to each sample well and to the Negative Control well.
- 6. Seal the plate with MicroAmp[™] Clear Adhesive Film, then shake the sealed plate at 1,050 rpm for 2 minutes.
- 7. Incubate the sealed plate at 65°C for 5 minutes (ensure the bottom of the plate is uncovered), then shake the plate at 1,050 rpm for 5 minutes.
- 8. Place the sealed plate on the magnetic stand for 10 minutes or until all of the beads have collected.

Wash the beads (400-µL sample input volume)

Test Home Collection Kit samples)".

1. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

IMPORTANT! Avoid disturbing the beads.

- 2. Remove the plate from the magnetic stand, then add 1 mL of Wash Solution to each sample.
- 3. Reseal the plate, then shake at 1,050 rpm for 1 minute.
- 4. Place the plate back on the magnetic stand for 2 minutes, or until all the beads have collected.
- 5. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

IMPORTANT! Avoid disturbing the beads.

- 6. Repeat step 2 to step 5 using 1 mL of 80% Ethanol.
- 7. Repeat step 2 to step 5 using 500 µL of 80% Ethanol.
- 8. Dry the beads by shaking the plate (uncovered) at 1,050 rpm for 2 minutes.

Elute the nucleic acid (400-µL sample input volume)

- Add 50 µL of Elution Solution to each sample, then seal the plate with MicroAmp[™] Clear Adhesive Film.
- 2. Shake the sealed plate at 1,050 rpm for 5 minutes.
- 3. Place the plate in an incubator at 65°C for 10 minutes.
- 4. Remove the plate from the incubator, then shake the plate at 1,050 rpm for 5 minutes.
- 5. Place the sealed plate on the magnetic stand for 3 minutes or until clear to collect the beads against the magnets.
- 6. Keeping the plate on the magnet, carefully remove the seal, transfer the eluates to a fresh standard (not deep-well) 96-well plate, then seal the plate with MicroAmp[™] Clear Adhesive Film.

IMPORTANT! To prevent evaporation, seal the plate containing the eluate immediately after the transfers are complete.

Note: Significant bead carry over may adversely impact RT-PCR performance. If bead carry over is observed, repeat the test by re-extracting a new aliquot of the sample.

Place the plate on ice for immediate use in real-time RT-PCR.



Prepare RT-PCR reactions— COVID-19 assay only

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Protocols in this chapter

This chapter covers preparation of RT-PCR reaction plates for samples that were collected using methods other than the Everlywell™ COVID-19 Test Home Collection Kit. The RT-PCR reactions in this chapter use the COVID-19 Real Time PCR Assay Multiplex or COVID-19 Assay Multiplex Advanced, but do not include the RNase P assay.

Note: For samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit, see Chapter 5, "Prepare RT-PCR reactions—COVID-19 assay plus RNase P assay (Everlywell[™] COVID-19 Test Home Collection Kit)".

IMPORTANT! Do not combine samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit with samples collected via other methods on the same RT-PCR plate. The plate setup and software template files are different.

In this chapter, select the appropriate RT-PCR preparation procedure based on the following criteria:

- The original sample input volume used for RNA extraction (200 μL or 400 μL)
- The size of the RT-PCR reaction plate (96–well or a 384–well)
- The volume of purified sample RNA that is added to the reaction

Guidelines for RT-PCR

IMPORTANT!

- For each RT-PCR reaction plate, include the following controls:
 - One Positive Control
 - One Negative Control from each extraction run.
 For example, if RNA samples from 4 extraction runs are combined on one 384-well RT-PCR reaction plate, then 4 Negative Control wells must be run on that 384-well reaction plate.
- Prepare the RT-PCR reaction plate on ice and keep it on ice until it is loaded into the real-time PCR instrument.
- Run the plate immediately after preparation. Failure to do so could result in degraded RNA samples.
- To prevent contamination, prepare reagents in a PCR workstation or equivalent amplicon-free area.
 Do not use the same pipette for controls and RNA samples, and always use aerosol barrier pipette tips.
- Maintain an RNase-free environment.
- Protect assays from light.
- · Keep RNA samples and components on ice during use.

Prepare the RT-PCR reactions (200-µL sample input, 96-well reaction plate, COVID-19 assay only)

Use this procedure under the following conditions:

- Original sample input volume of 200 µL was used for extraction
- Instrument is compatible with 96-well RT-PCR reaction plates
- Sample was not collected with the Everlywell[™] COVID-19 Test Home Collection Kit (no RNase P assay required)
- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 μL of TaqPath[™] COVID-19 Control. Mix well, then centrifuge briefly.
 - Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the dilution created in substep 3a. Mix well, then centrifuge briefly.

Note: The TagPath[™] COVID-19 Control does not contain the MS2 template.

4. Prepare the Reaction Mix:

a. For each run, combine the following components sufficient for the number of RNA samples to be tested plus one Positive Control and one Negative Control.

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in this table assume that you extracted sample RNA using an original sample input volume of 200 μ L.

Component	Volume per RNA sample or control	Volume for <i>n</i> RNA samples plus 2 controls	Volume for 94 RNA Samples plus 2 controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 µL	6.875 x (<i>n</i> + 2) μL	660 µL
COVID-19 Real Time PCR Assay Multiplex	1.25 μL	1.375 x (<i>n</i> + 2) μL	132 µL
Nuclease-free Water	7.50 µL	8.25 x (n + 2) μL	792 µL
Total Reaction Mix volume	15.0 µL	_	1584 µL

5. Set up the reaction plate:

a. Pipet 15.0 µL of the Reaction Mix prepared in step 4 into each well of a MicroAmp[™] Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate with Barcode, 0.2 mL.

For other reaction plates that can be used, see "Required materials not supplied" on page 11.

- b. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
- c. Unseal the plate containing the purified sample RNA and Negative Control from the RNA extraction procedure. Add either sample RNA, Negative Control, or Positive Control to each well of the reaction plate according to Table 3 on page 38.
- d. Seal the plate thoroughly with MicroAmp[™] Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

e. Vortex the plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.

IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.

f. Centrifuge the reaction plate for 1–2 minutes at \geq 650 × g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Table 3 Reaction plate volumes

Component	Volume per reaction		
Component	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix	15.0 µL	15.0 µL	15.0 μL
Purified sample RNA (from RNA extraction)	10.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control, from step 3)	_	2.0 µL	_
Nuclease-free Water	_	8.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	10.0 µL
Total volume	25.0 µL	25.0 μL	25.0 μL

Prepare the RT-PCR reactions (200-µL sample input, 384-well reaction plate, COVID-19 assay only)

Use this procedure under the following conditions:

- Original sample input volume of 200 µL was used for extraction
- Instrument is compatible with 384-well RT-PCR reaction plates
- Sample was not collected with the Everlywell[™] COVID-19 Test Home Collection Kit (no RNase P assay required)
- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 μL of TaqPath[™] COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the dilution created in substep 3a. Mix well, then centrifuge briefly.

Note: The TaqPath[™] COVID-19 Control does not contain the MS2 template.

4. Prepare the Reaction Mix.

a. For each run, combine the following components sufficient for the number of RNA samples, plus one Positive Control per 384-well real-time RT-PCR plate, and one Negative Control from each extraction run.

For example, if RNA samples from 4 extraction runs are being combined on one 384-well real-time RT-PCR plate, then 4 Negative Control wells need to be run on that 384-well real-time RT-PCR plate.

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in this table assume that you extracted sample RNA using an original sample input volume of 200 μ L.

Component	Volume per RNA Sample or control	Volume for nRNA Samples plus yNegative Controls plus 1 Positive Control	Volume for 379 RNA Samples plus 4 Negative Controls plus 1 Positive Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 µL	5.50 x (n + y + 1) μL	2112.0 μL
COVID-19 Real Time PCR Assay Multiplex	1.00 µL	1.10 x (n + y + 1) μL	422.4 µL
Nuclease-free Water	4.00 µL	4.40 x (n + y + 1) μL	1690.0 µL
Total Reaction Mix volume	10.0 μL	_	4224.4 µL

5. Set up the reaction plate:

- a. Pipet 10.0 µL of the Reaction Mix prepared in step 4 into each well of a MicroAmp[™] Optical 384-Well Reaction Plate with Barcode.
 - For other reaction plates that can be used, see "Required materials not supplied" on page 11.
- b. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
- c. Unseal the plate containing the purified sample RNA and Negative Control from the RNA extraction procedure. Add either sample RNA, Negative Control, or Positive Control to each well of the reaction plate according to Table 4 on page 40.

IMPORTANT! To prevent sample contamination, unseal one extraction plate at a time, then reseal it after adding the samples to the RT-PCR reaction plate.

d. Seal the plate thoroughly with MicroAmp[™] Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

- e. Vortex the plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.
 - IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.
- f. Centrifuge the reaction plate for 1–2 minutes at ≥650 x g (≥650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Table 4 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix	10.0 μL	10.0 μL	10.0 μL
Purified sample RNA (from RNA extraction)	10.0 µL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 µL	_
Nuclease-free Water	_	8.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	10.0 μL
Total volume	20.0 μL	20.0 μL	20.0 µL

Prepare the RT-PCR reactions (400-µL sample input, 96-well reaction plate, COVID-19 assay only)

Use this procedure under the following conditions:

- Original sample input volume of 400 µL was used for extraction
- Instrument is compatible with 96-well RT-PCR reaction plates
- Sample was not collected with the Everlywell[™] COVID-19 Test Home Collection Kit (no RNase P assay required)
- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TagPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 μL of TaqPath[™] COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the dilution created in substep 3a. Mix well, then centrifuge briefly.

Note: The TagPath[™] COVID-19 Control does not contain the MS2 template.

4. Prepare the Reaction Mix:

a. For each run, combine the following components sufficient for the number of RNA samples to be tested plus one Positive Control and one Negative Control.

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in this table assume that you extracted sample RNA using an original sample input volume of 400 μ L.

Component	Volume per RNA Sample or control	Volume for n RNA Samples plus 2 controls	Volume for 94 RNA Samples plus 2 controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 µL	6.875 x (<i>n</i> + 2) μL	660 µL
COVID-19 Real Time PCR Assay Multiplex	1.25 μL	1.375 x (n + 2) μL	132 µL
Nuclease-free Water	12.50 µL	13.75 x (n + 2) μL	1320 µL
Total Reaction Mix volume	20.0 μL	_	2112 μL

5. Set up the reaction plate:

a. Pipet 20.0 µL of the Reaction Mix prepared in step 4 into each well of a MicroAmp[™] Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate with Barcode, 0.2 mL.

For other reaction plates that can be used, see "Required materials not supplied" on page 11.

- b. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
- c. Unseal the plate containing the purified sample RNA and Negative Control from the RNA extraction procedure. Add either sample RNA, Negative Control, or Positive Control to each well of the reaction plate according to Table 5 on page 42.
- d. Seal the plate thoroughly with MicroAmp™ Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

e. Vortex the plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.

IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.

f. Centrifuge the reaction plate for 1–2 minutes at \geq 650 × g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Table 5 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix	20.0 μL	20.0 µL	20.0 μL
Purified sample RNA (from RNA extraction)	5.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 μL	_
Nuclease-free Water	_	3.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	5.0 µL
Total volume	25.0 μL	25.0 μL	25.0 µL

Prepare the RT-PCR reactions (400-µL sample input, 384-well reaction plate, COVID-19 assay only)

Use this procedure under the following conditions:

- Original sample input volume of 400 µL was used for extraction
- Instrument is compatible with 384-well RT-PCR reaction plates
- Sample was not collected with the Everlywell[™] COVID-19 Test Home Collection Kit (no RNase P assay required)
- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 μL of TaqPath[™] COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the dilution created in substep 3a. Mix well, then centrifuge briefly.

Note: The TaqPath[™] COVID-19 Control does not contain the MS2 template.



4. Prepare the Reaction Mix.

a. For each run, combine the following components sufficient for the number of RNA samples, plus one Positive Control per 384-well real-time RT-PCR plate, and one Negative Control from each extraction run.

For example, if RNA samples from 4 extraction runs are being combined on one 384-well real-time RT-PCR plate, then 4 Negative Control wells need to be run on that 384-well real-time RT-PCR plate.

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in this table assume that you extracted sample RNA using an original sample input volume of 400 μ L.

Component	Volume per RNA Sample or control	Volume for n RNA Samples plus y Negative Controls plus 1 Positive Control	Volume for 379 RNA Samples plus 4 Negative Controls plus 1 Positive Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 µL	5.50 x (n + y + 1) µL	2112.0 μL
COVID-19 Real Time PCR Assay Multiplex	1.00 μL	1.10 x (n + y + 1) μL	422.4 µL
Nuclease-free Water	9.00 µL	9.90 x (n + y + 1) μL	3802.0 µL
Total Reaction Mix volume	15.0 μL	_	6336.4 µL

5. Set up the reaction plate:

- a. Pipet 15.0 µL of the Reaction Mix prepared in step 4 into each well of a MicroAmp[™] Optical 384-Well Reaction Plate with Barcode.
 - For other reaction plates that can be used, see "Required materials not supplied" on page 11.
- b. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
- c. Unseal the plate containing the purified sample RNA and Negative Control from the RNA extraction procedure. Add either sample RNA, Negative Control, or Positive Control to each well of the reaction plate according to Table 6 on page 44.
 - IMPORTANT! To prevent sample contamination, unseal one extraction plate at a time, then reseal it after adding the samples to the RT-PCR reaction plate.
- d. Seal the plate thoroughly with MicroAmp[™] Optical Adhesive Film.
 - IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

- e. Vortex the plate at the highest setting speed for 10-30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.
 - IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.
- f. Centrifuge the reaction plate for 1–2 minutes at \geq 650 x g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Table 6 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix	15.0 µL	15.0 μL	15.0 µL
Purified sample RNA (from RNA extraction)	5.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 µL	_
Nuclease-free Water	_	3.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	5.0 µL
Total volume	20.0 μL	20.0 μL	20.0 µL

Prepare RT-PCR reactions with 17.5 µL of purified sample RNA (400-µL sample input, 96-well reaction plate, COVID-19 assay only)

IMPORTANT!

- The volumes provided in the TaqPath[™] COVID-19 Combo Kit Advanced (Cat. No. A47813) have been optimized for this procedure.
- The sample volumes used in this procedure are not compatible with samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

Use this procedure to prepare RT-PCR reactions under the following conditions.

- Original sample input volume of 400 µL was used for extraction
- 17.5 µL of purified sample RNA is added to the reaction
- Instrument is compatible with 96-well RT-PCR reaction plates
- Sample was not collected with the Everlywell™ COVID-19 Test Home Collection Kit (no RNase P assay required)

- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 µL of TaqPath™ COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 µL of TaqPath™ COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 µL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 µL of the dilution created in substep 3a. Mix well, then centrifuge briefly.

Note: The TaqPath[™] COVID-19 Control does not contain the MS2 template.

4. Prepare the Reaction Mix:

a. For each run, combine the following components sufficient for the number of RNA samples to be tested plus one Positive Control and one Negative Control.

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in this table assume that you extracted sample RNA using an original sample input volume of 400 μ L.

Component	Volume per RNA Sample or control	Volume for <i>n</i> RNA Samples plus 2 controls	Volume for 94 RNA Samples plus 2 controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 µL	6.875 x (n + 2) μL	660 µL
COVID-19 Assay Multiplex Advanced	1.25 μL	1.375 x (n + 2) μL	132 µL
Total Reaction Mix volume	7.5 μL	_	792 µL

5. Set up the reaction plate:

a. Pipet 7.5 µL of the Reaction Mix prepared in step 4 into each well of a MicroAmp[™] Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate with Barcode, 0.2 mL.

A MicroAmp[™] Fast Optical 96-Well Reaction Plate, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate, 0.2 mL can also be used (see "Required materials not supplied" on page 11).

IMPORTANT! The following plates are not compatible with this procedure:

- MicroAmp[™] EnduraPlate[™] Optical 96-Well Fast Clear Reaction Plates with Barcode
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Fast Clear Reaction Plates
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Clear Reaction Plates with Barcode
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Clear Reaction Plates
- b. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.



- c. Unseal the plate containing the purified sample RNA and Negative Control from the RNA extraction procedure. Add either sample RNA, Negative Control, or Positive Control to each well of the reaction plate according to Table 7 on page 46.
- d. Seal the plate thoroughly with MicroAmp™ Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

- e. Vortex the plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.
 - IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.
- f. Centrifuge the reaction plate for 1–2 minutes at \geq 650 × g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Table 7 Reaction plate volumes

	Volume per reaction		
Component	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix	7.5 μL	7.5 μL	7.5 µL
Purified sample RNA (from RNA extraction)	17.5 µL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 µL	_
Nuclease-free Water		15.5 µL	_
Purified Negative Control (from RNA extraction)	_	_	17.5 μL
Total volume	25.0 μL	25.0 μL	25.0 μL

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Prepare RT-PCR reactions with 14.0 µL of purified sample RNA (400-µL sample input, 384-well reaction plate, COVID-19 assay only)

IMPORTANT!

- The volumes provided in the TaqPath[™] COVID-19 Combo Kit Advanced (Cat. No. A47813) have been optimized for this procedure.
- The sample volumes used in this procedure are not compatible with samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

Use this procedure to prepare RT-PCR reactions under the following conditions

- Original sample input volume of 400 µL was used for extraction
- 14.0 µL of purified sample RNA is added to the reaction
- Instrument is compatible with 384-well RT-PCR reaction plates
- Sample was not collected with the Everlywell[™] COVID-19 Test Home Collection Kit (no RNase P assay required)
- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 µL of TaqPath™ COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 µL of TaqPath™ COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 µL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 µL of the dilution created in substep 3a. Mix well, then centrifuge briefly.

Note: The TagPath[™] COVID-19 Control does not contain the MS2 template.

- Prepare the Reaction Mix.
 - a. For each run, combine the following components sufficient for the number of RNA samples, plus one Positive Control per 384-well real-time RT-PCR plate, and one Negative Control from each extraction run.
 - For example, if RNA samples from 4 extraction runs are being combined on one 384-well real-time RT-PCR plate, then 4 Negative Control wells need to be run on that 384-well real-time RT-PCR plate.



All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in this table assume that you extracted sample RNA using an original sample input volume of $400 \, \mu L$.

Component	Volume per RNA Sample or control	Volume for nRNA Samples plus yNegative Controls plus 1 Positive Control	Volume for 379 RNA Samples plus 4 Negative Controls plus 1 Positive Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 μL	5.50 x (n + y + 1) μL	2112.0 μL
COVID-19 Assay Multiplex Advanced	1.00 μL	1.10 x (n + y + 1) μL	422.4 µL
Total Reaction Mix volume	6.0 μL	-	2534.4 µL

5. Set up the reaction plate:

a. Pipet 6.0 µL of the Reaction Mix prepared in step 4 into each well of a MicroAmp[™] Optical 384-Well Reaction Plate with Barcode.

A MicroAmp[™] Optical 384-Well Reaction Plate can be used (see "Required materials not supplied" on page 11).

IMPORTANT! MicroAmp[™] EnduraPlate[™] Optical 384-Well Clear Reaction Plates with Barcode and MicroAmp[™] EnduraPlate[™] Optical 384-Well Clear Reaction Plates are not compatible with this procedure.

- b. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
- c. Unseal the plate containing the purified sample RNA and Negative Control from the RNA extraction procedure. Add either sample RNA, Negative Control, or Positive Control to each well of the reaction plate according to Table 8 on page 49.

IMPORTANT! To prevent sample contamination, unseal one extraction plate at a time, then reseal it after adding the samples to the RT-PCR reaction plate.

d. Seal the plate thoroughly with MicroAmp™ Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

e. Vortex the plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.

IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.

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f. Centrifuge the reaction plate for 1–2 minutes at \geq 650 x g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Table 8 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix	6.0 µL	6.0 µL	6.0 µL
Purified sample RNA (from RNA extraction)	14.0 µL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 μL	_
Nuclease-free Water	_	12.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	14.0 µL
Total volume	20.0 μL	20.0 µL	20.0 μL



Prepare RT-PCR reactions— COVID-19 assay plus RNase P assay (Everlywell[™] COVID-19 Test Home Collection Kit)

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Protocols in this chapter

This chapter covers preparation of RT-PCR reaction plates for samples that were collected using the Everlywell[™] COVID-19 Test Home Collection Kit. These RT-PCR reactions must contain both the COVID-19 Real Time PCR Assay Multiplex and the TagMan[™] SARS-CoV-2 RNase P Assay.

Note: For samples that were collected using a method other than the Everlywell™ COVID-19 Test Home Collection Kit, see Chapter 4, "Prepare RT-PCR reactions—COVID-19 assay only".

IMPORTANT! Do not combine samples collected using the Everlywell™ COVID-19 Test Home Collection Kit with samples collected via other methods on the same RT-PCR plate. The plate setup and software template files are different.

In this chapter, select the appropriate RT-PCR preparation procedure based on the following criteria:

- The original sample input volume used for RNA extraction (200 μL or 400 μL)
- The size of the RT-PCR reaction plate (96-well or a 384-well)

Guidelines for RT-PCR

IMPORTANT!

- Prepare two RT-PCR reaction plate wells for each sample and Negative Control—one for the COVID-19 assay and one for the RNase P assay—plus a single RT-PCR reaction plate well for the Positive Control.
 - 96-well reaction plates can contain up to 46 samples (2 wells each), one Negative Control (2 wells), and one Positive Control (1 well).
 - 384-well reaction plates can contain up to 189 samples (2 wells each), 2 Negative Controls (one from each extraction plate; 4 wells total), and one Positive Control (1 well).
 - For example plate layouts, see Appendix A, "Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)".
- Prepare the reaction plate on ice and keep it on ice until it is loaded into the real-time PCR instrument.
- Run the plate immediately after preparation. Failure to do so could result in degraded RNA samples.
- To prevent contamination, prepare reagents in a PCR workstation or equivalent amplicon-free area.
 Do not use the same pipette for controls and RNA samples, and always use aerosol barrier pipette tips.
- · Maintain an RNase-free environment.
- Protect assays from light.
- Keep RNA samples and components on ice during use.

Prepare the RT-PCR reactions (200-µL sample input, 96-well reaction plate, includes RNase P assay)

Use this procedure under the following conditions:

- Original sample input volume of 200 µL was used for extraction
- Instrument is compatible with 96-well RT-PCR reaction plates
- Sample was collected with the Everlywell[™] COVID-19 Test Home Collection Kit (RNase P assay required)

IMPORTANT! In the following procedure, each sample or Negative Control on the 96-well extraction plate is added to two wells on the 96-well reaction plate (one for the COVID-19 assay and one for the RNase P assay). For this reason, only half of the extraction plate is used per reaction plate. The extraction plate can be stored at 2–8°C for up to 48 hours.

- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 µL of TaqPath™ COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 µL of TaqPath™ COVID-19 Control. Mix well, then centrifuge briefly.

Prepare the RT-PCR reactions (200-µL sample input, 96-well reaction plate, includes RNase Passay)

b. Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the previous dilution. Mix well, then centrifuge briefly.

Note: The TaqPath[™] COVID-19 Control does not contain the MS2 template.

- 4. Prepare separate reaction mixes for the COVID-19 assay and RNase P assay according to the following tables:
 - a. Prepare sufficient reaction mix based on the number of RNA samples, plus one Positive Control and 2 Negative Controls per reaction plate.

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in these tables assume that you extracted sample RNA using an original sample input volume of 200 μ L.

Table 9 COVID-19 Reaction Mix

Component	Volume per RNA sample or control	Volume for n RNA samples plus 2 controls	Volume for 46 RNA samples plus 2 controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 µL	6.875 x (n + 2) µL	330 µL
COVID-19 Real Time PCR Assay Multiplex	1.25 µL	1.375 x (<i>n</i> + 2) μL	66 µL
Nuclease-free Water	7.50 µL	8.25 x (n + 2) μL	396 µL
Total Reaction Mix volume	15.0 µL	_	792 µL

Table 10 RNase P Reaction Mix

Component	Volume per RNA sample or control	Volume for <i>n</i> RNA samples plus 1 Negative Control	Volume for 46 RNA samples plus 1 Negative Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 µL	6.875 x (n + 1) µL	324 µL
TaqMan [™] SARS-CoV-2 RNase P Assay	1.25 μL	1.375 x (n + 1) μL	65 µL
Nuclease-free Water	7.50 µL	8.25 x (n + 1) μL	388 µL
Total Reaction Mix volume	15.0 μL	_	777 μL

- 5. Add each reaction mix to a reaction plate:
 - a. Starting with row A, pipet 15.0 µL per well of the COVID-19 Reaction Mix into every other row of a MicroAmp™ Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL or a MicroAmp™ Optical 96-Well Reaction Plate with Barcode, 0.2 mL. See Figure 1 on page 53.

A MicroAmp[™] Fast Optical 96-Well Reaction Plate, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate, 0.2 mL can be used (see "Required materials not supplied" on page 11).

IMPORTANT! The following plates are not compatible with this procedure:

- MicroAmp[™] EnduraPlate[™] Optical 96-Well Fast Clear Reaction Plates with Barcode
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Fast Clear Reaction Plates
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Clear Reaction Plates with Barcode
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Clear Reaction Plates
- b. Pipet 15.0 µL per well of the RNase P Reaction Mix into the remaining rows of the plate.

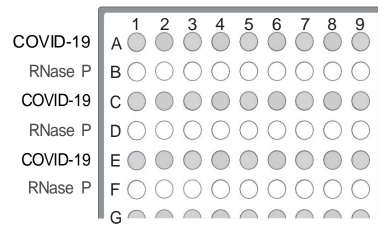


Figure 1 Assays in alternating plate rows

- 6. Add samples and controls to the reaction plate, using the volumes in Table 11 on page 54:
 - a. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
 - b. Unseal the extraction plate, then add the Negative Control from the single well of the extraction plate to wells A1 and B1 of the reaction plate, as shown in Figure 2 on page 54.
 - Add the Positive Control (from step 3) and Nuclease-free Water to an appropriate well of the reaction plate (containing the COVID-19 assay).

There is no RNase P well for the Positive Control.

Prepare the RT-PCR reactions (200-µL sample input, 96-well reaction plate, includes RNase Passay)

d. Add each RNA sample from the extraction plate to adjacent COVID-19 assay and RNase P assay wells of the reaction plate, as shown in the following figure.

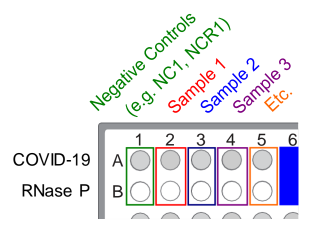


Figure 2 Paired sample and Negative Control wells for each assay

Table 11 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix (COVID-19 or RNase P)	15.0 μL	15.0 µL	15.0 µL
Purified sample RNA (from RNA extraction)	10.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control, from step 3)	_	2.0 μL	_
Nuclease-free Water	_	8.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	10.0 µL
Total volume	25.0 μL	25.0 µL	25.0 µL

- 7. Seal, vortex, and centrifuge the reaction plate:
 - a. Seal the reaction plate thoroughly with MicroAmp[™] Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

b. Vortex the reaction plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.

IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.

c. Centrifuge the reaction plate for 1–2 minutes at ≥650 x g (≥650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Prepare the RT-PCR reactions (200-µL sample input, 384-well reaction plate, includes RNase P assay)

Use this procedure under the following conditions:

- Original sample input volume of 200 µL was used for extraction
- Instrument is compatible with 384-well RT-PCR reaction plates
- Sample was collected with the Everlywell[™] COVID-19 Test Home Collection Kit (RNase P assay required)

Note: You can combine multiple 96-well extraction plates on a single 384-well RT-PCR reaction plate.

- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 µL of TaqPath™ COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 µL of TaqPath™ COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 µL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 µL of the previous dilution. Mix well, then centrifuge briefly.

Note: The TagPath[™] COVID-19 Control does not contain the MS2 template.

- 4. Prepare separate reaction mixes for the COVID-19 assay and RNase P assay according to the following tables:
 - a. Prepare sufficient reaction mix based on the number of RNA samples, plus one Positive Control per 384-well reaction plate, and 2 Negative Controls from each extraction run. For example, if RNA samples from 2 96-well extraction runs are being combined on one 384-well reaction plate, then prepare 4 Negative Control wells on that reaction plate (2 extraction plates x 2 assay types).



Chapter 5 Prepare RT-PCR reactions—COVID-19 assay plus RNase P assay (Everlywell™ COVID-19 Test Home Collection Kit)

Prepare the RT-PCR reactions (200-µL sample input, 384-well reaction plate, includes RNase P assay)

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in these tables assume that you extracted sample RNA using an original sample input volume of 200 μ L.

Table 12 COVID-19 Reaction Mix

Component	Volume per RNA Sample or Control	Volume for nRNA Samples plus yNegative Controls plus 1 Positive Control	Volume for 189 RNA Samples plus 2 Negative Controls plus 1 Positive Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 µL	5.50 x (n + y + 1) μL	1056 µL
COVID-19 Real Time PCR Assay Multiplex	1.00 μL	1.10 x (n + y + 1) μL	211 µL
Nuclease-free Water	4.00 μL	4.40 x (n + y + 1) μL	845 µL
Total Reaction Mix volume	10.0 μL	_	2112 µL

Table 13 RNase P Reaction Mix

Component	Volume per RNA Sample or control	Volume for nRNA Samples plus yNegative Controls	Volume for 189 RNA Samples plus 2 Negative Controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 µL	5.50 x (<i>n</i> + <i>y</i>) µL	1050.5 μL
TaqMan [™] SARS-CoV-2 RNase P Assay	1.00 μL	1.10 x (<i>n</i> + <i>y</i>) µL	210.1 μL
Nuclease-free Water	4.00 μL	4.40 x (<i>n</i> + <i>y</i>) μL	840.4 µL
Total Reaction Mix volume	10.0 μL	_	2101.0 µL

- 5. Add each reaction mix to a reaction plate:
 - a. Starting with row A, pipet 10.0 µL per well of the COVID-19 Reaction Mix into every other row of a MicroAmp[™] Optical 384-Well Reaction Plate with Barcode. See Figure 3 on page 57.
 A MicroAmp[™] Optical 384-Well Reaction Plate can be used (see "Required materials not supplied" on page 11).

IMPORTANT! MicroAmp[™] EnduraPlate[™] Optical 384-Well Clear Reaction Plates with Barcode and MicroAmp[™] EnduraPlate[™] Optical 384-Well Clear Reaction Plates are not compatible with this procedure.

b. Pipet 10.0 µL per well of the RNase P Reaction Mix into the remaining rows of the plate.

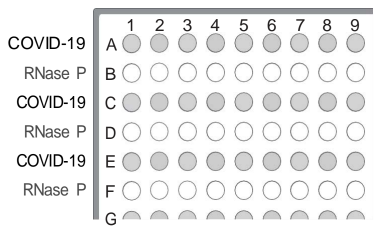


Figure 3 Assays in alternating plate rows

- 6. Add samples and controls to the reaction plate, using the volumes in Table 14 on page 58:
 - a. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.

IMPORTANT! To prevent sample contamination, unseal one extraction plate at a time, then reseal it after adding the samples to the reaction plate.

b. Unseal the extraction plate, then add the first Negative Control from the single well of the extraction plate to wells A1 and B1 of the reaction plate, as shown in Figure 4 on page 58.

Note: If you combine multiple extraction plates on the reaction plate, add the Negative Control from each extraction plate to adjacent COVID-19 assay and RNase P assay wells.

c. Add the Positive Control (from step 3) and Nuclease-free Water to an appropriate well of the reaction plate (containing the COVID-19 assay).

Note:

- There is no RNase P well for the Positive Control.
- Only one Positive Control is required per 384-well plate, even if you combine multiple extraction plates.

Prepare the RT-PCR reactions (200-µL sample input, 384-well reaction plate, includes RNase P assay)

d. Add each RNA sample from the extraction plate to adjacent COVID-19 assay and RNase P assay wells of the reaction plate, as shown in the following figure.

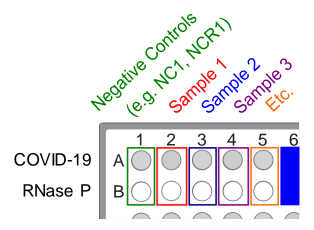


Figure 4 Paired sample and Negative Control wells for each assay

Table 14 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix (COVID-19 or RNase P)	10.0 µL	10.0 μL	10.0 μL
Purified sample RNA (from RNA extraction)	10.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 μL	_
Nuclease-free Water	-	8.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	10.0 μL
Total volume	20.0 μL	20.0 μL	20.0 μL

- 7. Seal, vortex, and centrifuge the reaction plate:
 - a. Seal the reaction plate thoroughly with MicroAmp[™] Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

- b. Vortex the reaction plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.
 - IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.

c. Centrifuge the reaction plate for 1–2 minutes at \geq 650 x g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Prepare the RT-PCR reactions (400-µL sample input, 96-well reaction plate, includes RNase P assay)

Use this procedure under the following conditions:

- Original sample input volume of 400 µL was used for extraction
- Instrument is compatible with 96-well RT-PCR reaction plates
- Sample was collected with the Everlywell[™] COVID-19 Test Home Collection Kit (RNase P assay required)

IMPORTANT! In the following procedure, each sample or Negative Control on the 96-well extraction plate is added to two wells on the 96-well reaction plate (one for the COVID-19 assay and one for the RNase P assay). For this reason, only half of the extraction plate is used per reaction plate. The extraction plate can be stored at 2–8°C for up to 48 hours.

- 1. If frozen, thaw the reagents on ice.
- 2. Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 µL of TaqPath™ COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 µL of TaqPath™ COVID-19 Control. Mix well, then centrifuge briefly.
 - b. Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the previous dilution. Mix well, then centrifuge briefly.

Note: The TaqPath[™] COVID-19 Control does not contain the MS2 template.

- 4. Prepare separate reaction mixes for the COVID-19 assay and RNase P assay according to the following tables:
 - a. Prepare sufficient reaction mix based on the number of RNA samples, plus one Positive Control and 2 Negative Controls per reaction plate.



Chapter 5 Prepare RT-PCR reactions—COVID-19 assay plus RNase P assay (Everlywell™ COVID-19 Test Home Collection Kit)

Prepare the RT-PCR reactions (400-µL sample input, 96-well reaction plate, includes RNase Passay)

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in these tables assume that you extracted sample RNA using an original sample input volume of 400 μ L.

Table 15 COVID-19 Reaction Mix

Component	Volume per RNA Sample or control	Volume for <i>n</i> RNA Samples plus 2 controls	Volume for 46 RNA samples plus 2 controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 μL	6.875 x (<i>n</i> + 2) μL	330 µL
COVID-19 Real Time PCR Assay Multiplex	1.25 µL	1.375 x (<i>n</i> + 2) μL	66 µL
Nuclease-free Water	12.50 µL	13.75 x (n + 2) μL	660 µL
Total Reaction Mix volume	20.0 μL	_	1056 µL

Table 16 RNase P Reaction Mix

Component	Volume per RNA sample or control	Volume for <i>n</i> RNA samples plus 1 Negative Control	Volume for 46 RNA samples plus 1 Negative Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	6.25 μL	6.875 x (<i>n</i> + 1) μL	323.1 μL
TaqMan [™] SARS-CoV-2 RNase P Assay	1.25 µL	1.375 x (n + 1) μL	64.6 µL
Nuclease-free Water	12.50 µL	13.75 x (<i>n</i> + 1) μL	646.3 µL
Total Reaction Mix volume	20.0 μL	_	1034.0 µL

5. Add each reaction mix to a reaction plate:

a. Starting with row A, pipet 20.0 µL per well of the COVID-19 Reaction Mix into every other row of a MicroAmp[™] Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate with Barcode, 0.2 mL. See Figure 5 on page 61.

A MicroAmp[™] Fast Optical 96-Well Reaction Plate, 0.1 mL or a MicroAmp[™] Optical 96-Well Reaction Plate, 0.2 mL can be used (see "Required materials not supplied" on page 11).

IMPORTANT! The following plates are not compatible with this procedure:

- MicroAmp[™] EnduraPlate[™] Optical 96-Well Fast Clear Reaction Plates with Barcode
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Fast Clear Reaction Plates
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Clear Reaction Plates with Barcode
- MicroAmp[™] EnduraPlate[™] Optical 96-Well Clear Reaction Plates

b. Pipet 20.0 µL per well of the RNase P Reaction Mix into the remaining rows of the plate.

Prepare the RT-PCR reactions (400-µL sample input, 96-well reaction plate, includes RNase P assay)

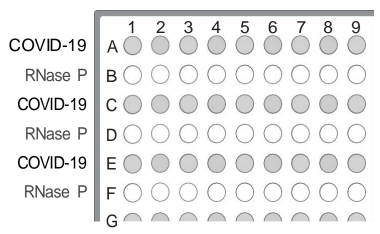


Figure 5 Assays in alternating plate rows

- 6. Add samples and controls to the reaction plate, using the volumes in Table 17 on page 62:
 - a. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.
 - b. Unseal the extraction plate, then add Negative Control from the single well of the extraction plate to wells A1 and B1 of the reaction plate, as shown in Figure 6 on page 61.
 - Add the Positive Control (from step 3) and Nuclease-free Water to an appropriate well of the reaction plate (containing the COVID-19 assay).

There is no RNase P well for the Positive Control.

 d. Add each RNA sample from the extraction plate to adjacent COVID-19 assay and RNase P assay wells of the reaction plate, as shown in the following figure.

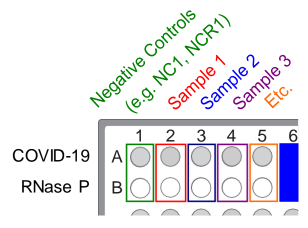


Figure 6 Paired sample and Negative Control wells for each assay



Chapter 5 Prepare RT-PCR reactions—COVID-19 assay plus RNase P assay (Everlywell™ COVID-19 Test Home Collection Kit)

Prepare the RT-PCR reactions (400-µL sample input, 384-well reaction plate, includes RNase P assay)

Table 17 Reaction plate volumes

	Volume per reaction		
Component	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix (COVID-19 or RNase P)	20.0 μL	20.0 μL	20.0 μL
Purified sample RNA (from RNA extraction)	5.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 μL	_
Nuclease-free Water	_	3.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	5.0 µL
Total volume	25.0 μL	25.0 µL	25.0 μL

- 7. Seal, vortex, and centrifuge the reaction plate:
 - a. Seal the reaction plate thoroughly with MicroAmp[™] Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

- b. Vortex the reaction plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.
 - IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.
- c. Centrifuge the reaction plate for 1–2 minutes at ≥650 x g (≥650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.

Prepare the RT-PCR reactions (400-µL sample input, 384-well reaction plate, includes RNase P assay)

Use this procedure under the following conditions:

- Original sample input volume of 400 µL was used for extraction
- Instrument is compatible with 384-well RT-PCR reaction plates
- Sample was collected with the Everlywell[™] COVID-19 Test Home Collection Kit (RNase P assay required)

Note: You can combine multiple 96-well extraction plates on a single 384-well RT-PCR reaction plate.

- 1. If frozen, thaw the reagents on ice.
- Gently vortex the reagents, then centrifuge briefly to collect liquid at the bottom of the tube.
- 3. Dilute TaqPath[™] COVID-19 Control (1 x 10⁴ copies/µL) to a working stock of 25 copies/µL:
 - a. Pipet 98 µL of TaqPath™ COVID-19 Control Dilution Buffer into a microcentrifuge tube, then add 2 µL of TaqPath™ COVID-19 Control. Mix well, then centrifuge briefly.

Prepare the RT-PCR reactions (400-µL sample input, 384-well reaction plate, includes RNase P assay)

b. Pipet 87.5 μL of TaqPath[™] COVID-19 Control Dilution Buffer into a second microcentrifuge tube, then add 12.5 μL of the previous dilution. Mix well, then centrifuge briefly.

Note: The TaqPath[™] COVID-19 Control does not contain the MS2 template.

- 4. Prepare separate reaction mixes for the COVID-19 assay and RNase P assay according to the following tables:
 - a. Prepare sufficient reaction mix based on the number of RNA samples, plus one Positive Control per 384-well reaction plate, and 2 Negative Controls from each extraction run. For example, if RNA samples from 2 96-well extraction runs are being combined on one 384-well reaction plate, then prepare 4 Negative Control wells on that reaction plate (2 extraction plates x 2 assay types).

All volumes include 10% overage for pipette error.

IMPORTANT! The volumes in these tables assume that you extracted sample RNA using an original sample input volume of 400 µL.

Table 18 COVID-19 Reaction Mix

Component	Volume per RNA Sample or control	Volume for n RNA Samples plus y Negative Controls plus 1 Positive Control	Volume for 189 RNA Samples plus 2 Negative Controls plus 1 Positive Control
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 μL	5.50 x (n + y + 1) µL	1056 μL
COVID-19 Real Time PCR Assay Multiplex	1.00 μL	1.10 x (n + y + 1) μL	211 μL
Nuclease-free Water	9.00 µL	9.90 x (n + y + 1) µL	1901 µL
Total Reaction Mix volume	15.0 μL	_	3168 µL

Chapter 5 Prepare RT-PCR reactions—COVID-19 assay plus RNase P assay (Everlywell™ COVID-19 Test Home Collection Kit)

Prepare the RT-PCR reactions (400-µL sample input, 384-well reaction plate, includes RNase P assay)

Table 19 RNase P Reaction Mix

Component	Volume per RNA Sample or control	Volume for <i>n</i> RNA Samples plus <i>y</i> Negative Controls	Volume for 189 RNA Samples plus 2 Negative Controls
TaqPath [™] 1-Step Multiplex Master Mix (No ROX [™]) (4X)	5.00 μL	5.50 x (<i>n</i> + <i>y</i>) μL	1050.5 μL
TaqMan [™] SARS-CoV-2 RNase P Assay	1.00 µL	1.10 x (<i>n</i> + <i>y</i>) µL	210.1 μL
Nuclease-free Water	9.00 µL	9.90 x (n + y + 1) μL	1890.9 µL
Total Reaction Mix volume	15.0 µL	_	3151.5 μL

- 5. Add each reaction mix to a reaction plate:
 - a. Starting with row A, pipet 15.0 µL per well of the COVID-19 Reaction Mix into every other row of a MicroAmp[™] Optical 384-Well Reaction Plate with Barcode. See Figure 7 on page 64. A MicroAmp[™] Optical 384-Well Reaction Plate can be used (see "Required materials not supplied" on page 11).

IMPORTANT! MicroAmp[™] EnduraPlate[™] Optical 384-Well Clear Reaction Plates with Barcode and MicroAmp[™] EnduraPlate[™] Optical 384-Well Clear Reaction Plates are not compatible with this procedure.

b. Pipet 15.0 µL per well of the RNase P Reaction Mix into the remaining rows of the plate.

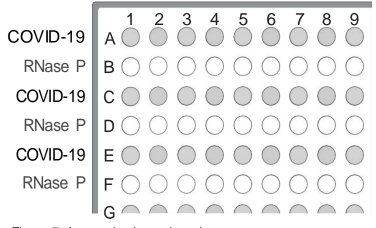


Figure 7 Assays in alternating plate rows

- 6. Add samples and controls to the reaction plate, using the volumes in Table 20 on page 66:
 - a. Gently vortex the sealed plate containing the purified sample RNA and Negative Control from the RNA extraction procedure, then centrifuge briefly to collect liquid at the bottom of the plate.

IMPORTANT! To prevent sample contamination, unseal one extraction plate at a time, then reseal it after adding the samples to the reaction plate.

b. Unseal the extraction plate, then add the Negative Control from the single well of the extraction plate to wells A1 and B1 of the reaction plate, as shown in Figure 8 on page 65.

Note: If you combine multiple extraction plates on the reaction plate, add the Negative Control from each extraction plate to adjacent COVID-19 assay and RNase P assay wells.

c. Add the Positive Control (from step 3) and Nuclease-free Water to an appropriate well of the reaction plate (containing the COVID-19 assay).

Note:

- There is no RNase P well for the Positive Control.
- Only one Positive Control is required per 384-well plate, even if you combine multiple extraction plates.
- d. Add each RNA sample from the extraction plate to adjacent COVID-19 assay and RNase P assay wells of the reaction plate, as shown in the following figure.

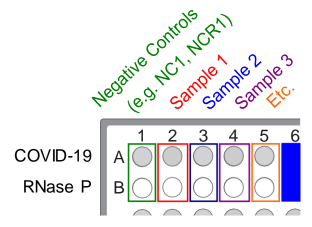


Figure 8 Paired sample and Negative Control wells for each assay



Chapter 5 Prepare RT-PCR reactions—COVID-19 assay plus RNase P assay (Everlywell™ COVID-19 Test Home Collection Kit)

Prepare the RT-PCR reactions (400-µL sample input, 384-well reaction plate, includes RNase P assay)

Table 20 Reaction plate volumes

Component	Volume per reaction		
	RNA Sample reaction	Positive Control reaction	Negative Control reaction
Reaction Mix (COVID-19 or RNase P)	15.0 µL	15.0 μL	15.0 μL
Purified sample RNA (from RNA extraction)	5.0 μL	_	_
Positive Control (diluted TaqPath™ COVID-19 Control from step 3)	_	2.0 μL	_
Nuclease-free Water	-	3.0 µL	_
Purified Negative Control (from RNA extraction)	_	_	5.0 µL
Total volume	20.0 μL	20.0 μL	20.0 μL

- 7. Seal, vortex, and centrifuge the reaction plate:
 - a. Seal the reaction plate thoroughly with MicroAmp[™] Optical Adhesive Film.

IMPORTANT! When applying the MicroAmp[™] Optical Adhesive Film, ensure that pressure is applied across the entire plate and that there is a tight seal across every individual well. Failure to do so runs the risk of an improperly sealed well, leading to potential well-to-well contamination during vortexing and evaporation during PCR.

- b. Vortex the reaction plate at the highest setting speed for 10–30 seconds with medium pressure. Move the plate around to ensure equal contact on the vortex mixer platform.
 - IMPORTANT! Vortex for 10–30 seconds to ensure proper mixing. Failure to do so might result in false classification of samples.
- c. Centrifuge the reaction plate for 1–2 minutes at \geq 650 x g (\geq 650 RCF) to remove bubbles and to collect the liquid at the bottom of the reaction plate.



Perform RT-PCR using the Applied Biosystems[™] 7500 Fast Dx Real-Time PCR Instrument

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Dye calibration for the 7500 Real-Time PCR Instrument series

A maintained instrument will be calibrated for many dyes. In addition to those dyes, the instrument operator must calibrate the instrument for $ABY^{\text{\tiny TM}}$ dye and $JUN^{\text{\tiny TM}}$ dye that are used with this kit. For all other assays, refer to the standard calibration process.

Transfer the template (SDT) file for the 7500 Fast Dx Real-Time PCR Instrument

The template (SDT) file contains the settings for the instrument run. It is installed on the computer with Applied Biosystems™ COVID-19 Interpretive Software, and must be transferred via a USB drive or other method to the computer on which SDS Software v1.4.1 is installed.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 1. On the computer with Applied Biosystems[™] COVID-19 Interpretive Software, navigate to one of the following directories, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell[™] COVID-19 Test Home Collection Kit samples):
 - <...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User Documents\COVID-19
 - ...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User
 Documents\COVID-19 + RNase P
 - <...> is the installation directory



2. Select the correct SDT file, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):

Assay type	Template name		
COVID-19 only	TaqPath COVID-19 Kit Template 7500fastDx sds1_4_1 v1-4.sdt		
	TaqPath COVID-19 and RNaseP Template 7500fastDx sdsl_4_1 v1-0.sdt		

Transfer the SDT file to the computer with SDS Software v1.4.1, using a USB drive or other method.

Set up and run the 7500 Fast Dx Real-Time PCR Instrument —COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay.

For plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the 7500 Fast Dx Real-Time PCR Instrument—COVID-19 plus RNase P assay" on page 70.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

1. Using SDS Software v1.4.1, open the SDT file that you transferred in "Transfer the template (SDT) file for the 7500 Fast Dx Real-Time PCR Instrument" on page 67.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 2. Confirm the run settings in the template and adjust as necessary.
 - Assay: Standard Curve (Absolute Quantitation)
 - Run mode: Standard 7500Passive reference: None

Sample volume: 25 µL

IMPORTANT! The passive reference must be set to None.

3. Confirm that the reporter dye and the detector pairs are correct in the **Detector Manager** in the **Tools** menu.

Reporter dye	Detector
FAM	ORF1ab
VIC	N gene
ABY	S gene
JUN	MS2



- 4. Confirm that the targets above are assigned to each well in the plate layout.
- 5. Confirm the labeling of the control wells.
 - The template has one positive control and one negative control assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 6. For wells with a positive control, confirm that **Task** is set to **Standard**.
- 7. For wells with a negative control, confirm that **Task** is set to **NTC**.
- 8. Edit the plate layout to assign a unique sample name to each well in the physical plate. For wells with a patient sample, confirm that **Task** is set to **Unknown** for all detectors.

Note: Wells that do not have a sample name will not be analyzed by the software.

9. Confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	70

- 10. Click Save As, enter a file name, then click Save.
- 11. Reopen the file to connect the computer to the instrument, load the plate, then start the run on the real-time PCR instrument.
- 12. After the instrument run is complete, open the SDS file in SDS Software v1.4.1. Analyze, then save the file.

Set up and run the 7500 Fast Dx Real-Time PCR Instrument —COVID-19 plus RNase P assay

This section covers instrument set-up and operation for reaction plates that contain both the COVID-19 assay and the RNase P assay. Both assays are required for samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

For plates that contain only the COVID-19 assay, see "Set up and run the 7500 Fast Dx Real-Time PCR Instrument—COVID-19 assay only" on page 68.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

1. Using SDS Software v1.4.1, open the SDT file that you transferred in "Transfer the template (SDT) file for the 7500 Fast Dx Real-Time PCR Instrument" on page 67.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 2. Confirm the run settings in the template and adjust as necessary.
 - Assay: Standard Curve (Absolute Quantitation)

Run mode: Standard 7500
Passive reference: None
Sample volume: 25 µL

IMPORTANT! The passive reference must be set to None.

Confirm that the reporter dye and the detector pairs are correct in the **Detector Manager** in the **Tools** menu.

Reporter dye	Detector
COVID-19 assay	
FAM	ORF1ab
VIC	N gene
ABY	S gene
JUN	MS2
RNase P assay	
VIC	RNaseP

Confirm that the detectors above are assigned to the appropriate wells in the plate layout.

Note:

- Wells containing the RNase P assay have only the RNaseP detector assigned.
- Wells containing the COVID-19 assay have all other detectors assigned.



- 5. Confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned to wells.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 6. For wells with a positive control, confirm that **Task** is set to **Standard**.
- 7. For wells with a negative control, confirm that **Task** is set to **NTC**.
- 8. Edit the plate layout to assign the same sample name to each paired COVID-19 assay well and RNase P assay well in the physical plate.

IMPORTANT! Paired sample assay wells that do not have the same sample name will not be analyzed by the software.

For wells with a patient sample, confirm that **Task** is set to **Unknown** for all detectors.

9. Confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	70

- 10. Click **Save As**, enter a file name, then click **Save**.
- 11. Reopen the file to connect the computer to the instrument, load the plate, then start the run on the instrument.
- 12. After the instrument run is complete, open the SDS file in SDS Software v1.4.1. Analyze, then save the file.



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Dye calibration for the 7500 Real-Time PCR Instrument series

A maintained instrument will be calibrated for many dyes. In addition to those dyes, the instrument operator must calibrate the instrument for ABY^{TM} dye and JUN^{TM} dye that are used with this kit. For all other assays, refer to the standard calibration process.



Transfer the template (SDT or EDT) file for the 7500 Fast Real-Time PCR Instrument

The template (SDT or EDT) file contains the settings for the instrument run. It is installed on the computer with Applied Biosystems[™] COVID-19 Interpretive Software, and must be transferred via a USB drive or other method to the computer on which instrument data collection software is installed.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 1. On the computer with Applied Biosystems™ COVID-19 Interpretive Software, navigate to one of the following directories, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):
 - <...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User Documents\COVID-19
 - ...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User
 Documents\COVID-19 + RNase P
 - <...> is the installation directory
- 2. Select the correct template file, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):

Table 21 COVID-19 assay only

Data collection software version	Template file	
SDS Software v1.5.1	TaqPath COVID-19 Kit Template 7500fast sds1_5_1 v1-4.sdt	
7500 Software v2.3	TaqPath COVID-19 Kit Template 7500fast sds2_3 v1-2.edt	

Table 22 COVID-19 assay and RNase P assay

Data collection software version	Template file
SDS Software v1.5.1	TaqPath COVID-19 and RNaseP Template 7500fast sds1_5_1 v1-0.sdt
7500 Software v2.3	TaqPath COVID-19 and RNaseP Template 7500fast sds2_3 v1-0.edt

3. Select the correct SDT or EDT file for the version of the instrument software that you are using:

Data collection software version	Template file	
SDS Software v1.5.1	TaqPath COVID-19 Kit Template 7500fast sds1_5_1 v1-4.sdt	
7500 Software v2.3	TaqPath COVID-19 Kit Template 7500fast sds2_3 v1-2.edt	

4. Transfer the appropriate SDT or EDT file to the computer with your data collection software, using a USB drive or other method.

Set up and run the 7500 Fast Real-Time PCR Instrument (SDS Software v1.5.1)—COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay, and is specific for the 7500 Fast Real-Time PCR Instrument using SDS Software v1.5.1.

For plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the 7500 Fast Real-Time PCR Instrument (SDS Software v1.5.1)—COVID-19 plus RNase P assay" on page 77.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. Using SDS Software v1.5.1, access the appropriate template file.
 - a. Create a new experiment.
 - b. In the **Template** field, browse to, then open the SDT file that you transferred in "Transfer the template (SDT or EDT) file for the 7500 Fast Real-Time PCR Instrument" on page 73.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 2. Confirm the run settings in the template and adjust as necessary.
 - Assay: Standard Curve (Absolute Quantitation)

Run mode: Standard 7500
 Passive reference: None
 Sample volume: 25 µL

IMPORTANT! The passive reference must be set to **None**.



3. Confirm that the reporter dye and the detector pairs are correct in the **Detector Manager** in the **Tools** menu.

Reporter dye	Detector
FAM	ORF1ab
VIC	N gene
ABY	S gene
JUN	MS2

- 4. Confirm that the targets above are assigned to each well in the plate layout.
- 5. Confirm the labeling of the control wells.
 - The template has one positive control (PC) and one negative control (NC) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 6. For wells with a positive control, confirm that **Task** is set to **Standard**.
- 7. For wells with a negative control, confirm that **Task** is set to **NTC**.
- 8. Edit the plate layout to assign a unique sample name to each well in the physical plate. For wells with a patient sample, ensure that **Task** is set to **Unknown** for all detectors.

Note: Wells that do not have a sample name will not be analyzed by the software.

9. Confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	70

- 10. Click Save As, enter a file name, then click Save.
- 11. Reopen the file, load the plate, then start the run on the instrument.
- 12. After the instrument run is complete, open the SDS file in SDS Software v1.5.1. Analyze, then save the file.

Set up and run the 7500 Fast Real-Time PCR Instrument (7500 Software v2.3)—COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay, and is specific for the 7500 Fast Real-Time PCR Instrument using SDS Software v2.3.

For plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the 7500 Fast Real-Time PCR Instrument (7500 Software v2.3)—COVID-19 plus RNase P assay" on page 79.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the 7500 Software v2.3 home page, click **Template**.
- 2. Browse to, then open the EDT file that you transferred in "Transfer the template (SDT or EDT) file for the 7500 Fast Real-Time PCR Instrument" on page 73.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Experiment Properties** window, enter or confirm the following information:
 - Experiment name: Enter a unique name
 - Instrument type: 7500 Fast (96 wells)
 - Type of experiment: Quantitation Standard Curve
 - Reagents: TaqMan[™]
 - Ramp Speed: Standard
- 4. In the **Plate Setup** window, in the **Define Targets and Samples** tab and the **Define Targets** pane, confirm that the targets, reporter dyes, and quenchers are listed correctly.

Target	Reporter dye	Quencher
ORF1ab	FAM	None
N gene	VIC	None
S gene	ABY	None
MS2	JUN	None

- 5. In the Plate Setup window, in the Define Targets and Samples tab and the Define Samples pane, confirm the labeling of the control wells.
 - The template has one positive control (PC) and one negative control (NC) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
 - To include additional controls, select Add New Sample.
- 6. Select Add New Sample to assign a unique sample name to each well in the physical plate.



7. In the **Plate Setup** window, in the **Assign Targets and Samples** tab, confirm that four targets are assigned to each well in the plate layout. To designate a target or sample to a well, select the well, then check the **Assign** checkbox.

Note: Wells that do not have a sample name will not be analyzed by the software.

- 8. For wells with a positive control, confirm that **Task** is set to **S** for Standard.
- 9. For wells with a negative control, confirm that **Task** is set to **N** for Negative.
- 10. For wells with a patient sample, confirm that **Task** is set to **U** for Unknown.
- 11. Confirm that the Passive Reference is set to None.
- In the Run Method window, confirm that the Reaction Volume Per Well is 25 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	40

- 13. Select **Start Run**, enter a file name, then click **Save**.
- 14. After the instrument run is complete, click **Analyze**, then save the file.

Set up and run the 7500 Fast Real-Time PCR Instrument (SDS Software v1.5.1)—COVID-19 plus RNase P assay

This section covers instrument set-up and operation for reaction plates that contain both the COVID-19 assay and the RNase P assay, and is specific for the 7500 Fast Real-Time PCR Instrument using SDS Software v1.5.1.

Both assays are required for samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

For plates that contain only the COVID-19 assay, see "Set up and run the 7500 Fast Real-Time PCR Instrument (SDS Software v1.5.1)—COVID-19 assay only" on page 74.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. Using SDS Software v1.5.1, access the appropriate template file.
 - a. Create a new experiment.

b. In the **Template** field, browse to, then open the SDT file that you transferred in "Transfer the template (SDT or EDT) file for the 7500 Fast Real-Time PCR Instrument" on page 73.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

2. Confirm the run settings in the template and adjust as necessary.

Assay: Standard Curve (Absolute Quantitation)

Run mode: Standard 7500
Passive reference: None
Sample volume: 25 µL

IMPORTANT! The passive reference must be set to None.

Confirm that the reporter dye and the detector pairs are correct in the **Detector Manager** in the **Tools** menu.

Reporter dye	Detector		
COVID-19 assay			
FAM	ORF1ab		
VIC	N gene		
ABY	S gene		
JUN	MS2		
RNase P assay			
VIC	RNaseP		

4. Confirm that the detectors above are assigned to each well in the plate layout.

Note:

- Wells containing the RNase P assay have only the RNaseP detector assigned.
- · Wells containing the COVID-19 assay have all other detectors assigned.
- 5. Confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 6. For wells with a positive control, confirm that **Task** is set to **Standard**.
- 7. For wells with a negative control, confirm that **Task** is set to **NTC**.

8. Edit the plate layout to assign the same sample name to each paired COVID-19 assay well and RNase P assay well in the physical plate.

IMPORTANT! Paired sample assay wells that do not have the same sample name will not be analyzed by the software.

For wells with a patient sample, ensure that **Task** is set to **Unknown** for all detectors.

9. Confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	40

- 10. Click Save As, enter a file name, then click Save.
- 11. Reopen the file, load the plate, then start the run on the instrument.
- 12. After the instrument run is complete, open the SDS file in SDS Software v1.5.1. Analyze, then save the file.

Set up and run the 7500 Fast Real-Time PCR Instrument (7500 Software v2.3)—COVID-19 plus RNase P assay

This section covers instrument set-up and operation for reaction plates that contain both the COVID-19 assay and the RNase P assay, and is specific for the 7500 Fast Real-Time PCR Instrument using SDS Software v2.3.

Both assays are required for samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit.

For plates that contain only the COVID-19 assay, see "Set up and run the 7500 Fast Real-Time PCR Instrument (7500 Software v2.3)—COVID-19 assay only" on page 76.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the 7500 Software v2.3 home page, click **Template**.
- 2. Browse to, then open the EDT file that you transferred in "Transfer the template (SDT or EDT) file for the 7500 Fast Real-Time PCR Instrument" on page 73.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Experiment Properties** window, enter or confirm the following information:
 - Experiment name: Enter a unique name
 - Instrument type: 7500 Fast (96 wells)
 - Type of experiment: Quantitation Standard Curve
 - Reagents: TaqMan[™]
 - Ramp Speed: Standard
- 4. In the **Plate Setup** window, in the **Define Targets and Samples** tab and the **Define Targets** pane, confirm that the targets, reporter dyes, and quenchers are listed correctly.

Target	Reporter dye	Quencher		
COVID-19 assaytargets				
ORF1ab	FAM	None		
N gene	VIC	None		
S gene	ABY	None		
MS2	JUN	None		
RNase P assaytarget				
RNaseP	VIC	None		

- 5. In the Plate Setup window, in the Define Targets and Samples tab and the Define Samples pane, confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
 - To include additional controls, select Add New Sample.
- 6. Select **Add New Sample** to create a unique sample name.
- 7. In the Plate Setup window, in the Assign Targets and Samples tab, confirm the following:
 - Wells containing the RNase P assay have only the RNaseP target assigned.
 - Wells containing the COVID-19 assay have all other targets assigned.
- 8. To assign a target or sample to a well, select the well, then check the **Assign** checkbox.

IMPORTANT! Paired sample assay wells that do not have the same sample name will not be analyzed.

- 9. For wells with a positive control, confirm that **Task** is set to **S** for Standard.
- 10. For wells with a negative control, confirm that **Task** is set to **N** for Negative.
- 11. For wells with a patient sample, confirm that **Task** is set to **U** for Unknown.
- 12. Confirm that the **Passive Reference** is set to **None**.



13. In the **Run Method** window, confirm that the **Reaction Volume Per Well** is 25 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	40

- 14. Select Start Run, enter a file name, then click Save.
- 15. After the instrument run is complete, click **Analyze**, then save the file.



Perform RT-PCR using the Applied Biosystems[™] 7500 Real-Time PCR Instrument

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Dye calibration for the 7500 Real-Time PCR Instrument series

A maintained instrument will be calibrated for many dyes. In addition to those dyes, the instrument operator must calibrate the instrument for ABY^{TM} dye and JUN^{TM} dye that are used with this kit. For all other assays, refer to the standard calibration process.

IMPORTANT! Use only the calibration plates listed in "Required materials not supplied" on page 11.



Transfer the template (EDT) file for the 7500 Real-Time PCR Instrument

The template (EDT) file contains the settings for the instrument run. It is installed on the computer with Applied Biosystems™ COVID-19 Interpretive Software, and must be transferred via a USB drive or other method to the computer on which 7500 Software ∨2.3 is installed.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 1. On the computer with Applied Biosystems™ COVID-19 Interpretive Software, navigate to one of the following directories, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):
 - <...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User Documents\COVID-19
 - ...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User
 Documents\COVID-19 + RNase P
 - <...> is the installation directory
- 2. Select the correct EDT file, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):

Assay type	Template name	
COVID-19 only	TaqPath COVID-19 Kit Template 7500std sds2_3 v1-2.edt	
COVID-19 and RNase P	TaqPath COVID-19 and RNaseP Template 7500std sds2_3 v1-0.edt	

Transfer the EDT file to the computer with your data collection software, using a USB drive or other method.

Set up and run the 7500 Real-Time PCR Instrument—COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay.

For plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the 7500 Fast Real-Time PCR Instrument (7500 Software v2.3)—COVID-19 plus RNase P assay" on page 79.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the 7500 Software v2.3 home page, click **Template**.
- 2. Browse to, then open the template file that you transferred in "Transfer the template (EDT) file for the 7500 Real-Time PCR Instrument" on page 83.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Experiment Properties** window, enter or confirm the following information:
 - Experiment name: Enter a unique name
 - Instrument type: 7500 (96 wells)
 - Type of experiment: Quantitation Standard Curve
 - Reagents: TaqMan[™]
 - Ramp Speed: Standard
- 4. In the **Plate Setup** window, in the **Define Targets and Samples** tab and the **Define Targets** pane, confirm that the targets, reporter dyes, and quenchers are listed correctly.

Target	Reporter dye	Quencher
ORF1ab	FAM	None
N gene	VIC	None
S gene	ABY	None
MS2	JUN	None

- 5. In the Plate Setup window, in the Define Targets and Samples tab and the Define Samples pane, confirm the labeling of the control wells.
 - The template has one positive control (PC) and one negative control (NC) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.

To include additional controls, select Add New Sample.

6. Select Add New Sample to assign a unique sample name to each well in the physical plate.

7. In the **Plate Setup** window, in the **Assign Targets and Samples** tab, confirm that four targets are assigned to each well in the plate layout. To designate a target or sample to a well, select the well, then check the **Assign** checkbox.

Note: Wells that do not have a sample name will not be analyzed by the software.

- 8. For wells with a positive control, confirm that **Task** is set to **S** for Standard.
- 9. For wells with a negative control, confirm that **Task** is set to **N** for Negative.
- 10. For wells with a patient sample, confirm that **Task** is set to **U** for Unknown.
- 11. Confirm that the **Passive Reference** is set to **None**.
- 12. In the **Run Method** window, confirm that the **Reaction Volume Per Well** is 25 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	40

- 13. Select **Start Run**, enter a file name, then click **Save**.
- 14. After the instrument run is complete, click **Analyze**, then save the file.

Set up and run the 7500 Real-Time PCR Instrument—COVID-19 and RNase P assay

This section covers instrument set-up and operation for reaction plates that contain both the COVID-19 assay and the RNase P assay. Both assays are required for samples collected using the Everlywell[™] COVID-19 Test Home Collection Kit.

For plates that contain only the COVID-19 assay, see "Set up and run the 7500 Real-Time PCR Instrument—COVID-19 assay only" on page 84.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the 7500 Software v2.3 home page, click **Template**.
- 2. Browse to, then open the template file that you transferred in "Transfer the template (EDT) file for the 7500 Real-Time PCR Instrument" on page 83.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Experiment Properties** window, enter or confirm the following information:
 - Experiment name: Enter a unique name
 - Instrument type: 7500 (96 wells)
 - Type of experiment: Quantitation Standard Curve
 - Reagents: TaqMan[™]
 - Ramp Speed: Standard
- 4. In the **Plate Setup** window, in the **Define Targets and Samples** tab and the **Define Targets** pane, confirm that the targets, reporter dyes, and quenchers are listed correctly.

Target	Reporter dye	Quencher		
COVID-19 assaytargets				
ORF1ab	FAM	None		
N gene	VIC	None		
S gene	ABY	None		
MS2	JUN	None		
RNase P assaytarget				
RNaseP	VIC	None		

- 5. In the Plate Setup window, in the Define Targets and Samples tab and the Define Samples pane, confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
 - To include additional controls, select Add New Sample.
- 6. Select **Add New Sample** to create a unique sample name.
- 7. In the Plate Setup window, in the Assign Targets and Samples tab, confirm the following:
 - Wells containing the RNase P assay have only the RNaseP target assigned.
 - Wells containing the COVID-19 assay have all other targets assigned.
- 8. To assign a target or sample to a well, select the well, then check the **Assign** checkbox.

IMPORTANT! Each paired COVID-19 assay well and RNase P assay well must have the same sample name, or they will not be correctly analyzed by the software.

- 9. For wells with a positive control, confirm that **Task** is set to **S** for Standard.
- 10. For wells with a negative control, confirm that **Task** is set to **N** for Negative.
- 11. For wells with a patient sample, confirm that **Task** is set to **U** for Unknown.
- 12. Confirm that the **Passive Reference** is set to **None**.

13. In the **Run Method** window, confirm that the **Reaction Volume Per Well** is 25 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	40

- 14. Select Start Run, enter a file name, then click Save.
- 15. After the instrument run is complete, click **Analyze**, then save the file.



Perform RT-PCR using the Applied Biosystems[™] QuantStudio[™] 5 Real-Time PCR Instrument

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Dye calibration for the QuantStudio[™] 5 Real-Time PCR Instrument

A maintained instrument will be calibrated for all dyes that are used with this kit. Ensure that the calibrations for FAM[™] dye, VIC[™] dye, ABY[™] dye, and JUN[™] dye are current. If calibration is required, refer to the standard calibration process in the instrument user guide.

Transfer the template (EDT) file for the QuantStudio[™] 5 Real-Time PCR Instrument

The template (EDT) file contains the settings for the instrument run. It is installed on the computer with Applied Biosystems[™] COVID-19 Interpretive Software, and must be transferred via a USB drive or other method to the computer on which QuantStudio[™] Design and Analysis Desktop Software v1.5.1 is installed.

IMPORTANT! Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

- 1. On the computer with Applied Biosystems™ COVID-19 Interpretive Software, navigate to one of the following directories, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):
 - <...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User Documents\COVID-19
 - ...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User
 Documents\COVID-19 + RNase P
 - <...> is the installation directory
- 2. Select the correct template file, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):

Table 23 COVID-19 assay only

Instrument	Template file
QuantStudio [™] 5 Real-Time PCR Instrument (96-well, 0.1-mL block)	TaqPath COVID-19 Kit Template QS5 0_1ml_da1_5_1 v2-2.edt
QuantStudio [™] 5 Real-Time PCR Instrument (96-well, 0.2-mL block)	TaqPath COVID-19 Kit Template QS5 0_2ml_da1_5_1 v2-2.edt
QuantStudio [™] 5 Real-Time PCR Instrument (384-well block)	TaqPath COVID-19 Kit Template QS5 384_da1_5_1 v2-2.edt



Table 24 COVID-19 assay and RNase P assay

Instrument	Template file
QuantStudio [™] 5 Real-Time PCR Instrument (96-well, 0.1-mL block)	TaqPath COVID-19 and RNaseP Template QS5 0_1ml_da1_5_1 v1-0.edt
QuantStudio [™] 5 Real-Time PCR Instrument (96-well, 0.2-mL block)	TaqPath COVID-19 and RNaseP Template QS5 0_2ml_da1_5_1 v1-0.edt
QuantStudio [™] 5 Real-Time PCR Instrument (384-well block)	TaqPath COVID-19 and RNaseP Template QS5 384_da1_5_1 v1-0.edt

3. Transfer the EDT file to the computer with QuantStudio[™] Design and Analysis Desktop Software v1.5.1, using a USB drive or other method.

Set up and run the QuantStudio[™] 5 Real-Time PCR Instrument (96-well plates)—COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay.

For 96-well plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the QuantStudio™ 5 Real-Time PCR Instrument (96-well plates)—COVID-19 and RNase P assay" on page 94.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the QuantStudio[™] Design and Analysis Desktop Software v1.5.1, in the **New Experiment** box, select **Create New Experiment** ▶ **Template**.
- Browse to, then open the EDT file that you transferred in "Transfer the template (EDT) file for the QuantStudio™ 5 Real-Time PCR Instrument" on page 89.

IMPORTANT! Be careful to select the appropriate template file for your instrument and block type. Failure to do so can cause errors in the analysis.

Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Properties** tab, enter or confirm the following.
 - Name: Enter a unique name
 - Instrument type: QuantStudio[™] 5 System
 - Block type: 96-Well 0.2-mL Block or 96-Well 0.1-mL Block
 - Experiment type: Standard Curve
 - Chemistry: TagMan[™] Reagents
 - Run Mode: Standard

4. In the **Method** tab, confirm that the **Volume** is 25 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	40

- 5. In the Plate tab, click Quick Setup.
- 6. In the Plate Attributes pane, confirm that the Passive Reference is set to None.
- 7. In the **Plate** tab, click **Advanced Setup**.
- 8. In the **Targets** table, confirm that the reporter dye and the target pairs are correct.

Reporter dye	Detector	Quencher
FAM	ORF1ab	None
VIC	N gene	None
ABY	S gene	None
JUN	MS2	None

- 9. Confirm that the targets above are assigned to each well in the plate layout.
- 10. In the plate layout pane, confirm the labeling of the control wells.
 - The template has one positive control (PC) and one negative control (NC) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 11. For all targets in the positive control well, confirm that **Task** is set to **S (Standard)**.
- 12. For all targets in the negative control well, confirm that **Task** is set to **N** (**Negative Control**).
- 13. In the **Samples** table, click **Add** to define the sample names. Create a unique sample name for each well in the physical plate.
- 14. To assign a sample to a well, select the well in the plate layout, then select the sample from the **Samples** table.

For all targets in the patient sample wells, confirm that **Task** is set to **U (Unknown)**.

Note: Wells that do not have a sample name will not be analyzed by the software.



- 15. In the Run tab, click Start Run, then select your instrument from the drop-down list.
- 16. Enter a file name in the dialog box that prompts you to save the run file, then save the file.

Set up and run the QuantStudio[™] 5 Real-Time PCR Instrument (384-well plates)—COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay.

For plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the QuantStudio™ 5 Real-Time PCR Instrument (384-well plates)—COVID-19 and RNase P assay" on page 96.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the QuantStudio[™] Design and Analysis Desktop Software v1.5.1, in the **New Experiment** box, select **Create New Experiment** ▶ **Template**.
- Browse to, then open the EDT file that you transferred in "Transfer the template (EDT) file for the QuantStudio™ 5 Real-Time PCR Instrument" on page 89.

IMPORTANT! Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

3. In the **Properties** tab, enter or confirm the following.

• Name: Enter a name

Instrument type: QuantStudio[™] 5 System

Block type: 384-well Block

• Experiment type: Standard Curve

Chemistry: TaqMan[™] Reagents

Run Mode: Standard

4. In the **Method** tab, confirm that the **Volume** is 20 µL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal/ extension	60°C	30 seconds	70

- 5. In the Plate tab, click Quick Setup.
- 6. In the Plate Attributes pane, confirm that the Passive Reference is set to None.

- 7. In the Plate tab, click Advanced Setup.
- 8. In the **Targets** table, confirm that the reporter dye and the target pairs are correct.

Reporter dye	Detector	Quencher
FAM	ORF1ab	None
VIC	N gene	None
ABY	S gene	None
JUN	MS2	None

- 9. Confirm that the targets above are assigned to each well in the plate layout.
- 10. In the plate layout pane, confirm the labeling of the control wells.
 - The template has one positive control (PC) and one negative control (NC) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.

Note: If samples from more than one RNA extraction run are being included on the same RT-PCR 384-well plate, then there must be one negative control for each RNA extraction run included on the RT-PCR 384-well plate (see "Guidelines for RT-PCR" on page 36). Label each negative control with a unique name, for example, *NC1*, *NC2*, *NC3*, and *NC4*.

- 11. For all targets in the positive control well, confirm that **Task** is set to **S** (**Standard**).
- 12. For all targets in the negative control well, confirm that **Task** is set to **N** (**Negative Control**).
- 13. In the **Samples** table, click **Add** to define the names. Create a unique sample name for each well in the physical plate.

The template has one positive control and one negative control assigned to wells for reference. If additional control wells are required, each control well must have a unique name.

14. To assign a sample to a well, select the well in the plate layout, then select the sample from the **Samples** table.

For all targets in the patient sample wells, confirm that **Task** is set to **U (Unknown)**.

Note: Wells that do not have a sample name will not be analyzed by the software.

- 15. In the Run tab, click Start Run, then select your instrument from the drop-down list.
- 16. Enter a file name in the dialog box that prompts you to save the run file, then save the file.



Set up and run the QuantStudio[™] 5 Real-Time PCR Instrument (96-well plates)—COVID-19 and RNase P assay

This section covers instrument set-up and operation for 96-well reaction plates that contain both the COVID-19 assay and the RNase P assay. Both assays are required for samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

For 96-well plates that contain only the COVID-19 assay, see "Set up and run the QuantStudio™ 5 Real-Time PCR Instrument (96-well plates)—COVID-19 assay only" on page 90.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the QuantStudio[™] Design and Analysis Desktop Software v1.5.1, in the **New Experiment** box, select **Create New Experiment** ▶ **Template**.
- 2. Browse to, then open the EDT file that you transferred in "Transfer the template (EDT) file for the QuantStudio™ 5 Real-Time PCR Instrument" on page 89.

IMPORTANT! Be careful to select the appropriate template file for your instrument, software version, and assay type. Failure to do so can cause errors in the analysis.

3. In the **Properties** tab, enter or confirm the following.

• Name: Enter a unique name

Instrument type: QuantStudio[™] 5 System

Block type: 96-Well 0.2-mL Block or 96-Well 0.1-mL Block

Experiment type: Standard Curve

Chemistry: TaqMan[™] Reagents

Run Mode: Standard

4. In the **Method** tab, confirm that the **Volume** is 25 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation 95°C		3 seconds	40
Anneal/ extension	60°C	30 seconds	70

- 5. In the Plate tab, click Quick Setup.
- 6. In the Plate Attributes pane, confirm that the Passive Reference is set to None.
- 7. In the Plate tab, click Advanced Setup.

8. In the **Targets** table, confirm that the reporter dye and the target pairs are correct.

Reporter dye	Detector	Quencher				
COVID-19 assaytargets	COVID-19 assaytargets					
FAM	ORF1ab	None				
VIC	N gene	None				
ABY	S gene	None				
JUN	MS2	None				
RNase P assaytarget						
VIC	RNaseP	None				

9. Confirm that the targets above are assigned to each well in the plate layout.

Note:

- · Wells containing the RNase P assay have only the RNaseP detector assigned.
- Wells containing the COVID-19 assay have all other detectors assigned.
- 10. In the plate layout pane, confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 11. For all targets in the positive control well, confirm that **Task** is set to **S (Standard)**.
- 12. For all targets in the negative control well, confirm that **Task** is set to **N** (Negative Control).
- 13. In the **Samples** table, click **Add** to define the sample names. The same sample name is used for each paired COVID-19 assay well and RNase P assay well in the physical plate.
- 14. To assign a sample to a well, select the well in the plate layout, then select the sample from the **Samples** table.

IMPORTANT! Paired sample assay wells that do not have the same sample name will not be analyzed by the software.

For all targets in the patient sample wells, confirm that **Task** is set to **U (Unknown)**.

- 15. In the Run tab, click Start Run, then select your instrument from the drop-down list.
- 16. Enter a file name in the dialog box that prompts you to save the run file, then save the file.

9

Set up and run the QuantStudio[™] 5 Real-Time PCR Instrument (384-well plates)—COVID-19 and RNase P assay

This section covers instrument set-up and operation for 384-well reaction plates that contain both the COVID-19 assay and the RNase P assay. Both assays are required for samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

For 384-well plates that contain only the COVID-19 assay, see "Set up and run the QuantStudio™ 5 Real-Time PCR Instrument (384-well plates)—COVID-19 assay only" on page 92.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- In the QuantStudio[™] Design and Analysis Desktop Software v1.5.1, in the New Experiment box, select Create New Experiment ➤ Template.
- 2. Browse to, then open the EDT file that you transferred in "Transfer the template (EDT) file for the QuantStudio™ 5 Real-Time PCR Instrument" on page 89.

IMPORTANT! Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

3. In the **Properties** tab, enter or confirm the following.

• Name: Enter a name

Instrument type: QuantStudio[™] 5 System

Block type: 384-well Block

Experiment type: Standard Curve
 Chemistry: TaqMan™ Reagents

Run Mode: Standard

4. In the **Method** tab, confirm that the **Volume** is 20 µL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	enaturation 95°C		40
Anneal/ extension	60°C	30 seconds	40

- 5. In the Plate tab, click Quick Setup.
- 6. In the Plate Attributes pane, confirm that the Passive Reference is set to None.
- 7. In the Plate tab, click Advanced Setup.

8. In the **Targets** table, confirm that the reporter dye and the target pairs are correct.

Reporter dye	Detector Quencher					
COVID-19 assaytargets						
FAM	ORF1ab	None				
VIC	N gene	None				
ABY	S gene	None				
JUN	MS2	None				
RNase P assaytarget						
VIC	RNaseP	None				

9. Confirm that the targets above are assigned to each well in the plate layout.

Note:

- · Wells containing the RNase P assay have only the RNaseP detector assigned.
- Wells containing the COVID-19 assay have all other detectors assigned.
- 10. In the plate layout pane, confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned. To assign additional negative control wells, see step 15.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.
- 11. For all targets in the positive control well, confirm that **Task** is set to **S (Standard)**.
- 12. For all targets in each negative control well, confirm that Task is set to N (Negative Control).
- 13. In the **Samples** table, click **Add** to define the sample names.

The same sample name is used for each paired COVID-19 assay well and RNase P assay well in the physical plate.

14. To assign a sample to paired COVID-19 and RNase P assay wells, select each well in the plate layout, then select the sample from the **Samples** table.

IMPORTANT! Paired sample assay wells that do not have the same name will not be analyzed by the software.

For all targets in the patient sample wells, confirm that **Task** is set to **U (Unknown)**.

15. If the 384-well plate includes additional negative controls (from combined extraction runs), define each additional negative control and assign it to paired control wells (COVID-19 and RNase P) to match the physical plate.

Give each negative control a unique name (for example, NC1, NC2, and so forth).



- 16. In the Run tab, click Start Run, then select your instrument from the drop-down list.
- 17. Enter a file name in the dialog box that prompts you to save the run file, then save the file.



Perform RT-PCR using the Applied Biosystems[™] QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well block)

Dye calibration for the QuantStudio[™] 7 Flex Real-Time PCR Instrument

A maintained instrument will be calibrated for many dyes. In addition to those dyes, the instrument operator must calibrate the instrument for ABY^{TM} dye and JUN^{TM} dye that are used with this kit. For all other assays, refer to the standard calibration process.

Transfer the template (EDT) file for the QuantStudio[™] 7 Flex Real-Time PCR Instrument (384–well block)

The template (EDT) file contains the settings for the instrument run. It is installed on the computer with Applied Biosystems[™] COVID-19 Interpretive Software, and must be transferred via a USB drive or other method to the computer on which the QuantStudio[™] Real-Time PCR Software v1.3 is installed.

IMPORTANT! Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

- 1. On the computer with Applied Biosystems™ COVID-19 Interpretive Software, navigate to one of the following directories, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):
 - <...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User Documents\COVID-19
 - ...>\Applied Biosystems\COVID-19 Interpretive Software\Client\docs\User
 Documents\COVID-19 + RNase P
 - <...> is the installation directory



Chapter 10 Perform RT-PCR using the Applied Biosystems[™] QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well block)

Set up and run the QuantStudio™ 7 Flex Real-Time PCR Instrument (384-well block)—COVID-19 assay only

2. Select the correct EDT file, based on whether the RT-PCR reaction contains only the COVID-19 assay, or both the COVID-19 assay and the RNase P assay (for Everlywell™ COVID-19 Test Home Collection Kit samples):

Assay type	Template name		
COVID-19 only	TaqPath COVID-19 Kit Template QS7 384 1_3 v2-2.edt		
COVID-19 and RNase P	TaqPath COVID-19 and RNaseP Template QS7 384 1_3 v1-0.edt		

3. Transfer the EDT file to the computer with QuantStudio[™] Real-Time PCR Software v1.3, using a USB drive or other method.

Set up and run the QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well block)—COVID-19 assay only

This section covers instrument set-up and operation for reaction plates that contain only the COVID-19 assay.

For plates that contain both the COVID-19 assay and RNase P assay, see "Set up and run the QuantStudio™ 7 Flex Real-Time PCR Instrument (384–well block)—COVID-19 and RNase P assay" on page 102.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- 1. In the QuantStudio[™] Real-Time PCR Software v1.3 home screen, click **Template**.
- Browse to, then open the EDT file that you transferred in "Transfer the template (EDT) file for the QuantStudio™ 7 Flex Real-Time PCR Instrument (384–well block)" on page 99.

IMPORTANT! Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Experiment Properties** tab, enter or confirm the following.
 - Experiment Name: Enter a unique name
 - Instrument type: QuantStudio[™] 7 Flex System
 - Block: 384-well
 - Type of Experiment: Standard Curve
 - Reagents: TaqMan[™]
 - Properties: Standard



4. In the **Define** tab, in the **Targets** pane, confirm that the targets, reporter dyes, and quenchers are listed correctly.

Target	Reporter dye	Quencher
ORF1ab	FAM	None
N gene	VIC	None
S gene	ABY	None
MS2	JUN	None

5. In the **Define** tab, in the **Samples** pane, define a unique sample name for each well in the physical plate.

Note: Wells that do not have a sample name will not be analyzed by the software.

- 6. In the **Define** tab, confirm that the **Passive Reference** is set to **None**.
- 7. In the **Assign** tab, confirm that four targets are assigned to each well in the plate layout. To assign a target to a well, select the well, then check the **Assign** checkbox.
- 8. In the **Assign** tab, in the **Samples** pane, confirm the labeling of the control wells.
 - The template has one positive control (PC) and one negative control (NC) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.

Note: If the 384-well plate includes additional negative controls (from combined extraction runs), define each additional negative control and assign it to a well to match the physical plate. Give each negative control a unique name (for example, *NC1*, *NC2*, *NC3*, and *NC4*).

- 9. In the **Assign** tab, confirm the **Task** assignments.
 - For wells with a Positive Control (PC), confirm that the Task is set to S for Standard for all of the targets.
 - For wells with a Negative Control (NC), confirm that the **Task** is set to **N** for Negative for all of the targets.
 - For the wells with a patient sample, confirm that the Task is set to U for Unknown for all of the targets.
- In the Assign tab, assign a sample name to each well to match the physical plate.
 To assign a sample to a well, select the well, then check the Assign checkbox.



Chapter 10 Perform RT-PCR using the Applied Biosystems[™] QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well block)

Set up and run the QuantStudio™ 7 Flex Real-Time PCR Instrument (384-well block)—COVID-19 and RNase Passay

11. In the **Run Method** tab, confirm that the **Reaction Volume Per Well** is 20 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles	
UNG incubation	25°C	2 minutes	1	
Reverse transcription	53°C	10 minutes	1	
Activation	95°C	2 minutes	1	
Denaturation	95°C	3 seconds	40	
Anneal / extension	60°C	30 seconds	70	

- 12. In the Run tab, click Start Run, then select your instrument from the drop-down list.
- 13. Enter a file name in the dialog box that prompts you to save the run file, then save the file.

Set up and run the QuantStudio[™] 7 Flex Real-Time PCR Instrument (384–well block)—COVID-19 and RNase P assay

This section covers instrument set-up and operation for 384-well reaction plates that contain both the COVID-19 assay and the RNase P assay. Both assays are required for samples collected using the Everlywell™ COVID-19 Test Home Collection Kit.

For 384-well plates that contain only the COVID-19 assay, see "Set up and run the QuantStudio™ 7 Flex Real-Time PCR Instrument (384–well block)—COVID-19 assay only" on page 100.

For more information about the instrument, see the documents listed in "Related documentation" on page 124.

- In the QuantStudio[™] Real-Time PCR Software v1.3 home screen, click Template.
- 2. Browse to, then open the EDT file that you transferred in "Transfer the template (EDT) file for the QuantStudio™ 7 Flex Real-Time PCR Instrument (384–well block)" on page 99.

IMPORTANT! Be careful to select the appropriate template file for your instrument, block type, and assay type. Failure to do so can cause errors in the analysis.

- 3. In the **Experiment Properties** tab, enter or confirm the following.
 - Experiment Name: Enter a unique name
 - Instrument type: QuantStudio[™] 7 Flex System
 - Block: 384-well
 - Type of Experiment: Standard Curve
 - Reagents: TaqMan[™]
 - Properties: Standard



4. In the **Define** tab, in the **Targets** pane, confirm that the targets, reporter dyes, and quenchers are listed correctly.

Target	Reporter dye	Quencher					
COVID-19 assaytargets	COVID-19 assaytargets						
ORF1ab	FAM	None					
N gene	VIC	None					
S gene	ABY	None					
MS2	JUN	None					
RNase P assaytarget							
RNaseP	VIC	None					

5. In the **Define** tab, in the **Samples** pane, define a sample name for each sample in the physical plate.

The same sample name is used for each paired COVID-19 assay well and RNase P assay well in the physical plate.

IMPORTANT! Paired sample assay wells that do not have the same name will not be analyzed by the software.

- 6. In the **Define** tab, confirm that the **Passive Reference** is set to **None**.
- 7. In the **Assign** tab, confirm that targets are assigned to each well in the plate layout.

Note:

- Wells containing the RNase P assay have only the RNaseP target assigned.
- Wells containing the COVID-19 assay have all other targets assigned.

To assign a target to a well, select the well, then check the **Assign** checkbox.

- 8. In the **Assign** tab, in the **Samples** pane, confirm the labeling of the control wells.
 - The template has one positive control and 2 negative controls (one each for the COVID-19 assay and RNase P assay) assigned to wells for reference.
 - Move the control well assignments by copying the existing control wells and pasting them according to their location on the physical plate.

Note: If the 384-well plate includes additional negative controls (from combined extraction runs), define each additional negative control and assign it to paired control wells (COVID-19 and RNase P) to match the physical plate.

Give each negative control a unique name (for example, NC1, NC2, and so forth).

Chapter 10 Perform RT-PCR using the Applied Biosystems[™] QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well block)

Set up and run the QuantStudio™ 7 Flex Real-Time PCR Instrument (384-well block)—COVID-19 and RNase Passay

- 9. In the **Assign** tab, confirm the **Task** assignments.
 - For wells with a Positive Control (PC), confirm that the Task is set to S for Standard for all of the targets.
 - For wells with a Negative Control (NC), confirm that the **Task** is set to **N** for Negative for all of the targets.
 - For the wells with a patient sample, confirm that the **Task** is set to **U** for Unknown for all of the targets.
- 10. In the **Assign** tab, assign a sample name to paired wells (COVID-19 and RNase P) to match the physical plate.

To assign a sample to a well, select the well, then check the **Assign** checkbox.

11. In the **Run Method** tab, confirm that the **Reaction Volume Per Well** is 20 μL, then confirm the thermal protocol.

Step	Temperature	Time	Number of cycles
UNG incubation	25°C	2 minutes	1
Reverse transcription	53°C	10 minutes	1
Activation	95°C	2 minutes	1
Denaturation	95°C	3 seconds	40
Anneal / extension	60°C	30 seconds	70

- 12. In the Run tab, click Start Run, then select your instrument from the drop-down list.
- 13. Enter a file name in the dialog box that prompts you to save the run file, then save the file.

Analysis and results

Obtain the Applied Biosystems[™] COVID-19 Interpretive Software

To perform data analysis and results interpretation, you must use the Applied Biosystems[™] COVID-19 Interpretive Software.

To obtain the software, contact your local instrument service team. Go to https://www.thermofisher.com/contactus.

Analyze the data

For detailed instructions about using the software, click the **Help** menu in the COVID-19 Interpretive Software.

- 1. Using a USB drive or other method, transfer the SDS or EDS files from the computer with the data collection software to the computer with the COVID-19 Interpretive Software.
- 2. Sign in to the COVID-19 Interpretive Software, then click the button that is appropriate for your test:
 - COVID-19 Test
 - COVID-19 + RNase P Test
- 3. In the **Home** screen, click the **Import Samples** button.
- 4. Select the SDS files or the EDS files to import, then click **Open**. After import, the software analyzes the run data, performs Quality Check (QC) analysis, and calculates the interpretive results for each sample and control.
- 5. In the **Home** screen, under **Batches**, click the <Batch ID> link for the batch of interest to display the **Batch Details** screen.
 - In the **Batch Details** screen, view the status and result for each sample in the **Sample**Information tab.
 - Optional: To identify signs of inadequate vortexing or centrifugation, select the Amplification Plot tab to view the amplification curve for each sample.
 - Optional: To enter a comment for the sample, select the Amplification Plot tab, select the Comment tab, enter a comment, then click Add.
 - The comment will appear in all exports (CSV files) and reports (PDF files). It will include the user name and date.

- 6. To generate a Batch Export (CSV file), return to the **Home** screen, select the checkbox for the batch, then click **Export Batch** at the top of the **Home** screen. Click **Open folder location** in the dialog box to locate and open the CSV file.
- 7. To generate a Batch Report (PDF file), select the checkbox for the batch, then click **Report Batch** at the top of the **Home** screen. Click **Open folder location** in the dialog box to locate and open the PDF file.

Interpretation of the results

Interpretation of the results is performed by the Applied BiosystemsTM COVID-19 Interpretive Software. For information about the C_t values that are used by the software to interpret results, see Appendix B, "Ct cutoff values for assay targets".

Quality control and validity of results

A minimum of one Negative Control and one Positive Control must be present for each run for the COVID-19 assay.

A minimum of two Negative Controls (one for the COVID-19 assay and one for the RNase P assay) and one Positive Control must be present for each run for the COVID-19 plus RNase P assay.

Additional Negative Control wells must be run for each extraction that is represented on a real-time RT-PCR plate. All control wells must pass for the real-time RT-PCR plate to be considered valid.

Validation of results is performed automatically by the Applied Biosystems[™] COVID-19 Interpretive Software based on performance of the Positive and Negative Controls.

Table 25 COVID-19 Test—Patient samples

ORF1ab	N gene	S gene	MS2	Status	Result	Action
NEG	NEG	NEG	NEG	INVALID	NA	Repeat test by re-extracting the original sample and repeating the RT-PCR. If the repeat result remains invalid, consider collecting a new specimen.
NEG	NEG	NEG	POS	VALID	SARS-CoV-2 Not Detected	Report results to the healthcare provider and appropriate public health authorities. Consider testing for other viruses.
Only one	SARS-Co ^V = POS	V-2 target	POS or NEG	VALID	SARS-CoV-2 Inconclusive	1. Repeat test by re-extracting the original sample and repeating the RT-PCR. 2. After retesting one time, report results to the healthcare provider and appropriate public health authorities. IMPORTANT! Samples with a result of SARS-CoV-2 Inconclusive shall be retested one time. If the repeat result remains inconclusive, the healthcare provider should conduct additional confirmation testing with a new specimen, if clinically indicated.
	nore SARS rgets = PC		POS or NEG	VALID	Positive SARS- CoV-2	Report results to the healthcare provider and appropriate public health authorities.

Table 26 COVID-19 + RNase P Test—Patient samples

ORF1ab	N gene	S gene	MS2	RNase P	Status	Result	Action
NEG	NEG	NEG	NEG	NEG	INVALID	NA	Repeat test by re-extracting the original sample and repeating the RT-PCR. If the repeat result remains invalid, consider collecting a new specimen.
NEG	NEG	NEG	NEG	POS	INVALID	NA	Repeat test by re-extracting the original sample and repeating the RT-PCR. If the repeat result remains invalid, consider collecting a new specimen.
NEG	NEG	NEG	POS	POS	VALID	SARS-CoV-2 Not Detected	Report results to the healthcare provider and appropriate public health authorities. Consider testing for other viruses.
NEG	NEG	NEG	POS	NEG	INVALID	NA	Repeat test by re-extracting the original sample and repeating the RT-PCR. If the repeat result remains invalid, consider collecting a new specimen.
	ne SARS- irget = PO		POS	or NEG	VALID	SARS-CoV-2 Inconclusive ^[1]	1. Repeat test by re-extracting the original sample and repeating the RT-PCR. 2. After retesting one time, report results to the healthcare provider and appropriate public health authorities. IMPORTANT! Samples with a result of SARS-CoV-2 Inconclusive shall be retested one time. If the repeat result remains inconclusive, the healthcare provider should conduct additional confirmation testing with a new specimen, if clinically indicated.
	nore SARS gets = PC		POS	or NEG	VALID	Positive SARS- CoV-2	Report results to the healthcare provider and appropriate public health authorities.

^[1] Samples with a result of SARS-CoV-2 Inconclusive shall be retested one time.



Conditions of authorization for labs

The TaqPath™ COVID-19 RT-PCR Kit and TaqPath™ COVID-19 Combo Kit Advanced Letter of Authorization, along with the authorized Fact Sheet for Healthcare Providers, the authorized Fact Sheet for Patients, and authorized labeling are available on the FDA website: https://www.fda.gov/emergency-preparedness-and-response/mcm-legal-regulatory-and-policy-framework/emergency-use-authorization#2019-ncov.

However, to assist clinical laboratories using the TaqPath[™] COVID-19 RT-PCR Kit and the TaqPath[™] COVID-19 RT-PCR Kit Advanced, the relevant Conditions of Authorization are listed below.

- Authorized laboratories^[1] using the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced will include with result reports of the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced 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 using the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced will perform the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced as outlined in the *TaqPath[™] COVID-19 Combo Kit Advanced Instructions for Use*. Deviations from the authorized procedures, including the authorized instruments, authorized extraction methods, authorized clinical specimen types, authorized control materials, authorized other ancillary reagents, and authorized materials required to perform the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced are not permitted.
- Authorized laboratories that receive the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19 Combo Kit Advanced must notify the relevant public health authorities of their intent to run the test prior to initiating testing.
- Authorized laboratories using the TaqPath[™] COVID-19 Combo Kit and the TaqPath[™] COVID-19
 Combo Kit Advanced will have a process in place for reporting test results to healthcare providers and relevant public health authorities, as appropriate.
- Authorized laboratories testing nasal swab specimens self-collected using the Everlywell[™]
 COVID-19 Test Home Collection Kit must be laboratories designated by Everlywell, Inc. and follow
 the *Receiving and Processing Everlywell Samples* standard operating procedure when accepting
 specimens for testing.
- Authorized laboratories will collect information on the performance of the test and report
 to DMD/OHT7-OIR/OPEQ/CDRH (via email: CDRH-EUA-Reporting@fda.hhs.gov) and Thermo
 Fisher Scientific (techservices@thermofisher.com; 1 800 955 6288) any suspected occurrence of
 false positive or false negative results and significant deviations from the established performance
 characteristics of the test of which they become aware.

^[1] For ease of reference, this letter will refer 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."

- All laboratory personnel using the test must be appropriately trained in RT-PCR techniques and use appropriate laboratory and personal protective equipment when handling this kit, and use the test in accordance with the authorized labeling.
- Thermo Fisher Scientific, its authorized distributor(s), and authorized laboratories using the TaqPath™ COVID-19 Combo Kit and the TaqPath™ COVID-19 Combo Kit Advanced 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

Analytical performance of the TaqPath[™] COVID-19 RT-PCR Kit was evaluated by determining limit of detection (LoD), characterizing the impact of interfering substances and cross-reactivity, as described in the following sections.

Limit of detection (LoD)

The LoD study established the lowest SARS-CoV-2 viral concentration (Genomic Copy Equivalents or GCE) that can be detected by the TaqPath™ COVID-19 Combo Kit in a particular specimen type at least 95% of the time. Banked Nasopharyngeal swab (NP) and Bronchoalveolar lavage (BAL) samples, obtained from U.S. patients in the years 2015-2019, were pooled, respectively, and spiked with purified SARS-CoV-2 viral genomic RNA at several concentrations and processed through the TaqPath™ COVID-19 Combo Kit workflow. A three-phase approach was used to determine the LoD for each specimen type. In phases I and II, the preliminary LoD was established and confirmed in phase III by testing 20 replicates.

Table 27 LoD determination in BAL

Effective	Replicate		Mea	n C _t		Interpretation	% Positive
Concentration	Replicate	ORF1ab	N	S	MS2	interpretation	% FOSITIVE
	1	29.9	29.1	28.5	23.1	Positive	
	2	30.1	29.3	29.7	24.0	Positive	
	3	30.0	29.7	29.3	24.0	Positive	
	4	30.3	29.7	29.1	23.8	Positive	
	5	30.2	29.6	29.6	23.7	Positive	
	6	30.3	29.3	29.7	23.5	Positive	
10 GCE/reaction	7	29.9	29.6	32.8	23.4	Positive	100%
	8	30.2	29.8	29.2	23.8	Positive	
	9	30.1	29.4	28.6	23.8	Positive	
	10	30.1	29.4	29.1	24.0	Positive	
	11	29.8	29.5	29.4	24.3	Positive	
	12	30.1	29.7	29.1	24.6	Positive	
	13	30.7	30.1	28.4	25.1	Positive	

Table 27 LoD determination in BAL (continued)

Effective	Replicate		Mea		Interpretation	% Positive	
Concentration	Replicate	ORF1ab	N	S	MS2	interpretation	70 1 OSIUVE
	14	30.4	29.8	29.1	24.8	Positive	
	15	30.2	29.8	29.7	24.9	Positive	
	16	30.3	29.8	29.4	24.6	Positive	
10 GCE/reaction	17	30.4	30.0	31.5	24.7	Positive	100%
	18	30.4	30.1	29.3	24.9	Positive	
	19	30.9	29.7	29.2	25.4	Positive	
	20	30.3	29.9	29.4	25.7	Positive	

Table 28 LoD determination in Nasopharyngeal specimens

Effective	Replicate		Mea	n C _t		Interpretation	% Positive
Concentration	Replicate	ORF1ab	N	S	MS2	interpretation	70 1 OSITIVE
	1	30.0	28.9	35.7	25.7	Positive	
	2	30.6	28.9	33.6	25.8	Positive	
	3	30.2	28.8	32.0	25.8	Positive	
	4	30.4	28.7	34.2	25.7	Positive	
	5	30.5	29.0	31.4	25.8	Positive	
	6	31.0	29.3	36.6	26.0	Positive	
	7	30.3	29.2	31.1	25.8	Positive	
	8	31.1	29.2	31.8	26.5	Positive	
10 GCE/reaction	9	30.5	28.9	33.0	26.2	Positive	100%
	10	30.3	28.8	34.7	26.8	Positive	
	11	30.5	29.8	38.7	27.4	Positive	
	12	31.6	29.7	35.0	27.6	Positive	
	13	30.7	29.3	36.4	27.4	Positive	
	14	31.6	28.8	31.3	27.2	Positive	
	15	31.0	29.3	36.0	27.0	Positive	
	16	30.5	29.1	35.7	27.0	Positive	
	17	30.7	29.4	34.8	27.4	Positive	



Table 28 LoD determination in Nasopharyngeal specimens (continued)

Effective	Replicate		Mea	Interpretation	% Positive		
Concentration	Replicate	ORF1ab	N	S	MS2	interpretation	% FOSILIVE
	18	30.7	29.3	34.6	27.5	Positive	
10 GCE/reaction	19	31.0	29.3	35.9	28.7	Positive	100%
	20	30.4	29.2	32.7	28.4	Positive	

Table 29 LoD results

Specimen type	Limit of Detection (GCE/reaction)
Bronchoalveolar lavage	10 GCE/reaction
Nasopharyngeal swab	10 GCE/reaction

Limit of detection (LoD) for RT-PCR with 17.5 μ L and 14.0 μ L of purified sample RNA

The LoD study with 17.5 µL and 14.0 µL of purified sample RNA established the lowest SARS-CoV-2 viral concentration (Genomic Copy Equivalents or GCE) that can be detected by the TaqPath[™] COVID-19 Combo Kit Advanced at least 95% of the time. Banked Nasopharyngeal swab (NP) samples were pooled and spiked with gamma-irradiated SARS-CoV-2 virus at several concentrations and processed through the TaqPath[™] COVID-19 Combo Kit Advanced workflow. A three-phase approach was used to determine the LoD for each specimen type. In phases I and II, the preliminary LoD was established and confirmed in phase III by testing 20 replicates.

Table 30 7500 Fast Dx Real-Time PCR Instrument (96-well), 17.5 µL of purified sample RNA

Concentration	Replicate	ORF1ab	N gene	S gene	MS2	Interpretation	% Positive
	1	32.975	33.685	32.953	22.847	Positive	
	2	33.166	33.585	35.330	22.603	Positive	
	3	33.935	33.298	33.235	22.608	Positive	
	4	33.950	33.259	33.290	22.646	Positive	
50 GCE/mL	5	35.133	Undetectable	33.382	22.605	Positive	100%
(≤7 GCE/ reaction)	6	32.808	34.349	36.963	22.422	Positive	100%
	7	33.917	33.736	33.330	22.164	Positive	
	8	34.014	33.378	33.422	22.589	Positive	
	9	34.266	37.113	34.356	22.373	Positive	
	10	35.234	33.341	33.372	22.726	Positive	

Table 30 7500 Fast Dx Real-Time PCR Instrument (96-well), 17.5 µL of purified sample RNA (continued)

Concentration	Replicate	ORF1ab	N gene	S gene	MS2	Interpretation	% Positive
	11	35.012	35.053	35.610	23.059	Positive	
	12	34.519	37.033	34.424	22.762	Positive	
	13	35.234	35.736	33.564	22.623	Positive	
	14	33.594	36.446	34.853	22.715	Positive	
50 GCE/mL	15	34.320	38.100	32.873	22.368	Positive	100%
(≤7 GCE/ reaction)	16	34.195	36.707	33.931	22.004	Positive	100%
	17	34.484	34.133	36.220	21.957	Positive	
	18	33.904	33.656	33.643	22.414	Positive	
	19	36.505	Undetectable	36.198	22.422	Positive	
	20	35.304	33.726	35.437	22.743	Positive	

Table 31 QuantStudio[™] 7 Flex Real-Time PCR Instrument (384-well), 14.0 µL of purified sample RNA

Concentration	Replicate	ORF1ab	N gene	S gene	MS2	Interpretation	% Positive
	1	33.064	33.380	34.410	22.847	Positive	
	2	34.727	33.488	33.894	22.465	Positive	
	3	33.364	33.584	33.596	22.569	Positive	
	4	32.872	33.518	34.151	22.291	Positive	
	5	33.359	34.923	33.267	22.099	Positive	
	6	33.379	Undetectable	35.909	22.514	Positive	100%
	7	33.738	33.190	36.431	21.908	Positive	
50 GCE/mL	8	33.430	34.375	33.968	21.933	Positive	
(≤5.6 GCE/ reaction)	9	34.121	35.457	34.250	21.829	Positive	
	10	32.608	34.032	34.510	21.825	Positive	
	11	32.958	Undetectable	33.301	22.152	Positive	
	12	33.760	33.723	33.910	22.473	Positive	
	13	33.657	33.837	35.406	22.137	Positive	
	14	Undetectable	34.200	33.961	21.745	Positive	
	15	33.560	35.388	33.713	22.517	Positive	
	16	34.338	34.091	34.007	21.838	Positive	

Table 31 QuantStudio 7 Flex Real-Time PCR Instrument (384-well), 14.0 µL of purified sample RNA (continued)

Concentration	Replicate	ORF1ab	N gene	S gene	MS2	Interpretation	% Positive
	17	33.350	34.315	34.410	21.657	Positive	
50 GCE/mL	18	34.025	33.251	34.292	21.584	Positive	100%
(≤5.6 GCE/ reaction)	19	33.277	34.955	34.468	21.541	Positive	100 /6
	20	33.562	33.137	34.488	21.551	Positive	

Reactivity (Inclusivity)

In silico analysis was updated on June 3, 2020, using 25,998 complete SARS-CoV-2 genomes in GISAID and GenBank databases. Based upon BLAST analysis, the TaqPath™ COVID-19 Combo Kit maps with 100% homology to >99.99% of known SARS-CoV-2 isolates in GISAID and 100% of known isolates in GenBank databases. Mapping was deemed successful for a given isolate if at least two of the three targets (ORF1ab, S gene, and N gene) showed 100% identity.

Interfering substances

Pooled SARS-CoV-2-negative nasopharyngeal swab and bronchoalveolar lavage specimens were spiked with purified SARS-CoV-2 viral RNA at 3X the limit of detection (30 GCE/reaction) and potential interfering substances at the concentrations listed in Table 32 on page 116. Each substance was tested with triplicate extractions. The results are presented in Table 32 on page 116.

Pooled SARS-CoV-2-negative nasopharyngeal swab and bronchoalveolar lavage specimens were spiked with potential interfering substances at the concentrations above. Each substance was tested with triplicate extractions. No false positive results were observed for any of the substances at the concentrations tested.

Table 32 Interfering substances

	Final concentration	Agreement with expected results					
Interfering substance	Final concentration in sample	Positive BAL samples	Positive NP samples	Negative BAL samples	Negative NP samples		
None	N/A	100% ^[1]	100%	100%	100%		
Mucin: bovine submaxillary gland, type I-S	0.1 mg/mL	100% ^[2]	100%	100%	100%		
Blood (human)	1% v/v	100% ^[3]	100%	100%	100%		
Nasal sprays or drops— Nasacort™	10% v/v	100% ^[4]	100% ^[4]	100%	100%		
Nasal corticosteroids—Dymista [™]	5 μg/mL	100% ^[2]	100%	100%	100%		
NeilMed™ Nasogel™	1% w/v	100% ^[2]	100%	100%	100%		
Influenza A H1N1 Brisbane/59/07	1 × 10 ⁵ TCID ₅₀ /mL	100% ^[2]	100%	100%	100%		
Throat lozenges, oral anesthetic and analgesic—Chloraseptic™	1% w/v	100% ^[3]	100%	100%	100%		
Oseltamivirphosphate	33 μg/mL	100% ^[2]	100%	100%	100%		
Antibiotic, nasal ointment— Bactroban [™]	5 μg/mL	100% ^[2]	100%	100%	100%		
Antibacterial, systemic— Tobramycin	0.6 mg/mL	100% ^[2]	100%	100%	100%		
Homeopathic allergy relief medicine—Similasan™ Nasal	10% v/v	100%	100%	100%	100%		

^[1] Two of six replicates produced a $C_t > 37$ or Undetermined for S Gene, but all replicates were called Positive based on the interpretation algorithm.

 $^{^{[2]}}$ Two of three replicates produced a $C_t > 37$ or Undetermined for S Gene, but all replicates were called Positive based on the interpretation algorithm.

 $^{^{[3]}}$ All three replicates produced a $C_t > 37$ or Undetermined for S Gene but were called Positive based on the interpretation algorithm.

^[4] One of three replicates produced a $C_t > 37$ or Undetermined for S Gene, but all replicates were called Positive based on the interpretation algorithm

Cross-reactivity

In silico analysis of the following forty-three (43) organisms was performed.

Table 33 Organisms used for in silico cross-reactivity analysis

Human coronavirus 229E	Rhinovirus/Enterovirus
Human coronavirus OC43	Parechovirus
Human coronavirus HKU1	Candida albicans
Human coronavirus NL63	Corynebacterium diphtheriae
SARS-coronavirus	Legionella (non-pneumophila)
MERS-coronavirus	Bacillus anthracis (Anthrax)
Adenovirus	Moraxella catarrhalis
Human Metapneumovirus (hMPV)	Neisseria elongata and Neisseria meningitidis
Parainfluenza 1	Pseudomonas aeruginosa
Parainfluenza 2	Staphylococcus epidermidis
Parainfluenza 3	Streptococcus salivarius
Parainfluenza 4	Leptospira sp.
Influenza A	Chlamydophila pneumoniae
Influenza B	Chlamydophila psittaci
Influenza C	Coxiella burnetii (Q-Fever)
Enterovirus	Staphylococcus aureus
Respiratory Syncytial Virus A	Haem ophilus influenzae
Respiratory Syncytial Virus B	Legionella pneumophila
Bordetella pertussis	Mycobacterium tuberculosis
Mycoplasma pneumoniae	Streptococcus pneumoniae
Pneumocystis jirovecii(PJP)	Streptococcus pyogenes

Among the tested organisms, *Neisseria elongata* showed homology for the forward and reverse primers and probe for the N gene. The forward primer showed ≥80% homology while the reverse primer and probe showed 36% homology. The N gene reverse primer and probe show low homology, therefore the risk of the non-specific amplification is low.

Blast analysis showed ≥80% homology for one assay component (forward primer, reverse primer, or probe) for select isolates. Despite ≥80% homology of one assay component for select isolates, there is no anticipated amplification because hybridization of all three assay components are necessary to generate a signal. We also found multiple instances where different assay components had ≥80%

homology to different isolates of the same species. For example, *Bacillus anthracis* strain AFS029987 had ≥80% homology to the ORF1ab forward primer while strain MCCC 1A01412 had ≥80% homology to the ORF1ab reverse primer. Since these are two different organisms, amplification is not likely to occur. The *in silico* analysis indicates that significant amplification of non-target sequences that result in cross-reactivity or potentially interfere with detection of SARS-CoV-2 is not likely to occur.

Clinical evaluation

A clinical evaluation study was performed to evaluate the performance of the TaqPath[™] COVID-19 RT-PCR Kit using nasopharyngeal swab (NP) and bronchoalveolar lavage (BAL) specimens.

A total of sixty (60) contrived positive specimens were tested:

- 30 contrived positive nasopharyngeal swab (NP) specimens
- 30 contrived positive bronchoalveolar lavage (BAL) specimens

Samples were contrived by spiking known concentrations of extracted SARS-CoV-2 viral genomic RNA, relative to the product LoD, into matrices which were determined to be negative by the TaqPath[™] COVID-19 RT-PCR Kit prior to spiking in the RNA.

In addition to the contrived positive specimens, sixty (60) negative specimens were tested:

- 30 negative nasopharyngeal swab (NP) specimens
- 30 negative samples bronchoalveolar lavage (BAL) specimens
- All negative samples yielded negative results

Results for positive samples are shown in the tables below:

Table 34 BAL Clinical Evaluation Study

Final RNA	Number of	Mean C _t				
Concentration in Sample	Positives	S gene	ORF1ab	N gene		
2X LoD	20/20 ^[1]	28.9	29.5	28.7		
3X LoD	5/5[1]	28.8	29.2	28.5		
5X LoD	5/5	27.4	28.2	27.4		

^[1] Two samples initially gave inconclusive results and were retested. The results were positive after the retest. Mean C_t values are calculated from the retest results.



Table 35 NP Clinical Evaluation Study

Final RNA	Number of	Mean C _t						
Concentration in Sample	Positives	S gene	ORF1ab	N gene				
2X LoD	20/20 ^[1]	30.9	30.6	29.3				
3X LoD	5/5	30.0	30.1	28.8				
5X LoD	5/5	28.7	29.0	27.9				

^[1] One sample initially gave an inconclusive result and was retested. The result was positive after the retest. Mean C_t values are calculated from the retest results.



Example extraction plate and RT-PCR reaction plate layouts (Everlywell[™] COVID-19 Test Home Collection Kit samples)

Everlywell™ COVID-19 Test Home Collection Kit samples require that users prepare two RT-PCR plate wells for each sample and Negative Control—one for the COVID-19 assay and one for the RNase P assay—plus a single RT-PCR plate well for the Positive Control. A single 96-well extraction plate contains enough samples for two 96-well RT-PCR reaction plates, or may be combined with another extraction plate onto a single 384-well RT-PCR reaction plate.

The following diagrams provide example layouts for extraction plates and RT-PCR plates, assuming that you want to fit the maximum number of samples onto each plate.

Clinical evaluation

Layout of RNA extraction plate

92 RNA samples, 2 Negative Controls

	1	2	3	4	5	6	7	8	9	10	11	12
Α	NC1	S1	S2	S3	S4	S5	S6	S7	S8	S9	S10	S11
В	S12	S13	S14	S15	S16	S17	S18	S19	S20	S21	S22	S23
С	S24	S25	S26	S27	S28	S29	S30	S31	S32	S33	S34	S35
D	S36	S37	S38	S39	S40	S41	S42	S43	S44	S45	S46	EMPTY
Е	NC2	S47	S48	S49	S50	S51	S52	S53	S54	S55	S56	S57
F	S58	S59	S60	S61	S62	S63	S64	S65	S66	S67	S68	S 69
G	S7 0	S71	S72	S73	S74	S75	S76	S77	S78	S7 9	S80	S81
Н	S82	S83	S84	S85	S86	S87	S88	S89	S90	S91	S92	EMPTY



Layout of 2 96-well RT-PCR plates from 1 RNA extraction plate

Each RT-PCR plate contains the following:

46 RNA samples

1 Negative Control into 2 wells (COVID-19 and RNase Passay wells)

1 Positive Control (COVID-19 assay well)

RNase P B NCR1 S1 S2 S3 S4 S5 S6 S7 S8 S9 S10 S1 COVID-19 C S12 S13 S14 S15 S16 S17 S18 S19 S20 S21 S22 S2 RNase P D S12 S13 S14 S15 S16 S17 S18 S19 S20 S21 S22 S2	RT-PCR plate #1	1	T-PCR plate #1	2	3	4	5	6	7	8	9	10	11	12
COVID-19 C S12 S13 S14 S15 S16 S17 S18 S19 S20 S21 S22 S2 RNase P D S12 S13 S14 S15 S16 S17 S18 S19 S20 S21 S22 S2	COVID-19 A	NC1	OVID-19 A	S1	S2	S3	S4	S5	S6	S7	S8	S9	S10	S11
RNase P D S12 S13 S14 S15 S16 S17 S18 S19 S20 S21 S22 S2	RNase P B	NCR1	Nase P B	S1	S2	S3	S4	S5	S6	S7	S8	S9	S10	S11
	COVID-19 C	S12	OVID-19 C	S13	S14	S15	S16	S17	S18	S19	S20	S21	S22	S23
00/40 - 004 005 000 007 000 000 004 000 004 000	RNase P D	S12	Nase P D	S13	S14	S15	S16	S17	S18	S19	S20	S21	S22	S23
COVID-19 E 524 525 526 527 528 529 530 531 532 533 534 53	COVID-19 E	S24	OVID-19 E	S25	S26	S27	S28	S29	S30	S31	S32	S33	S34	S35
RNase P F S24 S25 S26 S27 S28 S29 S30 S31 S32 S33 S34 S3	RNase P F	S24	Nase P F	S25	S26	S27	S28	S29	S30	S31	S32	S33	S34	S35
COVID-19 G S36 S37 S38 S39 S40 S41 S42 S43 S44 S45 S46 PC	COVID-19 G	S36	OVID-19 G	S37	S38	S39	S40	S41	S42	S43	S44	S45	S46	PC
RNase P H S36 S37 S38 S39 S40 S41 S42 S43 S44 S45 S46 EMP	RNase P H	S36	Nase P H	S37	S38	S39	S40	S41	S42	S43	S44	S45	S46	EMPTY

RT-PCR plate	#2												
		1	2	3	4	5	6	7	8	9	10	11	12
COVID-19	Α	NC2	S47	S48	S49	S50	S51	S52	S53	S54	S55	S56	S57
RNase P	В	NCR2	S47	S48	S49	S50	S51	S52	S53	S54	S 55	S 56	S57
COVID-19	С	S58	S59	S60	S 61	S 62	S 63	S64	S65	S 66	S 67	S 68	S69
RNase P	D	S58	S59	S60	S 61	S 62	S 63	S64	S65	S 66	S 67	S 68	S 69
COVID-19	Ε	S70	S71	S72	S73	S74	S75	S76	S77	S78	S79	S80	S81
RNase P	F	S70	S71	S72	S73	S74	S75	S76	S77	S78	S79	S80	S81
COVID-19	G	S 82	S 83	S 84	S 85	S 86	S87	S 88	S 89	S 90	S 91	S 92	PC
RNase P	Н	S 82	S 83	S84	S85	S 86	S87	S88	S 89	S90	S 91	S92	EMPTY

Figure 9 Example plate layouts for 96-well RT-PCR reactions (Everlywell™ COVID-19 Test Home Collection Kit samples)



Appendix A Example extraction plate and RT-PCR reaction plate layouts (Everlywell™ COVID-19 Test Home Collection Kit samples)

Clinical evaluation

Layout of 2 RNA extraction plates

RNA extraction plate #1

95 RNA samples, 1 Negative Control

	1	2	3	4	5	6	7	8	9	10	11	12
Α	NC1	S1	S2	S3	S4	S5	S6	S7	S8	S9	S10	S11
В	S12	S13	S14	S15	S16	S17	S18	S19	S20	S21	S22	S23
С	S24	S25	S26	S27	S28	S29	S30	S31	S32	S33	S34	S35
D	S36	S37	S38	S39	S40	S41	S42	S43	S44	S45	S46	S47
E	S48	S49	S50	S51	S52	S53	S54	S55	S56	S57	S58	S59
F	S60	S61	S62	S63	S64	S65	S66	S67	S68	S69	S70	S71
G	S72	S73	S74	S75	S76	S77	S78	S79	S80	S81	S82	S83
Н	S84	S85	S86	S87	S88	S89	S90	S91	S92	S93	S94	S95

RNA extraction plate #2

94 RNA samples, 1 Negative Control

	1	2	3	4	5	6	7	8	9	10	11	12
Α	NC2	S96	S97	S98	S 99	S100	S101	S102	S103	S104	S105	S106
В	S107	S108	S109	S110	S111	S112	S113	S114	S115	S116	S117	S118
С	S119	S120	S121	S122	S123	S124	S125	S126	S127	S128	S129	S130
D	S131	S132	S133	S134	S135	S136	S137	S138	S139	S140	S141	S142
E	S143	S144	S145	S146	S147	S148	S149	S150	S151	S152	S153	S154
F	S155	S156	S157	S158	S159	S160	S161	S162	S163	S164	S165	S166
G	S167	S168	S169	S170	S171	S172	S173	S174	S175	S176	S177	S178
Н	S179	S180	S181	S182	S183	S184	S185	S186	S187	S188	S189	EMPTY



RT-PCR plate contains the following:

189 RNA samples

2 Negative Controls into 4 wells (2 COVID-19 and 2 RNase Passay wells)

1 Positive Control (COVID-19 assay well)

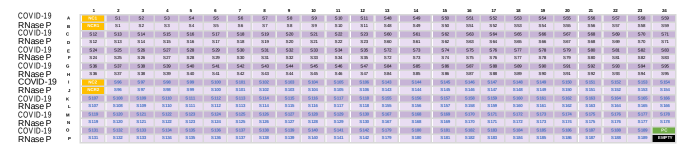


Figure 10 Example plate layouts for 384-well RT-PCR reactions (Everlywell™ COVID-19 Test Home Collection Kit samples)



Ct cutoff values for assay targets

The Applied Biosystems $^{\text{\tiny TM}}$ COVID-19 Interpretive Software uses the following C_t cutoff values for assay targets during interpretation of the results.

Table 36 COVID-19 Assay Ct cutoff values

Sample or Control	Target	C _t cutoff			
Positive Control	MS2	Valid C _t values are >37			
Fositive Control	Viral targets	Valid C _t values are ≤37			
Nogotive Control	MS2	Valid C _t values are ≤32			
Negative Control	Viral targets	Valid C _t values are >37			
Clinical complex	MS2	Valid C _t values are ≤32 ^[1]			
Clinical samples	Viral targets	Positive C _t values are ≤37			

^[1] If any of the viral targets is positive, the C_t for MS2 can be >32.

Table 37 RNase P Assay Ct cutoff values

Sample or Control	Target	C _t cutoff				
Negative Control RNase P		Valid C _t values are >33				
Clinical samples	RNase P	Valid C _t values are ≤33				



Documentation and support

Related documentation

Document	Publication Number
Applied Biosystems [™] 7500 Fast Dx Real-Time PCR Instrument Reference Guide	4406991
Applied Biosystems [™] 7500/7500 Fast Real-Time PCR System: Maintenance Guide	4387777
QuantStudio [™] 3 and 5 Real-Time PCR Systems Installation, Use, and Maintenance Guide	MAN0010407
QuantStudio [™] 6 and 7 Flex Real-Time PCR Systems Maintenance and Administration Guide	4489821
QuantStudio [™] 6 and 7 Flex Real-Time PCR Systems Quick Reference	4489826
QuantStudio™ Real-Time PCR Software Getting Started Guide	4489822
MagMAX [™] Viral/Pathogen Nucleic Acid Isolation Kit (automated extraction) User Guide	MAN0018073
MagMAX [™] Viral/Pathogen Nucleic Acid Isolation Kit (manual extraction) User Guide	MAN0018072
Thermo Scientific™ KingFisher™ Flex User Manual	N07669
COVID-19 Interpretive Software Installation Quick Reference	MAN0019257
TaqMan [™] SARS-CoV-2 RNase P Assay Kit Product Information Sheet	MAN0019524

Customer and technical support

For additional documentation and information about this kit, visit: https://www.thermofisher.com/covid19

For download instructions for the COVID-19 Interpretive Software, see "Obtain the Applied Biosystems™ COVID-19 Interpretive Software" on page 105.

Refer to the Read Me file provided with the COVID-19 Interpretive Software before contacting support for the software.

Visit: https://www.thermofisher.com/contactus for service and support information for this kit, including the following:

- Worldwide contact telephone numbers
- Product support information
- Order and web support

- Product documentation such as:
 - Certificates of Analysis
 - Safety Data Sheets (SDSs; also known as MSDSs)

Note: For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

Limited product warranty

Life Technologies Corporation and/or its affiliate(s) warrant their products as set forth in the Life Technologies' General Terms and Conditions of Sale at www.thermofisher.com/us/en/home/global/terms-and-conditions.html. If you have any questions, please contact Life Technologies at www.thermofisher.com/support.

