



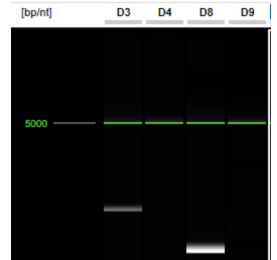
Feb 24, 2020

Version 1

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) RdRp nested RT-PCR V.1

DOI

<https://dx.doi.org/10.17504/protocols.io.bcmpiu5n>



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Coronavirus Method De...



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Hu D, Zhu C, Wang Y, Ai L, Yang L, Ye F, et al. Virome analysis for identification of novel mammalian viruses in bats from Southeast China. Scientific Reports. 2017;7(1):10917 . <https://www.nature.com/articles/s41598-017-11384-w>

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Protocol status: Working

We use this protocol and it's working

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Protocol Integer ID: 33167

Keywords: severe acute respiratory syndrome coronavirus, coronavirus rt, coronavirus, specific to sar, sar, pcr, nested rt, rdrp region, rdrp

Abstract

A nested RT-PCR targeting the RdRp region of the sub-genus *Sarbecovirus*. The primers are modified from the pan-coronavirus RT-PCR published by Hu *et al.* 2017 to be more specific to SARS-CoV-2.

Assay may be used in resource poor settings where real-time cyclers are not available.

Sanger sequencing can be used to confirm SARS-CoV-2 where WGS is not available or where WGS fails due to poor quality sample.

Materials

STEP MATERIALS

☒ SuperScript III One-Step RT-PCR System with Platinum Taq **Invitrogen - Thermo Fisher Catalog #12574026**

☒ MyFi Mix **Bioline Catalog #BIO-25049**

Protocol materials

☒ SuperScript III One-Step RT-PCR System with Platinum Taq **Invitrogen - Thermo Fisher Catalog #12574026**

☒ MyFi Mix **Bioline Catalog #BIO-25049**



Oligonucleotides

1

	Name	Purpose	5'-3'
	Wuhan15283-F	Round 1	ATG GGT TGG GAT TAT CCT AAA TG
	Wuhan15887-R	Round 1	TGT TGA GAG CAA AAT TCA TG
	Wuhan15286-F	Round 2	GGT TGG GAT TAT CCT AAA TGT GA
	Wuhan15725-R	Round 2	GCA TCG TCA GAG AGT ATC ATC AT

Round 1 amplified products will produce a band of 605bp.

Round 2 amplified products will produce a band of 440bp.

Mix

2

 SuperScript III One-Step RT-PCR System with Platinum Taq Invitrogen - Thermo
Fisher Catalog #12574026

 MyFi Mix Bioline Catalog #BIO-25049

Round 1 mix

	Reagent	Volume x1 (μ l)	Final Concentration
	Nuclease-free water	3.2	
	2 x Reaction mix*	10	1 X
	Wuhan15283-F (20p mol/ μ l)	0.5	500 nM
	Wuhan15887-R (20p mol/ μ l)	0.5	500 nM
	Superscript/Platinum Taq mix*	0.8	
	TOTAL	15	

*SuperScript III One-Step RT-PCR System with Platinum Taq #12574026

Dispense mix in 15 μ l amounts in 0.2ml PCR tubes suitable for thermocycler.

Mix maybe stored frozen if necessary.

Round 2 mix

	Reagent	Volume x1 (μl)	Final Concentration
	Nuclease-free water	4	
	MyFi 2X mix*	10	1X
	Wuhan15 286-F (20p mol/μl)	0.5	500 nM
	Wuhan15 725-R (20p mol/μl)	0.5	500 nM
	TOTAL	15	

*MyFi™ Mix Bioline #BIO-25049

Dispense mix in 15μl amounts in 0.2ml PCR tubes suitable for thermocycler. Mix may be stored frozen if necessary.

AMPLIFICATION ROUND 1

3

- Thaw required number of tubes with round 1 mix.
- Add 5μl of extract, control, or NTC (nuclease-free water) to round 1 mix above for a final reaction volume of 20μl.
- Label the tubes and record details.

The assay has been used with Eppendorf thermocyclers.

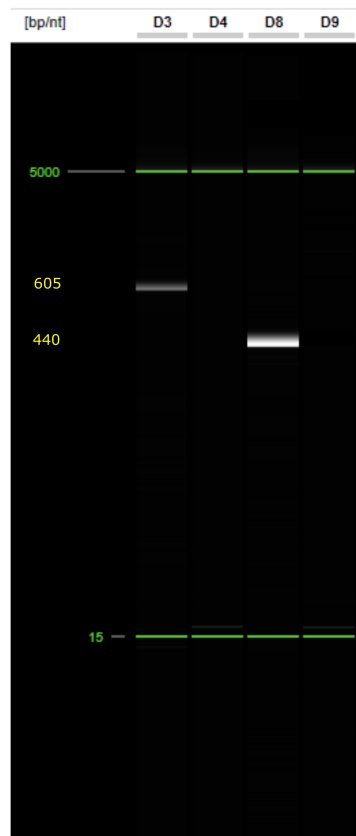
PCR cycling times

AMPLIFICATION ROUND 2

- 4
 - Dilute each of the round 1 amplicons 1/100 in nuclease-free water (2ul + 198ul is ideal)
 - Thaw required number of tubes with round 2 mix.
 - Add 5µl of the diluted round 1 amplicon.
 - Total reaction volume is 20µl.

GEL ELECTROPHORESIS

- 5 Amplified products are analysed by gel electrophoresis or equipment such as a QIAxcel or equivalent.
Round 1 amplified products will produce a band of 605bp.
Round 2 amplified products will produce a band of 440bp.
No template controls (NTC) should be not detected.



Example of result from a QIAxcel.
D3-Round 1 band, D8-round 2 band, D4 and D9 are NTC in round 1 and 2 respectively.

