

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) real-time RT-PCR E gene 2020

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ABSTRACT

A real-time RT-PCR designed to amplify a portion of the envelope gene of sequences from the Betacoronavirus sub-genus Sarbecovirus.

The probe and primers were published by Corman *et al.*, and we have slightly modified the protocol, increasing the concentration of the reverse primer, using a different kit and different cycling conditions.

This test has identified clinical positive cases of coronavirus disease 2019 (COVID-19).

THIS PROTOCOL ACCOMPANIES THE FOLLOWING PUBLICATION

Corman VM, Landt O, Kaiser M, Molenkamp R, Meijer A, Chu DK, et al. Detection of 2019 novel coronavirus (2019-nCoV) by real-time RT-PCR. Eurosurveillance. 2020;25(3):2000045. https://www.eurosurveillance.org/content/10.2807/1560-7917.ES.2020.25.3.2000045

GUIDELINES

- If using a different brand or model of real-time thermocycler, check the concentration of ROX is adequate.
- Method assumes the user is familiar with the thermocycler and software used to run the protocol.

MATERIALS

CATALOG # V	VENDOR V
1732088	Life Technologies
CATALOG #	vendor ~
1732088	Life Technologies
	1732088 ATALOG #

1

Oligo name	Sequence 5'-3'	Location *
E_Sarbeco_F1	ACAGGTACGTTAATAGTTAATAGCGT	26269- 26294
E_Sarbeco_R2	ATATTGCAGCAGTACGCACACA	26381- 26357
E_Sarbeco_P1	6FAM-ACACTAGCCATCCTTACTGCGCTTCG-BHQ1	26332- 26357

^{*}Based on numbering for GenBank accession NC_045512 Wuhan seafood market pneumonia virus isolate Wuhan-Hu-1

Reagents

2



Synthetic controls

3 Synthetic controls are produced using the <u>binary synthetic template oligonucleotide positive control for in-house diagnostic real-time RT-PCR method.</u>

The oligonucleotide sequences required to make controls for this assay are:

Probe control:

AAAATAATACGACTCACTATAGGGTGAAGAGAATCCACAAGGAATTGAAACACTAGCCATCCTTACTGCGCTTCGACAGTGTTCAGCAGGTCCTGTTGAAAA

Primer control:

Reaction Set-up

- Assay has been designed to be used on both a Rotor-Gene 6000 / Rotor-Gene Q 5-plex using 100-place rotor discs and a ABI 7500 Fast real-time machine.
 - Total reaction volume is 20µL.
 - Prepare sufficient for number of reaction plus a 'dead volume' usually 2 extra. Adjust as necessary if using a robotic dispenser.

Reagent	Volume (µI) x1	Final reaction concentration
Nuclease-Free water	4.39	
E_Sarbeco_F1	0.04	400nM
E_Sarbeco_R2	0.09	900nM
E_Sarbeco_P1	0.04	200nM
2X Reaction mix*	10	
Superscript III/Platinum Taq enzyme mix*	0.4	
ROX reference dye (25uM)*	0.04	50nM
TOTAL VOLUME	15	

^{*}Superscript®III Platinum® One-Step qRT-PCR kit

Dispense $15\mu l$ to each reaction well.

Add 5µl of template, extracted RNA, controls or NTC (nuclease-free water).

Total reaction volume is 20µl.

Amplification

5 PCR Amplification

1 cycle	40 cycles
50°C 5 minutes	95°C 3 seconds
95°C 2 minutes	60°C 30 seconds*

^{*}Florescence acquisition step

Result analysis

- 6 The definition used for a satisfactory positive result from a real-time fluorogenic PCR should include each of the following:
 - 1. A sigmoidal curve the trace travels horizontally, curves upward, continues in an exponential rise and followed by a curve towards a horizontal plateau phase
 - 2. A suitable level of fluorescence intensity as measured in comparison to a positive control (y-axis)
 - 3. A defined threshold cycle (C_T) value which the fluorescent curve has clearly exceeded (Fig.1 arrow) and which sits early in the log-linear phase and is <40 cycles
 - 4. A flat or non-sigmoidal curve or a curve that crosses the threshold with a C_T value >40 cycles is considered a negative result
 - 5. NTCs should not produce a curve.

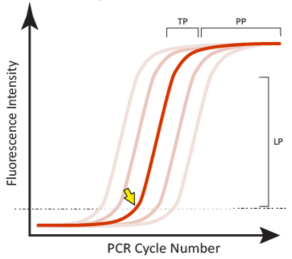


Figure 1. Examples of satisfactory sigmoidal amplification curve shape when considering an assay's fluorescent signal output. The crossing point or threshold cycle (C_T) is indicated (yellow arrow); it is the value at which fluorescence levels surpass a predefined (usually set during validation, or arbitrary) threshold level as shown in this normalized linear scale depiction. LP-log-linear phase of signal generated during the exponential part of the PCR amplification; TP-a slowing of the amplification and accompanying fluorescence signal marks the transition phase; PP-the plateau phase is reached when there is little or no increase in fluorescent signal despite continued cycling.

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