1	
2	
3	For use under the Emergency Use Authorization
4	(EUA) only
5	For in vitro diagnostic use
6	Rx Only
7	
8	
9	Lyra® SARS-CoV-2 Assay
10	Instructions for Use
11	
12	
13	For the qualitative detection of human coronavirus SARS-CoV-2 viral RNA extracted from
14	nasal, nasopharyngeal and oropharyngeal swab specimens.
15 16	
10 17	

18	Contents	
19	Intended Use	4
20	Summary and Explanation	4
21	Principle of the Procedure	4
22	Materials Provided	5
23	Materials Required But Not Provided	6
24	Warnings and Precautions	6
25	Storage and Handling of Kit Reagents	7
26	Specimen Collection, Storage and Handling	8
27	Nucleic Acid Extracts Storage	8
28	bioMérieux NucliSENS easyMAG Nucleic Acid Extraction Programming Instructions	8
29	Assay Procedure	10
30	Quality Control	11
31	CLINICAL PERFORMANCE	13
32	Level of Detection	14
33	Analytical Reactivity (Inclusivity)	19
34	Analytical Specificity (Cross-Reactivity)	19
35	Limitations	22
36	Customer and Technical Assistance	23
37	References	24
38	APPENDIX	25
39	Applied Biosystems 7500 Fast Dx Programming Instructions	25
40	Applied Biosystems® 7500 Fast Dx Thermocycler Test Procedure	27
41	Applied Biosystems 7500 Standard Programming Instructions	28
42	Applied Biosystems® 7500 Standard Thermocycler Test Procedure	30
43	Bio-Rad CFX96 Touch Thermocycler Programming Procedure	31

Quidel Corporation

Lyra® SARS-CoV-2 Assay 4/06/2020 Page 3 of 44

44	Bio-Rad CFX96 Touch Thermocycler Test Procedure	33
45	Qiagen Rotor-Gene Q Programming Instructions	33
46	Qiagen Rotor-Gene Q Test Run	35
47	Roche's LightCycler® 480 Instrument II Programming Instructions	36
48	Creating a LC 480 II Assay Run Template	36
49	Creating a LC 480 II Assay Test Procedure	37
50	ThermoFisher QuantStudio 7 Pro Programming Instructions	38
51	Creating a ThermoFisher QuantStudio 7 Pro Test Procedure	40
52	GLOSSARY	44
53		

Intended Use

The Lyra® SARS-CoV-2 Assay is a real-time RT-PCR assay intended for the qualitative detection of nucleic acid from SARS-CoV-2 in nasal, nasopharyngeal (NP), or oropharyngeal (OP) swab specimens from patients suspected of COVID-19 by their healthcare provider. Testing is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, to perform high complexity tests.

Results are for the identification of SARS-CoV-2 RNA. The SARS-CoV-2 is generally detectable in upper respiratory 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. Laboratories within the United States and its territories are required to report all

positive results to the appropriate public health authorities.

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

The Lyra SARS-CoV-2 Assay is 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 Lyra SARS-CoV-2 Assay is only for use under the Food and Drug Administration's Emergency Use Authorization.

Summary and Explanation

SARS-CoV-2, also known as the COVID-19 virus, was first identified in Wuhan, Hubei Province, China December 2019. This virus, as with the novel coronavirus SARS-1 and MERS, is thought to have originated in bats, however the SARS-CoV-2 may have had an intermediary host such as pangolins, pigs or civets. By the start of March 2020, human infection has spread to over 74 countries, infected over 92,000 people and has killed over 3100 people. On March 11, the WHO had declared the SARS-CoV-2 as a global pandemic.

The median incubation time is estimated to be 5.1 days with symptoms expected to be present within 12 days of infection.² The symptoms of COVID-19 are similar to other viral respiratory diseases and include fever, cough and shortness of breath.³

The Lyra SARS-CoV-2 Assay has been designed to specifically detect SARS-CoV-2 RNA.

Principle of the Procedure

The Lyra SARS-CoV-2 Assay detects SARS-CoV-2 viral RNA that has been extracted from a patient sample using either the bioMerieux NucliSENS® easyMAG® system or EMAG® system. A multiplex real-time RT-PCR reaction is carried out under optimized conditions in a single tube generating amplicons for the targeted virus (if present) and the Process Control (PRC) present in the sample. This reaction is performed utilizing one of six thermocyclers: Applied Biosystems 7500 Fast Dx, Applied Biosystems 7500 Standard, Roche LightCycler 480, Qiagen Rotor-Gene Q, Bio-Rad CFX96 Touch, Thermofisher QuantStudio 7 Pro. Identification of the SARS-CoV-2 virus occurs by the use of target specific primers and fluorescent-labeled probes that hybridize to a conserved region of the non-structural polyprotein of the SARS-CoV-2 virus.

Table 1. Lyra® SARS-CoV-2 Assay Probe Labels					
Target Dye					
Non-structural polyprotein (pp1ab)	FAM				
Process Control (PRC)	Quasar® 670				

The following is a summary of the procedure:

1. **Sample Collection:** Obtain nasopharyngeal, or opharyngeal, or nasal swabs using standard techniques from symptomatic patients. These specimens are transported, stored, and processed according to established laboratory procedures.

Nucleic Acid Extraction: Extract nucleic acids from the specimens with the NucliSENS® easyMAG® or EMAG®
 Systems following the manufacturer's instructions and using the appropriate reagents (See Materials
 Required but Not Provided).

Prior to the extraction procedure add 20 μ L of the Process Control (PRC) to each 180 μ L aliquot of specimen or controls. The PRC serves to monitor inhibitors in the extracted specimen, assures that adequate amplification has taken place and confirms that the nucleic acid extraction was sufficient.

3. **Rehydration of Master Mix:** Rehydrate the lyophilized Master Mix using 135µL of Rehydration Solution. The Master Mix contains oligonucleotide primers, fluorophore and quencher-labeled probes targeting conserved regions of the SARS-CoV-2 as well as the process control sequence. The probes are dual labeled with a reporter dye attached to the 5' end and a quencher attached to the 3' end. The rehydrated Master Mix is sufficient for eight reactions.

4. **Nucleic Acid Amplification and Detection:** Add 15 μ L of the rehydrated Master Mix to each plate well (Applied Biosystems® 7500 Fast Dx, Applied Biosystems 7500 Standard, the Roche LightCycler 480) or tube (Qiagen Rotor-Gene Q). 5 μ L of extracted nucleic acids (specimen with PRC) is then added to the plate well or tube. Place the plate or tube into the appropriate instrument.

Once the reaction plate or tubes are added to the instrument, the assay protocol is initiated. This protocol initiates reverse transcription of the RNA targets generating complementary DNA, and the subsequent amplification of the target sequences occurs. The Lyra SARS-CoV-2 Assay is based on TaqMan® chemistry, and uses an enzyme with reverse transcriptase, DNA polymerase, and 5'-3' exonuclease activities. During DNA amplification, this enzyme cleaves the probe bound to the complementary DNA sequence, separating the quencher dye from the reporter dye. This step generates an increase in fluorescent signal upon excitation by a light source of the appropriate wavelength. With each cycle, additional dye molecules are separated from their quenchers resulting in additional signal. If sufficient fluorescence is achieved, the sample is reported as positive for the detected target sequence.

Materials Provided

SKU # CE-M120

Table 2. Detection Kit (96 Reactions) – Store at 2°C to 8°C						
#	Component	Quantity				
0	Rehydration Solution Part M5003	1 vial/kit 1.9 mL				
0	Lyra SARS-CoV-2 Master Mix Part M5150 Lyophilized Contents:	12 vials/kit, 8 reactions/vial				

Table 2. Det	Table 2. Detection Kit (96 Reactions) – Store at 2°C to 8°C					
#	Component	Quantity				
	DNA polymerase enzyme with reverse transcriptase activity					
	Oligonucleotide primer pairs; Oligonucleotide probes					
	dNTPs (dATP, dCTP, dGTP, dUTP, dTTP)					
	Stabilizers					
CONTROL	Process Control Part M5005	1 vial/kit 2.0 mL				
CONTROL +	Positive Control containing SARS-CoV-2 Synthetic RNA, Part M5153	1 vial/kit 1.0 mL				
CONTROL -	Negative Control Part M5031	1 vial/kit 2.0 mL				

Lyra™ SARS-CoV-2 Assay Instructions for Use

146 147

158

159

Materials Required But Not Provided

- Micropipettors (range between 1 to 10 μL and 100 to 1000 μL)
- Non-aerosol pipette tips
- Applied Biosystems®7500Fast Dx, software version 1.4
- Applied Biosystems®Standard, software version 2.0.6
- Roche LightCycler® 480 Instrument II, software version 1.5.0.39
- Qiagen Rotor-Gene Q, software version 2.0.2.4
- Bio-Rad CFX96 Touch, software version 3.1
- Thermofisher QuantStudio 7 Pro, software version 2.0
- 156 96 well PCR plate #:
- 157 Applied Biosystems®7500Fast Dx: 4344906
 - Applied Biosystems®Standard: N8010560
 - Roche LightCycler® 480: 04729692001, foil included
- 160 Bio-Rad CFX96 Touch: HSP9631, seals MSB1001
- 161 Thermofisher Quantstudio 7 Pro: 4483354
- Optical plate films
- 163 Qiagen Rotor-Disc
- Qiagen Rotor-Disc Heat Sealing Film
- Plate centrifuge for 96 well plate
- bioMérieux NucliSENS easyMAG software version 2.0
- bioMérieux EMAG software version 2.0
- bioMérieux NucliSENS easyMAG Buffers 1, 2, 3
- bioMérieux NucliSENS easyMAG Lysis Buffer
- bioMérieux NucliSENS easyMAG Silica Magnetic Beads
- 171 bioMérieux NucliSENS easyMAG disposables
- 172 Biohit pipettor

173174

175

Warnings and Precautions

- For In Vitro Diagnostic Use under Emergency Use Authorization only.
- Positive results are indicative of the presence of SARS-CoV-2 RNA.
- Laboratories within the United States and its territories are required to report all positive results to the
 appropriate public health authorities.

- The assay has been validated using bioMérieux NucliSENS easyMAG software version 2.0. Please contact Quidel Technical Support prior to modifying or upgrading beyond this version of software.
- The assay has been validated using Applied Biosystems 7500Fast Dx software version 1.4. Please contact
 Quidel Technical Support prior to modifying or upgrading beyond this version of software.
- The assay has been validated using Applied Biosystems Standard software version 2.0.6. Please contact Quidel Technical Support prior to modifying or upgrading beyond this version of software.
- The assay has been validated using Roche LightCycler® 480 Instrument II, software version 1.5.0.39
 Please contact Quidel Technical Support prior to modifying or upgrading beyond this version of software.
- The assay has been validated using Qiagen Rotor-Gene Q, software version 2.0.2.4. Please contact Quidel
 Technical Support prior to modifying or upgrading beyond this version of software.
- The assay has been validated using Bio-Rad CFX96 Touch, software version 3.1. Please contact Quidel Technical Support prior to modifying or upgrading beyond this version of software.
 - The assay has been validated using Thermofisher QuantStudio 7 Pro, software version 2.0. Please contact Quidel Technical Support prior to modifying or upgrading beyond this version of software.
- Performance characteristics of this test have been established with the specimen types listed in the

 Intended Use Section only. The performance of this assay with other specimen types or samples has not
 been evaluated.
- Use of this product should be limited to personnel with sufficient training in PCR and RT-PCR techniques.
 - Treat all specimen/samples as potentially infectious. Follow universal precautions when handling samples, this kit and its contents.
- Proper sample collection, storage and transport are essential for correct results.
- Store assay reagents as indicated on their individual labels.
 - Wear suitable protective clothing, gloves, eye and face protection when using this kit.
- For accurate results, pipette carefully using only calibrated equipment.
 - Thoroughly clean and disinfect all surfaces with a 10% bleach solution followed by molecular grade water.
- Use micropipettes with an aerosol barrier or positive displacement tips for all procedures.
- Avoid microbial and cross contamination of the kit reagents. Follow Good Laboratory Procedures.
- Do not mix reagents from kits with different lot numbers.
- Do not use reagents from other manufacturers with this kit.
- Do not use product after its expiration date.

192

197

198

201

203

218

219

220 221

222

223

224

225

226227

228

- Proper workflow planning is essential to minimize contamination risk. Always plan laboratory workflow in a uni-directional manner, beginning with pre-amplification and moving through amplification and detection.
- Use dedicated supplies and equipment in pre-amplification and amplification areas.
- Do not allow cross movement of personnel or equipment between areas.
- Keep amplification supplies separate from pre-amplification supplies at all times.
- Do not open sample tubes or unseal plates post amplification.
- Dispose of amplified material carefully and in accordance with local laws and regulations in order to
 minimize the risk of amplicon contamination.
 - Do not use supplies dedicated for reagent or sample preparation for processing target nucleic acid.
 - MSDS is available upon request or can be accessed on the product website.

Storage and Handling of Kit Reagents

- Store the unopened kit at 2°C to 8°C until the expiration date listed on the outer kit box.
- The rehydrated Master Mix may be stored at room temperature (20°C to 25°C) for up to 24 hours. For longer storage the rehydrated Master Mix should be recapped, sealed with parafilm and stored in an upright position at ≤–20°C for up to 14 days. Protect the Master Mix from light during storage.

Indications of Instability or Deterioration of Reagents: Cloudiness of the Rehydration Solution, when within expiration, may indicate deterioration of this reagent. Contact Quidel Technical Assistance for a replacement.

229									
230	-			ge and Handling					
231				geal, or nasal specimens should be collected, transported, stored, and processed					
232	according to CLSI M41-A ² . Specimens should be stored at 2°C to 8°C until tested. If specimens cannot be								
233	tes	ted withi	in 72 hours of co	ellection, they should be frozen at -70°C or colder until tested.					
234 235	The following viral transport media (M4, M4-RT, M5, M6, MTM and UTM) (1 mL and 3 mL) are compatible								
236		-	a respiratory as	·					
237				https://www.cdc.gov/coronavirus/2019-ncov/downloads/Viral-Transport-					
238	Me	<u>:dium.pd</u>	<u>f</u>) is compatible	with the Lyra SARS-CoV-2 Assay.					
239									
240			cid Extracts S	_					
241				easyMAG can be stored at room temperature (20°C to 25°C) for 2 hours, at 2°C to					
242	8°C	for 24 h	ours and 1 mon	th at -20°C to -70°C.					
243									
244 245	bio	oMérie	ux NucliSENS	easyMAG Nucleic Acid Extraction Programming Instructions					
246			•	e. Lyra SARS-CoV-2, Positive Control #M5153), and a negative process control (i.e.					
247	Lyr	a SAKS-C	.ov-z, Negative (Control #M5031) should be included in each extraction run.					
248	1.	Turn or	the instrument	and wait for instrument light to appear orange. Then switch on the					
249		comput	ter/launch easyN	MAG software. Do not log into software until the light on the instrument has					
250		turned	green.						
251	2	Danaad		pressing the 'Instrument' and 'Reagent Inventory' buttons.					
251	2.	Barcou	e reagents after	pressing the 'Instrument' and 'Reagent Inventory' buttons.					
				THE STATE OF THE S					
252	3.	To ente	er samples, press	s the 'Daily Use' button, which will default to the 'Define Request'					
253		screen.	Select the follow	wing settings:					
254		a.	Sample ID:	Enter the sample name using the keyboard.					
255		b.	Matrix:	Select Other from the drop-down menu					
256		c.	Request:	Select Generic from the drop-down menu					
257		d.	Volume (mL):	Select 0.200 from the drop-down menu					
258		e.	Eluate (μL):	Select 50 from the drop-down menu					
259		f.	Type:	Primary					
260		g.	Priority:	Normal					
		J	,						
261	4.	Upon p	ressing the 'Save	e' button, the sample will appear in the 'Unassigned Sample' window on					
			J						
262		عام املاء	a: da af tha ann	button and report the					
262				en. Press the 'Enter New Extraction Request' button, and repeat the					
263		process	s for additional s	amples. Alternatively multiple samples can be entered by pressing the 'Auto					
264		Create	New Extraction	Requests' button.					
		J. Juic		· · · · · · · · · · · · · · · · · · ·					

Once all samples are created, go to 'Organize Runs' by clicking on the 265 icon near the top of the 266 page. Create a run by pressing the 'Create Run' button. Enter a run name or use the default. 267 6. Add samples to the run by using the 'Auto Fill Run' button (auto fills up to 24 samples from the 268 'Unassigned Sample list' on the left hand side of the screen). Alternatively, individual samples can be 269 moved into and out of the run by using the left and right 'Positioning icons' after selecting 270 the appropriate sample. The sample order within the run can be changed using the 'Move Extraction 271 Request Up/Down' buttons 272 7. Obtain 1 to 3 (for 8 to 24 samples, respectively) sample vessel(s), and add 20 μL of Process Control to each 273 sample well used. 274 Add 180 μL of each sample to the appropriate well as designated. 275 9. Go to 'Load Run' by pressing the button near the top of the screen. Insert tips and sample 276 vessel(s) into the instrument 277 10. Enter the barcode(s) of the sample vessel(s) 278 11. Enter the barcode(s) of silica beads to be used 279 12. Close the instrument lid. 280 13. Assign silica beads to samples as follows: 281 a. Click the reagents symbol below number 1 in the picture below. The lot number of the silica 282 beads should appear below the Silica tab at number 2 in the picture below. 283 b. Highlight and select the samples in the run for which beads need to be assigned (in the box 284 containing number 3 in the picture below) 285 positioning icon (below number 4 in the picture below) to assign the silica lot Click the 286 number to the selected samples 287 If the bead symbol to the right of number 5 in the picture below is selected, the silica bead lot 288 number should be displayed for each sample 匨





290 14. Print work list by touching 'Load Run' icon followed by pressing the 'Print Work List' icon

- 291 15. Press the 'Dispense Lysis' button. The on-board lysis will take approximately 12 minutes to complete.
 - 16. For each sample vessel, prepare magnetic particles using the Biohit pipettor and tips for up to eight reactions as follows:
 - a. Using 1 tip and Program 1, aspirate 550 μ L nuclease-free water and dispense into a 1.5 mL DNAse / RNAse free microfuge tube.
 - b. Vortex the magnetic silica. Using 1 tip and Program 1, aspirate 550 μ L of magnetic silica, dispense into the water and mix by vortexing.
 - c. Using 1 tip and Program 2, aspirate 1050 μ L of the magnetic silica mixture and dispense 25 μ L back into the same tube.
 - d. Dispense 125 µL magnetic silica mixture each into 8 wells of an ELISA strip plate. Discard tip.
 - e. After Lysis is complete (NB: the 'Instrument Status' at the bottom of the screen must be 'IDLE'!), using 8 tips and Program 3, aspirate 100 μ L of magnetic silica mixture in strip wells, dispense 100 μ L of magnetic silica mixture in strip wells, and aspirate 100 μ L of magnetic silica mixture in strip wells.
 - f. Insert tips into liquid within the sample vessels. Aspirate 800 μ L then dispense 900 μ L of magnetic silica mixture back into vessel. Aspirate 1000 μ L of magnetic silica mixture from vessel and dispense 1000 μ L of magnetic silica back into vessel. Repeat aspiration / dispensing of 1000 μ L two more times.
- 310 17. Close the instrument and press the 'Start' button to begin the run.
- 311 18. Upon completion of run, transfer purified nucleic acid to nuclease-free tubes. Eluates from the easyMAG can be stored at room temperature (20°C to 25°C) for 2 hours, at 2°C to 8°C for 24 hours and 1 month at 20°C to -70°C.

Assay Procedure

Run the following procedures at controlled room temperature of 20°C to 25°C.

316 317

318

319

320

321

322

323

324

327

314

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

Master Mix Rehydration Procedure

- 1. Determine the number of specimens extracted to be tested and obtain the correct number of eighttest lyophilized Master Mix vials for testing.
- 2. Return unused reagents to the appropriate storage conditions.
- 3. Open Master Mix carefully to avoid disruption of the pellet.
- 4. Add 135 μL of Rehydration Solution to the Master Mix.
- 5. Place vial at room temperature for 1 to 2 minutes to allow rehydration of pellet.
- 325 6. Gently pipette up and down 2 to 3 times avoiding the formation of bubbles prior to dispensing into the first plate well or tube.
 - **Note**: The rehydrated Master Mix is sufficient for 8 reactions.
- Note: The rehydrated Master Mix may be stored at room temperature (20°C to 25°C) for up to 24 hours. For longer storage the rehydrated Master Mix should be recapped, sealed with parafilm and stored in an upright position at ≤−20°C for up to 14 days. Protect the Master Mix from light during storage.

334

335

336

337

338

339

340

341

RT-PCR Set-up Procedure:

- 1. Add 15 μ L of the rehydrated Master Mix to each plate well or tube.
- 2. Add 5 μ L of extracted nucleic acid (specimen with the process control) into the plate well or tube. Mixing of reagents is not required.

Note: Use a new barrier micropipettor tip with each extracted specimen.

- 3. Seal the plate or tubes.
- 4. Centrifuge the plate or tubes for a minimum of 15 seconds. Ensure that all liquid is at the bottom of the plate wells or tubes.
- 5. Turn on the appropriate thermocycler.
- 6. Insert plate or tubes into the appropriate thermocycler.

342 343

NOTE: Refer to Appendix for specific programming and testing protocols of each thermocycler.

344345

346

Quality Control

The Lyra SARS-CoV-2 Assay incorporates several controls to monitor assay performance.

347348349

350

351

352

 The Process Control (PRC) consists of an inactivated and stabilized MS2 Bacteriophage that contains an RNA genome. It must be used during extraction <u>and</u> amplification in the assay. This control should be added to each sample aliquot prior to extraction. The PRC serves to monitor inhibitors in the extracted specimen, assures that adequate amplification has taken place and confirms that the nucleic acid extraction was sufficient.

353 354 355

2. The **Positive Control** (containing SARS-CoV-2 Synthetic RNA, Part M5153) must be treated as a patient specimen and be included in every extraction and RT-PCR run.

356357358

3. The **Negative Control** (Part M5031) must be treated as a patient specimen and be included in every extraction and PCR run.

359 360 361

362

363

4. Failure of either the **Positive Control** or the **Negative Control** invalidates the RT-PCR run and results should not be reported. The RT-PCR run should be repeated with the extracted controls and specimens first. Re-extract and retest another aliquot of the controls and the specimens or obtain new samples and retest if the controls fail again.

364 365 366

Table 3. Expected Results from Controls (Applied Biosystems 7500 Fast Dx, Applied Biosystems 7500 Standard, Bio-Rad Cfx96, Qiagen Rotor-Gene Q, or Thermofisher QS-7)

Control Type/ Name	Used to Monitor	SARS-CoV-2	Expected Ct Values	PRC	Expected Ct Values
Positive Control	Substantial reagent failure including primer and probe integrity	+	5.0≤ Ct ≤30.0	+/-	NA¹
Negative Control	Reagent and/or environmental contamination	-	None detected	+	5.0≤ Ct ≤30.0

Expected Results from Controls (Roche LightCycler 480)							
Control Type/ Name	Used to Monitor	SARS-CoV-2	Expected Ct Values	PRC	Expected Ct Values		
Positive Control	Substantial reagent failure including primer and probe integrity	+	5.0≤ Ct ≤40.0	+/-	NA ¹		
Negative Control	Reagent and/or environmental contamination	-	None detected	+	5.0≤ Ct ≤40.0		

¹No Ct value is required for the Process Control to make a positive call.

Interpretation of Results from Patient Specimens

368

369

370 371

Table 4. Interpretation of the Lyra SARS-CoV-2 Assay Results on the Applied Biosystems 7500 Fast Dx, Applied Biosystems 7500 Standard, Bio-rad Cfx96, Qiagen Rotor-Gene Q, or Thermofisher QS-7

Assay Result	Detector: SARS-CoV-2	Detector: Process Control	Interpretation of Results	Notes and Special Guidance
Negative	No Ct detected	5.0≤ Ct ≤30.0	No SARS-CoV-2 viral RNA detected; PRC Detected.	
SARS-CoV-2 Positive	5.0≤ Ct ≤30.0	NA ¹	SARS-CoV-2 Virus viral RNA detected.	
Invalid	No Ct detected	No Ct detected	No SARS-CoV-2 viral RNA and no PRC RNA detected.	Invalid test. Retest the same purified sample. If the test is also invalid, re-extract and retest another aliquot of the same specimen or obtain a new specimen and retest.

Interpretation of the Lyra SARS-CoV-2 Assay Results on the Roche LightCycler 480

Assay Result	Detector: SARS-CoV-2	Process Control	Interpretation of Results	Notes and Special Guidance
Negative	No Ct detected	5.0≤ Ct ≤40.0	No SARS-CoV-2 viral RNA detected; PRC Detected.	
SARS-CoV-2 Positive	5.0≤ Ct ≤40.0	NA ¹	SARS-CoV-2 Virus viral RNA detected.	
Invalid	No Ct detected	No Ct detected	No SARS-CoV-2 viral RNA and no PRC RNA detected.	Invalid test. Retest the same purified sample. If the test is also invalid, re-extract and retest another aliquot of the

		same specimen or obtain a
		new specimen and retest.

¹ No Ct value is required for the Process Control to make a positive call.

CLINICAL PERFORMANCE

The clinical performance of the Lyra SARS-CoV-2 Assay was evaluated using two different studies:

 A study using two hundred sixty-five fresh or frozen nasopharyngeal swab specimens collected in UTM (36 and 229, respectively) from patients located in the USA.

• A fully contrived positive specimen study using nasopharyngeal swab specimens.

Α ϵ

All two hundred sixty-five specimens were negative for SARS-CoV-2 when extracted with the easyMAG system and tested by the Lyra SARS-CoV-2 Assay.

One hundred twenty-four specimens included in this study were positive for other circulating respiratory viruses as identified by FDA-cleared assays:

Circulating Virus	# of positive specimens
Influenza A	30
RSV	4
Coronavirus Seasonal (not identified)	10
Coronavirus 229e	20
Coronavirus OC43	20
Coronavirus NL63	20
Coronavirus HKU1	20

Viral RNA was obtained from BEI Resources for use in the contrived clinical study. The genomic RNA was extracted from a preparation of cell lysate and supernatant from Cercopithecus aethiops kidney epithelial (Vero E6, ATCC® CRL-1586™) cells infected with SARS-related coronavirus 2 (SARS-CoV-2), isolate USA-WA1/2020, using QIAamp® Viral RNA Mini Kit (Qiagen 52904). The viral genomic RNA is in a background of cellular nucleic acid and carrier RNA. The Genome Copy Number was established using BioRad QX200 Droplet Digital PCR (ddPCR™) System.

Ninety-two positive contrived samples were created by spiking ninety-two individual clinical samples determined to be negative for SARS-CoV-2 by the Lyra SARS-CoV-2 Assay prior to spiking with one of three concentrations of genomic SARS-CoV-2 RNA. Forty-four specimens were spiked with 1x LoD (8.00E-01 cp/ μ L) of RNA. Twenty-four additional specimens were spiked with 3x LoD (2.40E00 cp/ μ L) of RNA. Twenty-four additional specimens were spiked with 5x LoD (4.00E00 cp/ μ L) of RNA. All samples were extracted and tested according to the Lyra SARS-CoV-2 Assay package insert.

All ninety-two contrived samples were positive in the Lyra SARS-CoV-2 Assay. The results for the contrived positive specimens are shown in the table below:

Table 5. Clinical evaluation in spiked nasopharyngeal swab specimens						
Sample RNA # Positives/# Tested Mean SARS-CoV-2 Ct %CV						
Concentration						
unspiked						

1 .0x LoD	44/44	26.9	5.7
3x LoD	24/24	22.8	3.4
5x LoD	24/24	22.4	3.0

Performance against the expected results are:

Positive Percent Agreement 92/92 = 100% (95% CI: 95.99%-100%) Negative Percent Agreement 92/92 = 100% (95% CI: 95.99%-100%)

405 406 407

403

404

ANALYTICAL PERFORMANCE

Level of Detection

408 409

412

413

414

419

420

The Limit of Detection of the Lyra SARS-CoV-2 Assay utilized limiting dilutions of genomic SARS-CoV-2 RNA in negative nasopharyngeal matrix. Each dilution was extracted using the NucliSENS

easyMAG System and tested on Applied Biosystems 7500 Fast Dx, Applied Biosystems 7500

Standard, Roche LightCycler 480, Qiagen Rotor-Gene Q, Bio-Rad CFX96 Touch, or Thermofisher

QuantStudio 7 Pro. Analytical sensitivity (LoD) is defined as the lowest concentration at which at

415 least 95% of all replicates tested positive.

The genomic RNA was extracted from a preparation of cell lysate and supernatant from

417 Cercopithecus aethiops kidney epithelial (Vero E6, ATCC® CRL-1586™) cells infected with SARS-

related coronavirus 2 (SARS-CoV-2), isolate USA-WA1/2020, using QIAamp® Viral RNA Mini Kit

(Qiagen 52904). The viral genomic RNA is in a background of cellular nucleic acid and carrier RNA.

The Genome Copy Number was established using BioRad QX200 Droplet Digital PCR (ddPCR™)

421 System.

This study established the LoD for the Lyra SARS-CoV-2 Assay as 8.00E-01 genomic RNA copies/ μ L, subsequently confirmed by testing 20 replicates.

423 424

Table 6. LoD in Nasopharyngeal specimens with Applied Biosystems 7500 Fast Dx						
Concentration	Replicate	SARS-CoV-2 Ct	PRC Ct	Interpretation		
8.00E-01	1	23.95	18.54	Positive		
genomic RNA	2	26.59	18.28	Positive		
copies/μL	3	26.19	18.32	Positive		
	4	25.13	18.41	Positive		
	5	24.88	18.74	Positive		
	6	24.84	19.18	Positive		
	7	25.51	18.82	Positive		
	8	25.20	18.58	Positive		
	9	24.69	18.71	Positive		
	10	24.57	18.67	Positive		
	11	23.86	18.75	Positive		
	12	24.58	18.91	Positive		
	13	25.19	19.03	Positive		

Table 6. LoD in Nasopharyngeal specimens with Applied Biosystems 7500 Fast Dx					
Concentration	Replicate	SARS-CoV-2 Ct	PRC Ct	Interpretation	
	14	25.84	19.05	Positive	
	15	26.58	19.10	Positive	
	16	26.72	19.15	Positive	
	17	24.16	19.06	Positive	
	18	25.15	18.91	Positive	
	19	25.51	19.05	Positive	
	20	24.41	19.07	Positive	

Table 7. LoD in Oropharyngeal specimens with Applied Biosystems 7500 Fast Dx					
Concentration	Replicate	SARS-CoV-2 Ct	PRC Ct	Interpretation	
8.00E-01	1	27.26	19.38	Positive	
genomic RNA	2	28.99	19.22	Positive	
copies/µL	3	27.3	19.51	Positive	
	4	26.09	19.27	Positive	
	5	26.88	19.61	Positive	
	6	26.02	19.19	Positive	
	7	26.37	19.21	Positive	
	8	25.01	19.30	Positive	
	9	25.14	19.06	Positive	
	10	26.21	19.03	Positive	
	11	27.79	19.27	Positive	
	12	28.83	19.12	Positive	
	13	28.83	19.19	Positive	
	14	26.81	19.50	Positive	
	15	25.1	19.30	Positive	
	16	26.2	19.11	Positive	
	17	26.74	19.00	Positive	
	18	25.28	19.13	Positive	
	19	26.27	19.31	Positive	
	20	26.37	19.24	Positive	

Table 8. LoD in Nasopharyngeal specimens with Applied Biosystems 7500 Standard							
Concentration	Concentration Replicate SARS-CoV-2 Ct PRC Ct Interpretation						
1 26.63 19.26 Positive							
2 29.15 19.28 Positive							

Table 8. LoD in Nasopharyngeal specimens with Applied Biosystems 7500 Standard					
Concentration	Replicate	SARS-CoV-2 Ct	PRC Ct	Interpretation	
8.00E-01	3	25.67	19.69	Positive	
genomic RNA	4	25.53	20.07	Positive	
copies/µL	5	26.15	20.50	Positive	
	6	26.71	20.50	Positive	
	7	26.11	19.14	Positive	
	8	26.94	19.18	Positive	
	9	25.62	18.64	Positive	
	10	25.80	18.80	Positive	
	11	26.76	19.15	Positive	
	12	26.15	19.63	Positive	
	13	27.42	19.44	Positive	
	14	27.51	19.99	Positive	
	15	26.07	19.9	Positive	
	16	25.92	18.81	Positive	
	17	27.95	20.02	Positive	
	18	27.71	19.27	Positive	
	19	26.51	18.86	Positive	
	20	Undetermined	19.11	Negative	

1	1	7
4	Z	/

Table 9. LoD in Nasopharyngeal specimens with Roche LightCycler 480*					
Concentration	Replicate	SARS-CoV-2 Ct*	PRC Ct*	Interpretation	
8.00E-01	1	32.91	31.73	Positive	
genomic RNA	2	34.54	32.9	Positive	
copies/μL	3	34.83	32.25	Positive	
	4	34.94	31.7	Positive	
	5	33.81	32.14	Positive	
	6	34.36	32.37	Positive	
	7	33.90	32.10	Positive	
	8	33.83	32.80	Positive	
	9	33.8	31.86	Positive	
	10	34.28	32.27	Positive	
	11	33.63	32.81	Positive	
	12	33.72	32.45	Positive	
	13	34.86	33.17	Positive	
	14	34.57	32.64	Positive	

Table 9. LoD in Nasopharyngeal specimens with Roche LightCycler 480*							
Concentration	Replicate SARS-CoV-2 Ct* PRC Ct* Interpretation						
	15	34.48	32.92	Positive			
	16	33.61	32.82	Positive			
	17	33.87	33.34	Positive			
	18	34.44	33.36	Positive			
	19	34.22	32.55	Positive			
	20	33.77	32.97	Positive			

* Results include 10 cycles not captured by the other instruments

Table 10. LoD in Nasopharyngeal specimens with Qiagen Rotor-Gene Q					
Concentration	Replicate	SARS-CoV-2 Ct	PRC Ct	Interpretation	
8.00E-01	1	24.01	19.08	Positive	
genomic RNA	2	24.04	19.36	Positive	
copies/μL	3	24.85	19.44	Positive	
	4	23.23	19.13	Positive	
	5	24.39	19.07	Positive	
	6	23.89	18.94	Positive	
	7	23.78	18.80	Positive	
	8	24.82	18.86	Positive	
	9	23.87	18.83	Positive	
	10	24.05	18.90	Positive	
	11	23.28	18.84	Positive	
	12	24.36	18.71	Positive	
	13	23.85	18.87	Positive	
	14	23.54	18.88	Positive	
	15	24.84	19.20	Positive	
	16	23.63	19.01	Positive	
	17	24.18	18.97	Positive	
	18	23.47	19.01	Positive	
	19	23.58	18.94	Positive	
	20	23.89	19.02	Positive	

Table 11. LoD in Nasopharyngeal specimens with Bio-Rad CFX96 Touch								
Concentration	Replicate	Replicate SARS-CoV-2 Ct PRC Ct Interpretation						
8.00E-01	1	27.19	21.25	Positive				
genomic RNA	2	25.57	21.35	Positive				
copies/μL	3	25.80	22.68	Positive				

Table 11. LoD in Nasopharyngeal specimens with Bio-Rad CFX96 Touch						
Concentration	Replicate	Replicate SARS-CoV-2 Ct P		Interpretation		
	4	27.93	21.3	Positive		
	5	29.03	21.09	Positive		
	6	25.79	21.45	Positive		
	7	25.65	21.19	Positive		
	8	26.26	21.16	Positive		
	9	29.46	21.41	Positive		
	10	25.09	21.45	Positive		
	11	25.68	21.36	Positive		
	12	28.51	21.49	Positive		
	13	25.5	21.97	Positive		
	14	26.81	21.36	Positive		
	15	26.17	21.1	Positive		
	16	25.04	21.91	Positive		
	17	25.47	22.08	Positive		
	18	25.54	21.26	Positive		
	19	25.77	22.29	Positive		
	20	25.59	22.16	Positive		

Table 12. LoD in Nasopharyngeal specimens with Thermofisher QS-7						
Concentration	Replicate	SARS-CoV-2 Ct	PRC Ct	Interpretation		
8.00E-01	1	24.25	20.21	Positive		
genomic RNA	2	26.7	20.9	Positive		
copies/μL	3	27.14	20.6	Positive		
	4	27.28	20.81	Positive		
	5	29.60	20.78	Positive		
	6	26.99	20.65	Positive		
	7	28.75	20.82	Positive		
	8	27.63	20.76	Positive		
	9	29.80	20.65	Positive		
	10	26.60	20.55	Positive		
	11	27.23	20.54	Positive		
	12	29.81	20.73	Positive		
	13	26.59	20.88	Positive		
	14	27.23	20.87	Positive		
	15	26.63	20.62	Positive		

Table 12. LoD in Nasopharyngeal specimens with Thermofisher QS-7									
Concentration	Replicate	Replicate SARS-CoV-2 Ct PRC Ct Interpretation							
	16 26.07 20.84 Positive								
	17	25.14	20.81	Positive					
	18	27.34	20.6	Positive					
	19	29.22	20.67	Positive					
	20	26.37	20.38	Positive					

Analytical Reactivity (Inclusivity)

The inclusivity of the Lyra SARS-CoV-2 Assay was established by testing Genomic RNA from the SARS-related coronavirus 2 (SARS-CoV-2), isolate USA-WA1/2020, via *in-silico* analysis. The *in-silico* analysis demonstrated the Lyra SARS-CoV-2 primers are 100% conserved to 257 SARS-CoV-2 sequences available from NCBI and GISAID as of March 5, 2020.

,

Analytical Specificity (Cross-Reactivity)

The Analytical Specificity of the assay was established by both direct testing of organisms in the assay ("wet" testing) and *in silico* analysis. The wet testing used 25 micro-organisms, in high concentrations, identified by the FDA as high priority for evaluation due to the reasonable likelihood they may be present in upper respiratory samples. All micro-organisms were undetectable with the Lyra SARS-CoV-2 Assay when wet tested as shown below.

Table 11. Cross-reactivity test results								
-		Source/						
Virus/Bacteria/Parasite	Strain	Sample type	Concentration	Results				
Adenovirus	Type 1	Isolate	1 x 10 ^{7.53} U/mL	Neg, Neg, Neg				
Coronavirus	229e	Isolate	1 x 10 ^{6.10} U/mL	Neg, Neg, Neg				
		Isolate	9.55 x 10 ⁶					
Coronavirus	OC43		$TCID_{50}/mL$	Neg, Neg, Neg				
Coronavirus	NL63	Isolate	1 x 10 ^{4.67} U/mL	Neg, Neg, Neg				
	Florida/USA-	Isolate						
	2_Saudia		4.17×10^5					
MERS-CoV (heat-inactivated)	Arabia_2014		TCID ₅₀ /mL	Neg, Neg, Neg				
		Inactivated						
SARS -1	2003-00592	virus	Not available	Neg, Neg, Neg				
Mycoplasma pneumoniae	M129	Isolate	3 x 10 ⁷ CCU/mL	Neg, Neg, Neg				
Streptococcus pyogenes	Z018	Isolate	3.8 x 10 ⁹ cfu/mL	Neg, Neg, Neg				
Influenza A H3N2	Brisbane/10/07	Isolate	1 x 10 ^{5.07} U/mL	Neg, Neg, Neg				
	New	Isolate						
Influenza A H1N1	Caledonia/20/99		1 x 10 ^{6.66} U/mL	Neg, Neg, Neg				
Influenza B	Brisbane/33/08	Isolate	1 x 10 ^{5.15} U/mL	Neg, Neg, Neg				

Table 11. Cross-reactivity test	Tesuits	Source/		
Virus/Bacteria/Parasite	Strain	Sample type	Concentration	Results
Parainfluenza	Type 1	Isolate	1 x 10 ^{8.01} U/mL	Neg, Neg, Neg
Parainfluenza	Type 2	Isolate	1 x 10 ^{6.34} U/mL	Neg, Neg, Neg
		Isolate	8.51 x 107	
Parainfluenza	Type 3		TCID50/mL	Neg, Neg, Neg
Parainfluenza	Type 4b	Isolate	1 x 10 ^{7.53} U/mL	Neg, Neg, Neg
Enterovirus	Type 68	Isolate	1 x 10 ^{6.5} U/mL	Neg, Neg, Neg
Human Metapneumovirus	A1 (IA10-s003)	Isolate	1 x 10 ^{5.55} U/mL	Neg, Neg, Neg
Respiratory Syncytial Virus	Type A (3/2015 Isolate #3)	Isolate	1 x 10 ^{5.62} U/mL	Neg, Neg, Neg
		Inactivated		
Human Rhinovirus	N/A	virus	Not available	Neg, Neg, Neg
Chlamydophila pneumoniae	AR-39	Isolate	2.9 x 107 IFU/mL	Neg, Neg, Neg
Haemophilus influenzae	Type b; Eagan	Isolate	7.87 x 10 ⁸ cfu/mL	Neg, Neg, Neg
Legionella pneumophila	Philadelphia	Isolate	6.82 x 10 ⁹ cfu/mL	Neg, Neg, Neg
Streptococcus pneumoniae	Z022; 19f	Isolate	2.26 x 10 ⁹ cfu/mL	Neg, Neg, Neg
Bordetella pertussis		Isolate		Neg, Neg, Neg
Pneumocystis jirovecii-S. cerevisiae Recombinant	W303-Pji	Isolate	1.56 x 10 ⁸ cfu/mL	Neg, Neg, Neg
Negative Nasopharyngeal Matrix	MTM	N/A	N/A	Neg, Neg, Neg
Negative Nasopharyngeal		N/A		
Matrix	MTM		N/A	Neg, Neg, Neg
Negative Nasal Matrix	CDC Viral Transport	N/A	N/A	Neg, Neg, Neg
Negative Oropharyngeal Matrix	CDC Viral Transport	N/A	N/A	Neg, Neg, Neg

The *in silico* analysis focused on 32 micro-organisms identified by the FDA as high priority for assessment due to their potential presence in upper respiratory samples.

TABLE 12. Cross-Reactivity Or	gamsms	T	
Organism	Total # Sequences	# Complete Genomes	# WGS Strains
Adenovirus	532	532	0
Coronavirus (Seasonal)	288	288	0
Enterovirus ^B	2708	2674	34
		21444 (+39	
Influenza A Virus ^{A B}	172455	A/Mexico/4108/2009)	108
		6755 (+16	
Influenza B Virus ^{A B}	53952	B/Florida/4/2006)	0

Organism	Total # Sequences	# Complete Genomes	# WGS Strains
Influenza C Virus ^B	2205	N/A	N/A
Human Metapneumovirus	145	145	0
Human Parainfluenza Virus 1-4	439	439	0
Human Parechovirus	124	124	0
Human Respiratory Syncytial Virus ^B	1275	1275	0
Rhinovirus	214	214	0
SARS-1	236 ^c	232 (+4 pp1ab sequences)	0
Bacillus anthracis	4152	69	86
Candida albicans	1541	59	34
Chlamydia pneumoniae	466	5	20
Chlamydia psittaci	11179	23	45
Corynebacterium diptheriae	20797	17	194
Coxiella burnetii	419	28	3
Haemophilus influenzae	45267	61	692
Legionella ^B	4843	98	65
Leptospira ^B	64456	133	266
Moraxella catarrhalis ^B	8333	11	184
Mycobacterium tuberculosis	194	194	0
Mycoplasma pneumoniae	808	51	45
Neisseria elongata & N. meningitidis ^B	312050	116	1318
Pneumocystis jirovecii	487	15	3
Pseudomonas aeruginosa	195	195	0
Staphylococcus aureus	634	634	0
Staphylococcus epidermidis ^B	61880	23	508
Streptococcus pneumoniae B	1633369	107	8526
Streptococcus pyogenes B	46153	201	1733
Streptococcus salivarius ^B	9417	18	48

^A Genome counts for Influenza A and Influenza B were attained for strains that included all 8 segments, except for A/Mexico/4108/2009(H1N1) and B/Florida/4/2006; all available gene sequences were included.

The in-silico analysis demonstrated < 80% homology with all organisms except for the following:

three Enterovirus sequences are 80.9% conserved to the reverse primer, however, the forward

453 primer is only 76% conserved and the probe alignment had an overall homology of 56%. The SARS-1

^B For BLAST, 'Max Target Seqs' was set to 5000. See Table 2.

^c 4 polyprotein cds sequences were also included.

sequences are ≥80% conserved to both primers, however, the last base on the 3' ends of both primers are not conserved. The wet testing of the only available SARS-1 strain was non-detectable.

456 457

458

459

460

461

462

463

464

465

466

467

468

469

472

473

474

475 476

477

478

454

455

Limitations

- Negative results do not preclude infection with SARS-CoV-2 and should not be the sole basis of a patient treatment decision.
- This test is intended to be used for the detection of SARS-CoV-2 RNA in nasopharyngeal and oropharyngeal swab samples. Testing of other sample types may result in inaccurate results.
- Nasal swabs and mid-turbinate nasal swabs are considered acceptable specimen types for use
 with the Lyra SARS-CoV-2 Assay but performance with these specimen types has not been
 established. Testing of nasal and mid-turbinate nasal swabs (self-collected under supervision of
 or collected by a healthcare provider) is limited to patients with symptoms of COVID-19. Please
 refer to FDA's FAQs on Diagnostic Testing for SARS-CoV-2 for additional information.
- Improper collection, storage or transport of specimens may lead to false negative results.
- Inhibitors present in the sample and/or errors in following the assay procedure may lead to false negative results.
- A trained health care professional should interpret assay results in conjunction with the
 patient's medical history, clinical signs and symptoms, and the results of other diagnostic tests.
 - Analyte targets (viral sequences) may persist *in vivo*, independent of virus viability. Detection of analyte target(s) does not imply that the corresponding virus(es) are infectious, nor are the causative agents for clinical symptoms.
 - There is a risk of false positive values resulting from cross-contamination by target organisms, their nucleic acids or amplified product, or from non-specific signals in the assay.
 - There is a risk of false negative values due to the presence of sequence variants in the viral targets of the assay.
 - The assay performance was not established in immunocompromised patients.

479 480 481

482

483

484

Conditions of Authorization for the Labs

The Lyra® SARS-CoV-2 Assay 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/MedicalDevices/Safety/ EmergencySituations/ucm161496.htm.

485 H 486 A

However, to assist clinical laboratories using the Lyra® SARS-CoV-2 Assay, the relevant Conditions of Authorization are listed below.

487 488 489

- Authorized laboratories¹ using the Lyra SARS-CoV-2 Assay will include with result reports of the Lyra SARS-CoV-2 Assay test, 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 Lyra SARS-CoV-2 Assay will perform the Lyra SARS-CoV-2 Assay as outlined in the Lyra SARS-CoV-2 Assay 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 Lyra SARS-CoV-2 Assay are not permitted.

- Authorized laboratories that receive the Lyra SARS-CoV-2 Assay must notify the relevant public health
 authorities of their intent to run the test prior to initiating testing.
 - Authorized laboratories using the Lyra SARS-CoV-2 Assay will have a process in place for reporting test results to healthcare providers and relevant public health authorities, as appropriate.
 - 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 Quidel
 (QDL.COV2.test.event.report@quidel.com) 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.
 - 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.
 - Quidel, its authorized distributor(s) and authorized laboratories using the Lyra SARS-CoV-2 Assay 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.

¹For ease of reference, the letter of authorization refers to, "United States (U. S.) laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, to perform high complexity tests" as "authorized laboratories."

Customer and Technical Assistance

To place an order or for technical support, please contact a Quidel Representative at (800) 874-1517 (toll-free in the U.S.) or (858) 552-1100 (outside of U.S.), Monday through Friday, between 8:00 a.m. and 5:00 p.m., Eastern Time. Orders may also be placed by fax at (740) 592-9820. For e-mail support contact: customer service@dhiusa.com or technical_services@dhiusa.com. For services outside the U.S., please contact your local distributor. Additional information about Quidel, our products, and our distributors can be found on our website quidel.com.

NucliSENS and easyMAG are registered trademarks of bioMérieux, Inc. TaqMan is a registered trademark of Roche. Applied Biosystems® is a registered trademark of Life Technologies. LightCycler® 480 is a registered trademark of Roche. Rotor-Gene is a registered trademark of Qiagen. Q Dye compounds in this product are sold under license from BioSearch Technologies, Inc. and protected by U.S. and world-wide patents either issued or under application. The license covers R&D use and human in vitro diagnostic (IVD) applications.

Quidel Corporation

Lyra® SARS-CoV-2 Assay 4/06/2020 Page 24 of 44

528	References
529	1. Baker, S., Frias, L., and Bendix, A. Coronavirus live updates: More than 92,000 people have been infected
530	and at least 3,100 have died. The US has reported 6 deaths. Here's everything we know. Business Insider.
531	March 03, 2020.
532	2. Clinical and Laboratory Standards Institute. Viral Culture; Approved Guidelines. CLSI document M41-A
533	[ISBN 1562386239] Clinical and Laboratory Standards Institute, 940 West Valley Road, Suite 1400, Wayne,
534	Pennsylvania 19087-1898, USA 2006.
535	3. Lauer, S.A., et. al. The incubation period of Coronavirus disease 2019 (COVID-19) from publicly reported
536	confirmed cases: estimation and application, Ann Intern Med. 2020
537	4. <u>www.cdc.gov/coronavirus/2019-ncov/about/symptoms.html</u>

538

540 **APPENDIX** Applied Biosystems 7500 Fast Dx Programming Instructions 541 542 Refer to User Manual Part Number 4406991 for additional information. 543 1. Launch the 7500 Fast Dx software package. 544 2. The Quick Startup document dialog window will open. Select the Create New Document button to start 545 the New Document Wizard. Follow each step to initiate the Lyra™ SARS-CoV-2 Assay protocol. 546 <u>Define Document</u>: Most of the following should be the default setting. If not, change accordingly. 547 i. Confirm or enter the following information. Standard Curve (Absolute Quantitation) Assay: Container: 96-Well Clear Template: Blank Document Run Mode: Fast 7500 Operator: your operator name SDS v1.4 Comments: Plate Name: 'Lyra SARS-CoV-2 Assay' 548 ii. Select the Next button. 549 550 b. Select Detectors: New detectors for SARS-CoV-2 and the process control (PRC) must be added. For 551 each target, select the New Detector button to open the New Detector pop-up window. 552 Alternatively, use the Create Another button from within the New Detector pop-up window for 553 the last two detectors. 554 555 i. Enter the following information for each detector. Quencher Dve Name Reporter Dve Color SARS-CoV-2 FAM (Select) (none) PRC Quasar 670 (none) (Select) 556 557 ii. Select a unique color to represent each detector. 558 iii. Highlight the new detectors and add to the **Detectors in Document** column using the **Add** 559 button. 560 iv. Select (none) from the Passive Reference drop-down menu. 561 v. Select the Next button. 562 vi. Select the Finish button without setting any wells. 563 c. The wizard will close and the software will open, starting with the **Setup** tab. This will show the 564 sample plate that was set up during the quick start. For the initial set up, nothing needs to be 565 changed here. 566 d. <u>Defining the Thermocycler Protocol:</u> Select the **Instrument** tab to set up the Lyra™ SARS-CoV-2 567 Assay RT-PCR cycling times and temperatures. Under Thermal Profile there should be a default 2-568 stage protocol. Each stage will have 3 user-editable text boxes. The top box value represents the 569 number of reps or cycles for that stage. The middle box value represents the temperature (°C) and 570 the lowest box value represents the time (minutes: seconds). 571

i. Make the following changes to the default **Thermal Cycler Protocol**:

1. Stage 1

572

-										
1			a.	Reps:	1					
5			b.	Temp:	55					
5			c.	Time:	5:0	0				
		2.	Select tl	he bar be	etween S	tage 1 an	d Stage 2	. Select th	e Add Hold	button to add
			another							
		3.	Stage 2	_						
			a.	Reps:	1					
			b.	Temp:	60					
			c.	Time:	5:0	0				
		4.	Select tl	he bar be	etween S	tage 2 an	d Stage 3	. Select th	e Add Hold	button to add
			another							
		5.	Stage 3	J						
			a.	Reps:	1					
			b.	Temp:	65					
			c.	Time:	5:0	0				
		6.	Stage 4			on Stage)				
			a.	Reps: 1		σ,				
			b.	Step 1						
				i.	Temp:	92				
				ii.	Time:	0:0	5			
			c.	Step 2						
				i.	Temp:	57				
				ii.	Time:	0:40	0			
		7.	Select th	he bar to	the right	of Stage	4. Select t	he Add C y	cle button t	o add another
			stage.					-		
		8.		(2-Step [Dissociati	on Stage)				
			a.	Reps: 3		σ.				
			b.	Step 1						
				i.	Temp:	92				
				ii.	Time:	0:0	5			
			c.	Step 2						
					Temp:	57				
				ii.	Time:	0:40	0			
		9.	If a wro	ng stage i				noved by i	oressing the	Delete button
								tical lines		
	ii.	Under S	ettings e			-				
			ample Vo			20 (defa	uul+)			
			un Mode		-1.		st (defaul	+1		
							-	-	11	
			ata Colle		مام ماهدام			7.0 @ 0:40		
		N	OTE: DO	not cne	ck the ch	еск рох п	ext to Ex	pert Mod	e.	
	6.1.11	ا اداما	!-							
			each and	-						
			he Result		Diat tele					
	ii.	select ti	he Ampli	tication l	Pi ot tab.					

iii. Select SARS-CoV-2 from the Detector tab in the top right corner.

615		iv.	In the Analysis Settings block, set the Threshold to 7.5e+004 .
616		٧.	Select the Auto Baseline radio button.
617		vi.	Select PRC from the Detector tab in the top right corner.
618		vii.	In the Analysis Settings block, set the Threshold to 1.0e+004 .
619		viii.	Select the Auto Baseline radio button.
620			
621	f.	Save the	e new protocol as a template for future use.
622		i.	At the top of the screen select File and then Save As.
623		ii.	Save In: D:\Applied Biosystems\7500 Fast System\Templates\
624		iii.	File name: 'Lyra SARS-CoV-2'
625		iv.	Save as type: 'SDS Templates (*.sdt)'
626	g.	Exit the	software.

Applied Biosystems® 7500 Fast Dx Thermocycler Test Procedure

- 1. Launch the Applied Biosystems® 7500 Fast Dx software v1.4 package.
- 2. The **Quick Startup document** dialog window will open.
- 630 3. Click on Create a new document.

627

628

631

632

633

634

635

636

637

638

639 640

641

642

643

644

645

646

647

648

4. Most of the following should be the default setting. If not, change accordingly.

Assay:	Standard Curve (Absolute Quantitation)
Container:	96-Well Clear
Template:	Lyra SARS-CoV-2
Run Mode:	Fast 7500
Operator:	your operator name
Comments:	SDS v1.4
Plate Name:	YYMMDD- Lyra SARS-CoV-2

- Set Up Sample Plate
 - a. Under the **Setup** and **Plate** tabs the plate setup will appear.
 - Select all wells that will contain sample, right-click and select the Well Inspector from the dropdown menu. When the Well Inspector pop-up window opens, select the detectors for SARS-CoV-2 and PRC.
 - c. Use the **Well Inspector** to enter the sample names. Patient IDs can be entered in the Well Inspector window. However, it is recommended that this is done prior to re-suspending the lyophilized master mix, post run or using the import function to minimize the time the PCR reactions will sit at room temperature prior to starting the run.
 - d. Save the run as YYMMDD- Lyra SARS-CoV-2.sds.
 - e. A window will open asking for the "Reason for change of entry". Enter "**Setup**" and any other comments relevant to the run.
 - 6. Starting the PCR
 - a. Select the **Instrument** tab.
 - b. Insert the 96 well PCR plate into the machine.
- c. Under **Instrument Control**, select the **Start** button to initiate the run.
- Post PCR
- 649 **IMPORTANT:** When the run is finished press OK.
- 650 a. Analyze the data by pressing the "Analyze" button in the top menu and save the file.

651	b.	Save the file	by pressing Sa	ave Document in	the task bar. A win	dow will o	pen asking for the
652		"Reason for	change of enti	ry".			
653	C.	Enter " Data	analysis post	run" and any othe	er comments relev	ant to the	run.
654	Applied B	iosystems 7	7500 Standa	rd Programmi	ng Instructions		
655	Refer to Us	ser Manual P	art Number 4	1387783 rev C fo	or additional info	rmation.	
656	1. Launch t	he ABI 7500	software pag	ckage.			
657	2. Select th	e Advanced	Setup buttor	n to open Setup	and Experiment	Propertie	s. Follow each step to
658	initiate the	Lyra SARS-C	CoV-2 protoco	ol.			
659	a. l	Experiment I	Name: Enter I	Experiment Nan	ne as SARS-CoV-2	. Leave th	ne Barcode, User
660	Na	me, and Con	nments fields	blank			
661	b.	Define Exper	iment Setup:	Select 7500 (96	Wells), Quantita	ition- Star	ndard Curve, TaqMan [©]
662	Re	agents, and	Standard (~2	hours to comple	ete a run)		
663		•	nu select Plat	•			
664		•	ts: New dete	ctors for SARS-C	oV-2, and the pro	ocess con	trol (PRC) must be
665	ad	ded.					
666		i. Enter	the following	g information fo	r each detector.		
667		i				0.1	l
			Name	Reporter Dye	Quencher Dye	Color	
			SARS-CoV-2 PRC	FAM Quasar 670	(none) (none)	(Select)	
668			FNC	Quasar 070	(Horie)	(Select)	
669		ii. Seled	ct Add New T	arget button fo	r each target.		
670				_	ct reporter, quer	ncher and	1 color
671			-	olor to represen		icrici, aric	7 60101
672	h		•	-		left corne	er, select none as the
673		ssive Refere	-	cs. Officer this to	ib iii tiic bottoiii	icit corric	ir, sciect none as the
674			rom the uppe	er left menu			
675					L under the Gra	hical or 1	Гabular View
676				•	•		View the default
677	pro	ofile should b	oe 2 holding s	stages and a 2-st	ep cycling protoc	col. Each s	stage will have 3 user-
678	ed	itable text bo	oxes. The first	t box value repr	esents the Ramp	Rate (%)	for that stage, the
679			•	s the temperatu	re (°C) and the th	ird box va	alue represents the
680	tim	ne (minutes:	seconds).				
681							
682					e default Thermo	cycler pro	otocol:
683		1. Stag	e 1 First Hold	•			
684			a. Ramp Rat				
685			b. Temp: 55				
686			c. Time: 5:00				
687		2. Step	1 Second Ho				
688			a. Ramp Rat				
689			b. Temp: 60				
690			c. Time: 5:00	0			

691 692	3. Highlight the second Holding Stage and select the Add Stage button. In the drop down menu select Holding
693	
694	4. Step 1 Third Holding Stage a. Ramp Rate: 100%
695	b. Temp: 65
696	c. Time: 5:00
697	5. First 2-Step Cycling Stage
698	a. Number of cycles: 10
699	b. Do NOT check Enable Auto Delta
700	c. Step 1
701	i. Ramp Rate: 100%
702	ii. Temp: 92
703	iii. Time: 0:05
704	d. Step 2
705	i. Ramp Rate: 100%
706	ii. Temp: 57
707	iii. Time: 0:40
708	iv. Turn data collection "Off" by selecting the Data Selection button
709	at the bottom of the step.
710	6. Highlight step 2 and select the Add Stage button. In the drop down menu select
711	Cycling
712	7. Second 2-Step Cycling Stage
713	a. Number of cycles: 30
714	b. Do NOT check Enable Auto Delta
715	c. Step 1
716	i. Ramp Rate: 100%
717	ii. Temp: 92
718	iii. Time: 0:05
719	d. Step 2
720	i. Ramp Rate: 100%
721	ii. Temp: 57
722	iii. Time: 0:40
723	iv. Ensure the data collection has been turned "On" for this step
724	(default setting)
725	8. If a wrong stage is added the stage can be removed by pressing the Undo "Add
726	Stage" button immediately after adding the stage or highlight the stage between
727 728	the vertical lines and select the Delete Selected button
729	5. Set threshold for each analyte
730	a. Select the Analysis tab in the upper left menu.
731	b. Select Analysis Settings button in the top right corner.
732	c. Highlight SARS-CoV-2 and deselect the Use Default Settings box. De-select Automatic
733	Threshold and change threshold to 75,000. Leave Automatic Baseline selected.

734	d. Highlight PRC and de-select the Use Default Settings box. De-select Automatic Threshold
735	and change threshold to 10,000. Leave Automatic Baseline selected.
736	e. At the bottom of the box select Apply Analysis Settings button
737	

Target	Threshold	Baseline Start	Baseline End
SARS-CoV-2	75,000	Auto	Auto
PRC	10,000	Auto	Auto

740

741

742

744

747

748

750

752

753

754

755

756

757

758

759

760

761

762

763

764

765

i. Save the new protocol as a template for future use.

- i. At the top of the screen select the drop down menu next to Save
- ii. Choose **Save as Template**
 - iii. Save in an appropriate folder
- 743 iv. **File name:** 'Lyra SARS-CoV-2'
 - v. Save as type: 'Experiment Document Template files (*.edt)'
- 745 vi. Exit the software.

746 Applied Biosystems® 7500 Standard Thermocycler Test Procedure

- 1. Launch the Applied Biosystems® 7500 Standard software v2.06 package.
- 2. The **Quick Startup document** dialog window will open.
- 749 3. Click on Create a new document.
 - 4. Most of the following should be the default setting. If not, change accordingly.

Assay:	Standard Curve (Absolute Quantitation)
Container:	96-Well Clear
Template:	Lyra SARS-CoV-2
Run Mode:	Fast 7500
Operator:	your operator name
Comments:	SDS v1.4
Plate Name:	YYMMDD- Lyra SARS-CoV-2

- 751 5. Set Up Sample Plate
 - a. Under the **Setup** and **Plate** tabs the plate setup will appear.
 - Select all wells that will contain sample, right-click and select the Well Inspector from the dropdown menu. When the Well Inspector pop-up window opens, select the detectors for SARS-CoV-2 and PRC.
 - c. Use the **Well Inspector** to enter the sample names. Patient IDs can be entered in the Well Inspector window. However, it is recommended that this is done prior to re-suspending the lyophilized master mix, post run or using the import function to minimize the time the PCR reactions will sit at room temperature prior to starting the run.
 - d. Save the run as YYMMDD- Lyra SARS-CoV-2.sds.
 - e. A window will open asking for the "Reason for change of entry". Enter "**Setup**" and any other comments relevant to the run.
 - 6. Starting the PCR
 - a. Select the **Instrument** tab.
 - b. Insert the 96 well PCR plate into the machine.
- 766 c. Under **Instrument Control**, select the **Start** button to initiate the run.

/6/	7. Po	ost PCR
768	IN	IPORTANT: When the run is finished press OK.
769	a.	Analyze the data by pressing the "Analyze" button in the top menu and save the file.
770	b.	Save the file by pressing Save Document in the task bar. A window will open asking for the
771		"Reason for change of entry".
772	C.	Enter "Data analysis post run" and any other comments relevant to the run.
773	Bio-Rad (CFX96 Touch Thermocycler Programming Procedure
774		Iser Manual Part Number 10010424 Rev D for additional information.
//4	helel to C	sel Mandal Fait Number 10010424 Nev D for additional information.
775	Programm	ing Instructions:
776	1. Launcl	n the CFX96 Touch software package
777	2. In the	Startup Wizard pop-up window Select instrument to be CFX96 from the drop down menu
778	3. Under	Select Run Type press the User-defined button
779	4. Create	a new thermocycler protocol by selecting Create New from the Run Setup window
780	5. Make	the following changes to the cycling conditions in the Protocol Editor:
781	a.	Change the Sample Volume to 20ul
782	b.	Under Tools in the top left toolbar select Run Time Calculator and check 96 Wells-All Channels
783	C.	Step 1 (Hold)
784		i. Reps: 1
785		ii. Temp: 55C
786		iii. Time: 5:00
787	d.	Step 2 (Hold)
788		i. Reps: 1
789		ii. Temp: 60C
790		iii. Time: 5:00
791	e.	Step 3 (Hold)
792		i. Reps: 1
793		ii. Temp: 65C
794		iii. Time: 5:00
795		iv. Remove the plate read from this stage by selecting the Remove Plate Read button on the
796		lower left
797	f.	Step 4 (2-Step Amplification Stage)
798		i. Highlight step 3 and go to the lower left of the window and select Insert Step for a total
799		of 2 times until step 5 is reached (ensure in the upper left of the window the drop-down
800		menu for Insert Step has After selected).
801		ii. Highlight step 4 and set as follows:
802		1. Temp: 92C
803		2. Time: 0:05
804		iii. Highlight step 5 and set as follows:
805		1. Temp: 57C
806		2. Time: 0:40
807		3. Go to the left of the screen and select Remove Plate Read button

808		iv. Select step 6, the GOTO step, and change to state GOTO step 4 and change the times to
809		repeat to 9
810	g.	Step 7 (2-Step Amplification Stage)
811		i. With step 6 highlighted select Insert Step button, on the lower left of the window, for a
812		total of 2 times (until step 8 is reached)
813		ii. Highlight step 7 and set as follows:
814		1. Temp: 92C
815		2. Time: 0:05
816		iii. Highlight step 8 and set as follows:
817		1. Temp: 57C
818		2. Time: 0:40
819		3. In the left of the window select Add Plate Read to Step button
820		4. Highlight step 8 and select Insert GOTO button on the lower left of the window
821		iv. Select step 9, the GOTO step, and change to GOTO step 7 and times to repeat to 29
822	h.	Save the new cycling conditions as protocol for future use
823		i. At the upper left of the screen select the Save button
824		ii. Save in the ExpressLoad folder
825		iii. Name the file 'Lyra SARS-CoV-2'
826		iv. Save as type 'Protocol File (*.prcl)'
827		v. Select Save
828		vi. Click Ok in the protocol editor window
829	6. Define	the plate setup
830	a.	In the Run Setup window select the Plate tab
831	b.	Under Express Load in the drop-down menu select Quick Plate 96 wells All Channels.pltd
832	C.	Select the Edit Selected button to customize the plate setup
833	d.	In the upper toolbar select Settings. The default settings need to be set.
834		i. Plate Size select 96 Wells
835		ii. Plate Type select BR Clear
836		iii. Number Convention select Scientific Notation
837		iv. Units select Copy Number
838	e.	Leave the Scan Mode set to All Channels at the top of the window
839	f.	Select the Select Fluorophores button on the upper right of the Plate Editor window
840		i. De-select all default fluorophores
841		ii. Select FAM , and Cy5 and click Ok
842	g.	In the Plate Editor window highlight the whole plate and click the check box in front of all
843		fluorophores: FAM and Cy5
844	h.	Select the Experiment Settings button in order to define the Targets
845		i. In the lower left of the Experiment Settings window in the New box type in SARS-CoV-2
846		and select Add
847		ii. Repeat this for the PRC
848		iii. Select Ok
849	i.	In the Plate Editor window next to FAM in the drop-down menu under Target Name select SARS
850		CoV-2 and for Cy5 select PRC
851	j.	Save the new plate setup for future use
852		i. At the upper left of the screen select the Save button

853	ii. Save in the ExpressLoad folder
854	iii. Name the file 'Lyra SARS-CoV-2 plate'
855	iv. Save as type 'Plate File (*.pltd)'
856	v. Select Save
857	vi. Click Ok in the Plate Editor window
858	k. Exit the software
859	Bio-Rad CFX96 Touch Thermocycler Test Procedure
860	Analysis Instructions:
861	1. Open the run file that needs to be analyzed
862	2. In the upper left select the Quantification Tab
863	3. On the Amplification curve check the box in front of Log Scale
864	4. Select Settings in the toolbar in the upper left of the screen
865	a. For the Cq Determination Mode select Single Threshold
866	b. Under the Baseline Setting choose Baseline Subtracted Curve Fit
867	c. For Analysis Mode select Target
868	d. Under Cycles to Analyze choose 1-30 and then click Ok
869	e. The baseline cycles and the threshold for each target need to be set
870	i. Ensure that only the SARS-CoV-2 box is checked in the amplification plot
871	Go up to Settings in the toolbar and select Baseline Threshold
872	 At the top of the box select Auto Calculated for the Baseline Cycles
873	2. For the Single Threshold at the bottom of the box select User Defined
874	a. Set this to 164
875	b. Select Ok
876	iii. Uncheck the SARS-CoV-2 box and check the PRC box in the amplification plot
877	iv. Go up to Settings in the toolbar and select Baseline Threshold
878	1. At the top of the box select Auto Calculated for the Baseline Cycles
879	2. For the Single Threshold at the bottom of the box select User Defined
880	a. Set this to 100
881	b. Select Ok
882	5. Exit the software
883	Qiagen Rotor-Gene Q Programming Instructions
884	Refer to User Manual Part Number 1065453EN for additional information.
885	Programming Instructions:
886	Launch the Rotor-Gene Q software package
887	2. In the New Run pop-up window select the Advanced tab on the top of the screen
888	3. Select Empty Run and then New on the lower right of the pop-up window to start the Advanced Run Wizard
889	a. Select the appropriate rotor size in the Advanced Run Wizard on the upper left of the screen
890	b. Check the box that states the Locking Ring is Attached and select Next

c. Leave the **Operator** and **Notes** sections empty

892	d.	Enter 20ul as the Reaction Volume in the lower left of the screen
893	e.	For the Sample Layout choose 1, 2, 3 and then select Next
894	f.	Under Channel Setup select Create New to enter information for each detector
895		i. Under Name enter SARS-CoV-2
896		ii. Source select 470nm
897		iii. Detector select 510nm
898		iv. Do not adjust the default Gain setting of 7 as this will be set in a later step
899		v. Select OK
900	g.	Repeat the step above by selecting Create New
901		i. Under Name enter PRC
902		ii. Source select 625nm
903		iii. Detector select 660nm
904		iv. Do not adjust the default Gain setting of 7 as this will be set in a later step
905		v. Select OK
906	h.	Select the Edit Profile button in the middle of the window to setup a cycling profile
907		i. In the Edit Profile window go to the upper left of the screen to New and in the drop-down
908		menu select Cycling. A hold and three step cycling stage should appear.
909		ii. Modify the hold stage to have a temperature at 55°C and a time of 5:00 minutes
910		iii. Select the Insert After button in the middle of the pop-up window and then select New
911		Hold at Temperature
912		iv. Modify the second hold stage to have a temperature at 60°C and a time of 5:00 minutes
913		v. Select the Insert After button in the middle of the pop-up window and then select New
914		Hold at Temperature to insert a third hold stage
915		vi. Modify the third hold stage to have a temperature at 65°C and a time of 5:00 minutes
916		vii. Highlight the first cycling stage and modify it as follows:
917		1. This cycle repeats 10 time(s)
918		2. Select Timed Step from the drop-down menu in the middle left of the screen
919		3. Do not select Long Range or Touchdown on the left of the screen
920		4. The first step:
921		a. 92°C
922		b. 5 seconds
923		c. Not Acquiring
924		5. Select step two and set as follows:
925		a. 57°C
926		b. 40 seconds
927		c. Not Acquiring
928		6. Highlight step three and delete it by selecting the "-" button in the middle of the
929		window
930		7. Select the Insert After button in the middle of the pop-up window and then
931		select New Cycling
932		viii. Highlight the second cycling stage and modify it as follows:
933		1. This cycle repeats 30 time(s)
934		2. Select Timed Step from the drop-down menu in the middle left of the screen
935		3. Do <u>not</u> select Long Range or Touchdown on the left of the screen
936		4. The first step:

937	a. 92°C
938	b. 5 seconds
939	c. Not Acquiring
940	5. Select step two and set as follows:
941	a. 57°C
942	b. 40 seconds
943	c. Select Acquiring to Cycling A
944	i. Under Acquiring Channels highlight the default channel name
945 946	(Green) and select the < button to move it over to the Available Channels list
947	
	ii. In the Available Channels list select SARS-CoV-2 and select the
948	> button to move it over to the Acquiring Channels list
949	iii. Repeat the step above for the PRC and then select OK
950 951	6. Highlight step three and delete it by selecting the "-" button in the middle of the window
952	ix. In the Edit Profile window select OK
953	i. In the New Run Wizard window select Gain Optimisation
954	i. In the middle of the Auto-Gain Optimisation Setup window select the drop-down menu
955	under Channel Settings and select SARS-CoV-2.
956	ii. Select the Add button on the right
957	1. In the Auto-Gain Optimisation Channel Settings window ensure that the SARS-
958	CoV-2 Tube Position is set to 1. This requires that a positive control, containing
959	SARS-CoV-2 and PRC, be tested with each PCR run and placed in the first tube.
960	Failure to do so may cause the gain to be incorrectly set.
961	2. Leave the Target Sample Range and the Acceptable Gain Range set to the
962	defaults, 5-10Fl and -10 to 10 respectively.
963	3. Select OK
964	4. Repeat steps 3. j. ii. 1-3. for the PRC
965	iii. In the Auto-Gain Optimisation Setup window check the box next to Perform
966	Optimisation Before 1 st Acquisition
967	iv. Select Close
968	j. In the New Run Wizard window select the Next button
969	k. Save the new protocol as a template for future use
970	 i. On the bottom right of the window select the Save Template button
971	ii. Save In: C:\Program Files\Rotor-Gene Q Software\Templates
972	iii. File name: 'Lyra SARS-CoV-2'
973	<pre>iv. Save as type: 'Template (*.ret)'</pre>
974	I. Exit the software
975	Qiagen Rotor-Gene Q Test Run
976	Analysis Instructions:
977	1. In the New Run Wizard load the SARS-CoV-2 Template.
	2. Press Start.
979	
977 978 979	2. Press Start.

4. In the upper menu toolbar select the **Analysis** button

981		a.	Select Quantitation, then Cycling A. SARS-CoV-2, and Show
982		b.	The threshold needs to be set for SARS-CoV-2
983			i. In the far right bottom of the screen under CT Calculation enter 0.03 for the SARS-CoV-2
984			Threshold
985			ii. In the Eliminate Cycles before box ensure the default of 1 is entered
986			iii. Ensure the amplification graph is set to Log Scale (toggle button on the bottom left of the
987			graph states Linear Scale or Log Scale)
988		c.	Select Quantitation, then Cycling A. PRC, and Show
989		d.	The threshold needs to be set for PRC
990			i. In the far right bottom of the screen under CT Calculation enter 0.05 for the PRC
991			Threshold
992			ii. In the Eliminate Cycles before box ensure the default of 1 is entered
993			iii. Ensure the amplification graph is set to Log Scale (toggle button on the bottom left of the
994			graph states Linear Scale or Log Scale)
005	р.	ala a/a 1:	
995			ghtCycler® 480 Instrument II Programming Instructions
996	Re	fer to Us	ser Manual Part Number 05152062001 0208 for additional information.
997	Cre	eating a	LC 480 II Assay Run Template
998	1.	Launch	the LightCycler (LC) 480 software package
999	2.	The Det	tection Format must be established to specify the channels in which fluorescence will be read
1000		a.	Select Tools in the startup screen in the lower right of the screen
1001		b.	Select Detection Formats then choose New
1002		c.	Name the format Lyra ® SARS-CoV-2
1003		d.	In the Filter Combination Selection window select 465-510 and 618-660
1004		e.	In the Selected Filter Combination List window under name type in SARS-CoV-2 for 465-510 and
1005			PRC for 618-660
1006		f.	Leave all default setting values to 1 under Melt Factor, Quant Factor, and Max Integration Time
1007		g.	Select Close to save the new detection format and return to startup screen
1008		h.	To access this newly created Detection Format , the LC 480 software must be closed, then reloaded
1009	3.	After cl	osing and reloading the software select White Plates and New Experiment under Experiment
1010		Creation	n window
1011	4.	On the	next screen select "Lyra ® SARS-CoV-2" from the pull-down menu under Detection Formats
1012	5.	Enter 20	Oul as the Reaction Volume in the upper right of the screen
1013	6.	Enter th	ne names for each of the RT-PCR programs
1014		a.	Under Program Name enter Stage 1, under Cycles enter 1, and in Analysis Mode select none
1015		b.	Select the "+" icon to add a program
1016		c.	Name the next program Stage 2 , under Cycles enter 1 , and in the Analysis Mode select none
1017		d.	Select the "+" icon to add a program
1018		e.	Name the next program Stage 3 , under Cycles enter 1 , and in the Analysis Mode select none
1019		f.	Select the "+" icon to add a program
1020		g.	Name the next program Stage 4, under Cycles enter 40, and in the Analysis Mode select
1021		-	quantification

1022	7.	Set the RT-PCR cycling times and temperatures
1023		a. Highlight Stage 1 under Program Name and change Stage 1 Temperature Targets as follows:
1024		i. Target (°C) set to 55
1025		ii. Acquisition Mode select none
1026		iii. Hold (hh:mm:ss) set to 5:00
1027		iv. Ramp Rate (°C/s) to 4.4
1028		v. Sec Target (°C), Step Size (°C), and Step Delay (cycles) will be left at 0 for stages 1-4.
1029		b. Highlight Stage 2 under Program Name and change Stage 2 Temperature Targets as follows:
1030		i. Target (°C) set to 60
1031		ii. Acquisition Mode select none
1032		iii. Hold (hh:mm:ss) set to 5:00
1033		iv. Ramp Rate (°C/s) to 4.4
1034		c. Highlight Stage 3 under Program Name and change Stage 3 Temperature Targets as follows:
1035		i. Target (°C) set to 65
1036		ii. Acquisition Mode select none
1037		iii. Hold (hh:mm:ss) set to 5:00
1038		iv. Ramp Rate (°C/s) to 4.4
1039		d. Highlight Stage 4 under Program Name and change Stage 4 Temperature Targets as follows:
1040		i. The first step:
1041		1. Target (°C) set to 92
1042		2. Acquisition Mode select none
1043		3. Hold (hh:mm:ss) set to 0:05
1044		4. Ramp Rate (°C/s) to 4.4
1045		ii. Select the "+" icon to add a step and set the second step:
1046		1. Target (°C) set to 57
1047		2. Acquisition Mode select single
1048		3. Hold (hh:mm:ss) set to 0:40
1049		4. Ramp Rate (°C/s) to 2.2
1050	8.	Save the new protocol as a run template for future use.
1051		a. In the lower left corner of the screen select the pull-down menu next to the Apply Template button
1052		b. Choose Save As Template
1053		c. Select the Templates Folder
1054		d. Highlight Run Templates Folder
1055		e. Name the template Lyra ® SARS-CoV-2 run template and click the "check" button
1056	9.	Exit the software.
1057	Cro	eating a LC 480 II Assay Test Procedure
1058	1.	Load the Lyra SARS-CoV-2 run template.
1059	2.	Press Start.
1060	3.	The analysis template can only be established after the initial experiment has completed
1061	4.	On the Lyra ® SARS-CoV-2 run select the Analysis button in the module bar
1062		a. Choose Abs Quant/Fit Points
		· ·

b. In the Create New Analysis pop-up window select your pre-defined subset from the subset drop

down menu and then select the "check" button

1063

1065	c. Set the Background to 2-10 for all analytes	
1066	i. Set Min Offset to 1	
1067	ii. Set Max Offset to 9	
1068	d. In the center bottom of the screen ensure that Color Compensation is off for all	analytes
1069	e. Leave the default settings as First Cycle 1 and Last Cycle 40	
1070	5. At the top middle of the screen select Noise Band	
1071	6. Choose the pull-down menu next to the Noise Band button and select Noise Band Fluore	scence
1072	7. For each analyte under the Filter Comb button, set the noise band as follows:	
1073	a. SARS-CoV-2set to 1.95	
1074	b. PRC set to 1.4619	
1075	8. Choose Calculate in the bottom left of the screen	
1076	9. Save the new analysis protocol as a template for future use	
1077	a. In the lower left corner of the screen select the pull-down menu next to the Apply	Template button
1078	b. Choose Save As Template	
1079	c. Select the Templates Folder	
1080	d. Highlight Analysis Templates Folder	
1081	e. Name the template Lyra ® SARS-CoV-2 analysis template and click the "check" be	utton
1082	10. Create a report	
1083	a. Select the Save icon on the global action bar on the right side of the screen	
1084	b. Choose the Report button on the module bar on the left of the screen	
1085	c. Select the appropriate settings and press the Generate button	
1086	11. To apply an Analysis Template to subsequent runs	
1087	a. Once the run has finished select the Analysis button in the module bar	
1088	b. Choose Abs Quant/Fit Points	
1089	c. In the Create New Analysis pop-up window select your pre-defined subset from	n the subset drop
1090	down menu and then select the "check" button	
1091	d. Select the Apply Template button on the far left of the screen and choose the L	yra ® SARS-CoV-2
1092	analysis template from the Analysis Templates Folder	
1093	e. Select yes in the pop-up window	
1094	12. Interpretation of results (See Table 4)	
1095	ThermoFisher QuantStudio 7 Pro Programming Instructions	
1096	Refer to User Manual Part Number 4489822 Revision A for additional information.	
1030	Refer to oser Manual Fart Number 4405022 Revision A for additional information.	
1097	ThermoFisher QS7 Test Run Programming Instructions:	
1098	1. Open the Design and Analysis Software	
1099	2. Select the "SET UP PLATE" option	
1100	3. From the side bar on the screen, select the following properties to filter:	
1101	a. Instrument – QuantStudio 7 Pro	
1102	b. Block – 96-Well 0.2 mL	
1103	c. Run Mode – Fast	

d. Analysis options are left blank

1105	4.	From the plate selections present on the screen, select the System Template "PCR Only" and
1106		the system will automatically navigate to the "Run Method" tab
1107	5.	
1108	•	a. Change the Reaction Volume to 20.0 uL
1109		b. The temperature of the enabled heated cover will remain at 105.0 degrees C
1110		c. Scroll over the Hold stage present in the cycling parameters and
1111		addition/subtraction buttons will become visible at both the top and bottom of the
1112		first stage.
1113		d. Left click the right addition button at the top and a list of Stage choices will become
1114		visible. Scroll down and choose Hold.
1115		e. Repeat the previous steps so there are three Hold stages present in the cycling
1116		parameters.
1117		f. Scroll over to the PCR stage and addition/subtraction buttons will become visible at
1118		both the top and bottom. Left click the right addition button at the top and a list of
1119		Stage choices will become visible. Scroll down and choose PCR.
1120		
1121		g. Going back to the first stage enter the following parameters:
1122		i. Stage 1 Hold
1123		1. 2.63 ramp rate
1124		2. 55°C
1125		3. 5 minutes
1126		ii. Stage 2 Hold
1127		1. 2.63 ramp rate
1128		2. 60°C
1129		3. 5 minutes
1130		iii. Stage 3 Hold
1131		1. 2.63 ramp rate
1132		2. 65°C
1133		3. 5 minutes
1134		iv. Stage 4 PCR
1135		1. Step 1:
1136		a. 2.63 ramp rate
1137		b. 92°C
1138		c. 5 seconds
1139		2. Step 2:
1140		a. 2.32 ramp rate
1141		b. 57°C
1142		c. 40 seconds
1143		d. Click on the camera icon under Step 2. A window will pop
1144		up asking for confirmation to turn off data collection during
1145		this step. Click "Ok".
1146		v. Located at the bottom of Stage 4 PCR change the number of cycles to 10
1147		vi. Stage 5 PCR
1148		1. Step 1:

1149		a. 2.63 ramp rate
1150		b. 92°C
1151		c. 5 seconds
1152		2. Step 2:
1153		a. 2.32 ramp rate
1154		b. 57°C
1155		c. 40 seconds
1156		d. Ensure the camera icon image is bold/on for data collection
1157		during the 30 cycles of Stage 5, Step 2.
1158	vii. Lo	cated at the bottom of Stage 4 PCR change the number of cycles to 30
1159	h. Scroll up a	nd choose the "Plate Setup" tab near the top of the screen.
1160	6. Plate Setup	
1161	a. Change th	e Passive Reference to "NONE"
1162	b. On the lov	ver right side of screen, ensure the Targets Tab is chosen then highlight
1163	and press	the addition button to add "Target 1". Press again to add "Target 2"
1164	c. Click on th	e "Target 1" box and change the name to CoV-2.
1165	d. Click the a	ssociated reporter box below the Reporter tab and, from the drop down
1166	menu, cho	ose FAM.
1167	e. Click on th	e "Target 2" box and change the name to PRC.
1168	f. Click the a	ssociated reporter box below the Reporter tab and, from the drop down
1169	menu, cho	ose CY5.
1170	g. Highlight	the "Actions" button located in the upper right side of the screen and
1171	press the	drop down button. In the drop down menu choose "Analysis Setting"
1172	h. Under An	alysis Setting, disable the following for all targets:
1173	i. U	se Default Column
1174	ii. A	uto Threshold Column
1175	iii. A	uto Baseline Column
1176	iv. Th	ne Baseline Start and Baseline End should default to 3 and 15
1177	i. Under "Th	reshold" click on the box associate with the CoV target and enter 70000.
1178		reshold" click on the box associated with the PRC target and enter 20000
1179	k. Click "Sav	2"
1180	l. Navigate	back to the "Actions" button and press the drop down button, choosing
1181	"Save As"	This will save your template to a location of choice. Save the template as
1182		S Cov-2 Assay".
1102	Cupating a Therma Fish	v OventStudio 7 Dve Test Dvessdu
1183	creating a inermofishe	r QuantStudio 7 Pro Test Procedure
1184	Note: These instructions	are based upon the user not having the QuantStudio 7 Real-Time PCR
1185		•
1186		esign and Analysis 2.2 software connected. The user must open the Lyra
1187	SARS CoV-2 template crea	ted previously with the software and save any newly created sample run

template onto a USB and transfer the template to the instrument.

1189	For connectivity related to the software and the instrument please contact your Thermo Fisher/ABI		
1190	luantStudio representative.		
1101			
1191			
1192	1) Open the Lyra SARS CoV-2 Assay Template previously generated.		
1193	2) Click on the Plate Setup Tab located near the top of the screen.		
1194	3) On the right side of the screen ensure the "Samples" tab is highlighted and press the	ne	
1195	addition button to add the number of samples being tested.		
1196	4) Click on the "Sample 1" box to rename the sample. Repeat this step for all subsequences.	uent	
1197	samples being entered.		
1198	5) Click the well located in the plate map then check the box next to the sample name	e from the	
1199	right side bar to associate the name to the well.		
1200	a. User also has the option to highlight the well location in the plate map and	click on	
1201	the "Enter sample" box. Enter the sample ID and press tab to continue to	the next	
1202	well in the plate map. This will automatically load the sample name into the	ıe	
1203	sidebar.		
1204	6) Once samples names have been entered, the wells may be highlighted by left clicking	ng the	
1205	mouse over starting well and dragging the mouse across all wells associated in run	. The	
1206	targets are then chosen by clicking the check boxes next to each target in the side	oar.	
1207	7) Click on the Actions button located top right of the screen and choose "Save As" in	the	
1208	dropdown menu.		
1209	 A pop-up window will appear directing the user to title the file according to 	כ	
1210	information pertaining to the sample run and the location of the file to be	saved.	
1211	b. Save the newly named (.edt) run file to a USB that is inserted into the com	puter.	
1212	8) Transfer the USB to the port on the front of the instrument.		
1213	9) From the options on the instrument's screen press "Load plate file". The QuantStu	dio 7 is a	
1214	touchscreen device.		
1215	10) From the "Run Queue" screen, press "USB drive" on the right side. This will bring u	ıp any	
1216	plate files saved on the USB.		
1217	11) Press the plate file associated with the run to be performed.		
1218	12) A new window will appear requesting location of results once the run is complete.		
1219	a. Press the "USB drive Connected" if the icon is not already highlighted and p	oress	
1220	"Done".		
1221	13) Centrifuge the 96-well sample plate to ensure all liquid is toward the bottom of each	ch well.	
1222	 Ensure the centrifuge is properly balanced. 		
1223	b. Gently pull the plate from the centrifuge to ensure all liquids remain at the	bottom	
1224	of the wells.		
1225	14) Press the double-arrow icon located at the top right sided corner of the screen on	the	
1226	instrument.		
1227	a. The instrument drawer will open from the front.		
1228	15) Place the centrifuged plate into the plate holder ensuring proper orientation of the	e plate.	

a. A1 well should be in the position of the top left corner

Quidel Corporation

Lyra® SARS-CoV-2 Assay 4/06/2020 Page 42 of 44

1230	b. The plate will appear slightly suspended above the block due to two silicone strips
1231	above and below this plate. This is to be expected and the instrument lid will press
1232	the plate down once the drawer has closed.
1233	16) Press "Start Run" on the screen of the instrument.
1234	a. A pop-up window will appear asking the user to confirm the plate has been loaded.
1235	b. If the plate has been loaded, press "Start Run" again or press "Open Drawer" to
1236	place the plate into the block and then press "Start Run"
1237	



M120 – Lyra SARS-CoV-2 Assay kit



Quidel Corporation 2005 East State Street, Suite 100 Athens, OH 45701 USA quidel.com

GLOSSARY

