Diagnostic detection of Wuhan coronavirus 2019 by real-time RT-PCR

-Protocol and preliminary evaluation as of Jan 13, 2020-

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Positive control material is available from Charité, Berlin, via EVAg (https://www.european-virus-archive.com/).

We acknowledge the originators of sequences in GISAID (www.gisaid.org): National Institute for Viral Disease Control and Prevention, China, Institute of Pathogen Biology, Chinese Academy of Medical Sciences, Peking Union Medical College, China, and Wuhan Jinyintan Hospital Wuhan Institute of Virology, Chinese Academy of Sciences, China). We acknowledge Professor Yong-Zhen Zhang, Shanghai Public Health Clinical Center & School of Public Health, Fudan University, Shanghai, China for release of another sequence (MN908947).

Abbreviations and taxonomy related to the Wuhan virus are not used in any systematic way, i.e., there are multiple different designations and abbreviations for the "Wuhan virus" in this document. They all relate to the same viral agent. We use the term "SARS-related Coronavirus" to include the SARS virus as well as the clade of betacoronaviruses known to be associated with (mainly) rhinolophid bats across the Palearctic. The latest taxonomy classifies these viruses in a subgenus termed Sarbecovirus.

Background

We used known SARS- and SARS-related coronaviruses (bat viruses from our own studies as well as literature sources) to generate a non-redundant alignment (excerpts shown in Annex). We designed candidate diagnostic RT-PCR assays before release of the first sequence of the Wuhan virus. Upon sequence release, three assays were selected based on their matching to the Wuhan virus as per inspection of the sequence alignment (Figures 1 and 2).

All assays can use SARS-CoV genomic RNA as positive control. Synthetic control RNA for Wuhan virus will be provided shortly.

First line screening assay: E gene assay Confirmatory assay: RdRp gene assay Additional confirmatory assay: N gene assay



Figure 1 relative positions of amplicon targets on SARS-CoV ad Wuhan-CoV genome. N: nucleocapsid; ORF: open reading frame; RdRp: RNA-dependent RNA polymerase. Numbers below amplicon are genome positions according to SARS-CoV, NC_004718.

Materials and assay formulation

Clinical samples and CoV cell culture supernatants

Respiratory samples were obtained during 2019 from patients hospitalized at Charité medical center and tested by the NxTAG® Respiratory Pathogen Panel (Luminex) or in cases of MERS-CoV by the MERS-CoV upE assay as published before (1).

Cell culture supernatants from typed coronaviruses were available at our research and clinical laboratories. The typed avian influenza virus RNA (H5N1) was obtained from the German Society for Promotion of Quality Assurance in Medical Laboratories (INSTAND) proficiency testing panels. RNA was extracted from clinical samples by using the MagNA Pure 96 system (Roche) and from cell culture supernatants by the viral RNA mini kit (Qiagen).

Assay design

For oligonucleotide design and in-silico evaluation we downloaded all complete and partial (if >400 nucleotides) SARS-related virus sequences available at GenBank by January 1st, 2020. The list (n=729 entries) was manually checked and artificial sequences (lab-derived,

synthetic etc.), as well as sequence duplicates removed, resulting in a final list of 375 sequences. These sequences were aligned and the alignment used for assay design. The alignment was later complemented by sequences released from the Wuhan cluster. All presently release sequences match the amplicons (Figure 2). An overview of oligonucleotide binding sites in all unique sequences of bat-associated SARS-related viruses is shown in the appendix.

		pSARS-	
RdRP gene	pSARS-F P1:	$\cdots \cdots \cdots W \cdots R \cdots \cdots M \cdots T \cdots \cdots$	pSARS-R
	P2		
WH-Human_1 China 2019-Dec	GTGAAATGGTCATGTGTGGCGG	CCAGGTGGAACCTCATCAGGAGATGC	TATGCTAATAGTGTTTTTAACATTTG
BetaCoV/Wuhan/IPBCAMS-WH-01/2019[EP] [SL 402123 BetaCoV/Wuhan/IVDC-HB-01/2019[EP] [SL 402119 BetaCoV/Wuhan/IVDC-HB-04/2020][EP] [SL 402120 BetaCoV/Wuhan/IVDC-HB-05/2019[EP] [SL 402120 BetaCoV/Wuhan/IVDC-HB-05/2019[EP] [SL 402121 BetaCoV/Wuhan/IVD/H2/019[EP] [SL 402124 Mg772333 Bat SARS-related CoV [GutSL-CoVZC45] NC_004718 Human SARS-related CoV [GutSL-CoVZC45] NC_014470 Bat SARS-related CoV (BM48-31/BGR/2008)	G G T	A C T A C T . C T . A T	e c
	E_Sarbeco_F	E_Sarbeco_P	E_Sarbeco_R
E gene			
WH-Human_1 China 2019-Dec	ACAGGTACGTTAATAGTTAATAGC	GT ACACTAGCCATCCTTACTGCGCT	TCGATTGTGTGCGTACTGCTGCAATAT
BetaCoV/Wuhan/IPBCAMS-WH-01/2019 EPI_ISL_402123			
BetaCoV/Wuhan/IVDC-HB-01/2019 EPI_ISL_402119 BetaCoV/Wuhan/IVDC-HB-04/2020 EPI_ISL_402120			
BetaCoV/Wuhan/IVDC-HB-05/2019 EPI_ISL_402121			
BetaCoV/Wuhan/WIV04/2019 EPI_ISL_402124			
NC_004718 Human SARS-related CoV (bat-SL-CoV2C45)			
NC_014470 Bat SARS-related CoV (BM48-31/BGR/2008)	· · C · · · · · · · · · · · · · · · · ·	· · · C · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·
	N_Sarbeco_F	N_Sarbeco_P	N_Sarbeco_R
N gene			
WH-Human_1 China 2019-Dec	CACATTGGCACCCGCAAT	C ACTTCCTCAAGGAACAACATTG	CCA CAAGCCTCTTCTCGTTCCTC
BetaCoV/Wuhan/IPBCAMS-WH-01/2019 EPI_ISL_402123			
BetaCoV/Wuhan/IVDC-HB-01/2019 EPI_ISL_402119 BetaCoV/Wuhan/IVDC-HB-04/2020 EPI_ISL_402120			
BetaCoV/Wuhan/IVDC-HB-05/2019[EPI_ISL_402121			
BetaCoV/Wuhan/WIV04/2019 EPI_ISL_402124			
NG 004718 Human SARS-related CoV (bat-SL-CoV2C45)			· · · · · · · · · · · · · · · · · · ·
NC_014470 Bat SARS-related CoV (BM48-31/BGR/2008)	· · · · · · · · · · · · · · · G · · · ·	GT · A · · A · · · · T · · T · · C · · ·	· · · · · · · · · · · · C · · G · · · ·

Figure 2 Partial alignments of oligonucleotide binding regions. Panels show six available sequences of the Wuhan-CoV, aligned to the corresponding partial sequences of SARS-CoV strain Frankfurt 1, which can be used as a positive control for all three RT-PCR assays. The alignment also contains the most closely-related bat virus (Bat SARS-related CoV isolate bat-SL-CoVZC45, GenBank Acc.No. MG772933.1) as well as the most distant member within the SARS-related bat CoV clade, detected in Bulgaria (GenBank Acc. No. NC_014470). Dots represent identical nucleotides compared to Wuhan-Hu 1. Substitutions are specified. More comprehensive alignments in the Appendix.

Real-time reverse-transcription polymerase chain reaction

All assays used the same conditions. Primer and probe sequences, as well as optimized concentrations are shown in Table 1. A 25- μ l reaction was set up containing 5 μ l of RNA, 12.5 μ l of 2 X reaction buffer provided with the Superscript III one step RT-PCR system with Platinum Taq Polymerase (Invitrogen; containing 0.4 mM of each deoxyribonucleotide triphosphates (dNTP) and 3.2 mM magnesium sulfate), 1 μ l of reverse transcriptase/Taq mixture from the kit, 0.4 μ l of a 50 mM magnesium sulfate solution (Invitrogen – not provided with the kit), and 1 μ g of nonacetylated bovine serum albumin (Roche). All oligonucleotides were synthesised and provided by Tib-Molbiol, Berlin. Thermal cycling was performed at 55°C for 10 min for reverse transcription, followed by 95°C for 3 min and then 45 cycles of 95°C for 15 s, 58°C for 30 s.

Table 1. Primers and probes

Optimized concentrations are mol per liter of final reaction mix.

(e.g., 1.5 microliters of a 10 micromolar (uM) primer stock solution per 25 microliter (ul) total reaction volume yields a final concentration of 600 nanomol per liter (nM) as indicated in the table)

-note that standard, non-optimized reaction conditions as indicated by suppliers of one-step RT-PCR kits will generally yield sufficient sensitivity-

Assay/ Use	Oligonucleotide ID	Sequence (5'–3')	Comment
RdRP gene	RdRP_SARSr-F2	GTGARATGGTCATGTGTGGCGG	use 600 nM per reaction
	RdRP_SARSr-R1	CARATGTTAAASACACTATTAGCATA	use 800 nM per reaction
	RdRP_SARSr-P2	FAM-CAGGTGGAACCTCATCAGGAGATGC- BBQ	Specific for Wuhan-CoV, will not detect SARS- CoV use 100 nM per reaction and mix with P1
	RdRP_SARSr-P1	FAM- CCAGGTGGWACRTCATCMGGTGATGC- BBQ	Pan Sarbeco-Probe, will detect Wuhan virus, SARS-CoV and bat-SARS-related CoVs use 100 nM per reaction and mix with P2
E gene	E_Sarbeco_F1	ACAGGTACGTTAATAGTTAATAGCGT	use 400 nM per reaction
	E_Sarbeco_R2	ATATTGCAGCAGTACGCACACA	use 400 nM per reaction
	E_Sarbeco_P1	FAM-ACACTAGCCATCCTTACTGCGCTTCG- BBQ	use 200 nM per reaction
N gene	N_Sarbeco_F1	CACATTGGCACCCGCAATC	use 600 nM per reaction
	N_Sarbeco_R1	GAGGAACGAGAAGAGGCTTG	use 800 nM per reaction
	N_Sarbeco_P1	FAM-ACTTCCTCAAGGAACAACATTGCCA- BBQ	use 200 nM per reaction

W is A/T; R is G/A; M is A/C ; FAM, 6-carboxyfluorescein; BBQ, blackberry quencher

Technical sensitivity testing

Preliminary assessment of analytical sensitivity for RdRp assay.

We tested purified cell culture supernatant containing SARS-CoV strain Frankfurt-1 virions grown on Vero cells, and quantified by real-time RT-PCR assay as described in Drosten et al. (2) using a specific *in-vitro* transcribed RNA quantification standard. The results are shown in Figure 3. All assays are highly sensitive.



Figure 3A First line assay: E gene

Technical limit of detection (LOD) = 5.2 RNA copies/reaction, at 95% hit rate; 95% CI: 3.7-9.6 RNA copies/reaction.



Legend to these figures: X-axis shows input RNA copies per reaction. Y-axis shows positive results in all parallel reactions performed, squares are experimental data points resulting from replicate testing of given concentrations (x-axis) in parallels assays (8 replicate reactions per datum point). The inner line is a probit curve (dose-response rule). The outer dotted lines are 95% confidence intervals.

Figure 3B. Confirmatory assay: RdRP gene Technical LOD = 3.8 RNA copies/reaction, at 95% hit rate; 95% CI: 2.7-7.6 RNA copies/reaction.



Figure 3C: Second confirmatory assay: N gene Technical LOD = 8.3 RNA copies/reaction, at 95% hit rate; 95% CI: 6.1-16.3 RNA copies/reaction.

Breadth of detection

To show that the assays will detect other bat-associated SARS-related viruses, we tested bat-derived fecal samples available from Drexler et al., (3) und Muth et al., (4) using the novel assays.

KC633203, Betacoronavirus BtCoV/Rhi_eur/BB98-98/BGR/2008 KC633204, Betacoronavirus BtCoV/Rhi_eur/BB98-92/BGR/2008 KC633201, Betacoronavirus BtCoV/Rhi_bla/BB98-22/BGR/2008 GU190221 Betacoronavirus Bat coronavirus BR98-19/BGR/2008 GU190222 Betacoronavirus Bat coronavirus BM98-01/BGR/2008 GU190223, Betacoronavirus Bat coronavirus BM98-13/BGR/2008

All samples were successfully tested positive by the E gene assay. Detection of these relatively distant members of the SARS-related CoV clade suggests that all Asian viruses are likely to be detected.

Specificity testing

1. Chemical stability

To exclude non-specific reactivity of oligonucleotides among each other, all assays were tested 40 times in parallel with water and no other nucleic acid except the provided oligonucleotides. In none of these reactions was any positive signal detected.

2. Cross-reactivity with other coronaviruses

Cell culture supernatants containing human coronaviruses (HCoV)-229E, -NL63, -OC43, and -HKU1 as well as MERS-CoV were tested in all three assays (Table 2). For the non-cultivable HCoV-HKU1, supernatant from human airway culture was used. Virus RNA concentration in all samples was determined by specific real-time RT-PCRs and in-vitro transcribed RNA standards designed for absolute viral load quantification.

Cell culture supernatants	Tested concentration	Result
Alphacoronaviruses		
Human coronavirus NL63	4x10^9 RNA copies/ml	No reactivity with any of three assays
Human coronavirus 229E	3x10^9 RNA copies/ml	No reactivity with any of three assays
Betacoronaviruses		
Betacoronavirus 1 (strain HCoV-OC43)	1x10^10 RNA copies/ml	No reactivity with any of three assays
Human coronavirus HKU1 (HCOV-HKU1)	1x10^5 RNA copies /ml	No reactivity with any of three assays
Middle East respiratory syndrome-related coronavirus (strain EMC/2012)	1x10^8 RNA copies/ml	No reactivity with any of three assays

Table 2. Cell-culture supernatants tested by all assays

3. Tests of human clinical samples previously tested to contain respiratory viruses

All assays were applied on human clinical samples from our own diagnostic services, previously tested positive for the viruses listed in Table 3. All tests returned negative results.

Clinical samples with known viruses	Number of samples tested in all three assays
HCoV-HKU1	2
HCoV-OC43	5
HCoV-NL63	5
HCoV-229E	5
MERS-CoV	5
Influenza A (H1N1/09)	6
Influenza A (H3N2)	5
Influenza A(H5N1)	1
Influenza B	3
Rhinovirus/Enterovirus	3
Respiratory syncytial virus (A/B)	6
Parainfluenza 1 virus	3
Parainfluenza 2 virus	3
Parainfluenza 3 virus	3
Parainfluenza A or -B virus	5
Human metapneumovirus	3
Adenovirus	3
Human Bocavirus	3
Legionella spp.	3
Mycoplasma spp.	3
Total clinical samples	75

Table 3. Tests of known respiratory viruses and bacteria in clinical samples

References

1. Corman VM, Eckerle I, Bleicker T, Zaki A, Landt O, Eschbach-Bludau M, et al. Detection of a novel human coronavirus by real-time reverse-transcription polymerase chain reaction. Euro Surveill. 2012;17(39).

2. Drosten C, Gunther S, Preiser W, van der Werf S, Brodt HR, Becker S, et al. Identification of a novel coronavirus in patients with severe acute respiratory syndrome. N Engl J Med. 2003;348(20):1967-76.

3. Drexler JF, Gloza-Rausch F, Glende J, Corman VM, Muth D, Goettsche M, et al. Genomic characterization of severe acute respiratory syndrome-related coronavirus in European bats and classification of coronaviruses based on partial RNA-dependent RNA polymerase gene sequences. J Virol. 2010;84(21):11336-49.

4. Muth D, Corman VM, Roth H, Binger T, Dijkman R, Gottula LT, et al. Attenuation of replication by a 29 nucleotide deletion in SARS-coronavirus acquired during the early stages of human-to-human transmission. Sci Rep. 2018;8(1):15177.

Annex:

RdRP_SARSr-P2	RHBP SARS-P2
RdRP_SARSr_Oligos	GTGARATGGTCATGTGTGGCGGCCAGGTGGWACRTCATCMGGTGATGCTATGCTAATAGTGTSTTTAACATYTG RdB SABS-62
RdRP_SAR5r_Oligos WH-Human,1 [China] 2019-Dec BetaCoVWuhan/WECAMS-WH-01/2019 [EPI_ISL_402123 BetaCoVWuhan/WECAMS-WH-01/2019 [EPI_ISL_402119 BetaCoVWuhan/WDC-HB-01/2020 [EPI_ISL_402119 BetaCoVWuhan/WDC-HB-01/2020 [EPI_ISL_402121 BetaCoVWuhan/WDC-HB-01/2020 [EPI_ISL_402124 NC_004718 [SAR5 coronavirus (RdBg gene for RNA dependent RNA polymerase, p BA889995 (SAR5 bat coronavirus RdBg gene for RNA dependent RNA polymerase, p DQ112031 [Bat-SAR5-cineated bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254452 (SAR5-related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254453 [SAR5-related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254453 (SAR5-related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254453 [SAR5-related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254453 [SAR5-Related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254453 [SAR5-related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF254453 [SAR5-related coronavirus isolate Aniong:11 ortTab polyprotein and KF25455 [SAR5-related bat coronavirus isolate Aniong:11 ortTab polyprotein and KF25453 [SAR5-Related SAR5-Related SAR5-Relat	STGARATGGTCATEGTGGGGGCCAGGTGGWACRTCATCATCMGGTGATGCTATGCTAATAGTGTSTTTAACATYTG RdP_SAS5F2 RdP_SAS5F2
KC633020 (Betacoronavirus BCOV/Rh. urv/RB99-04/BGR/2009 RNA-dependent RNA KC633020 (Betacoronavirus BCOV/Rh. urv/RB99-04/BGR/2008 RNA-dependent RNA KC633021 (Betacoronavirus BCOV/Rh. jrv/R15/174/2009 RNA-dependent RNA p.n. KC633231 (Betacoronavirus BCOV/Rh. jrv/R15-03/SPA/2010 RNA-dependent RNA p.n. KC633232 (Betacoronavirus BCOV/Rh. jr)/R156-23/SPA/2010 RNA-dependent RNA p.n.	Ατ.Α
E_Sarbeco_assay	ACAGGTACGTTAATAGTTAATAGCGTACACTAGCCATCCTTACTGCGCTTCGTGTGTGCGTACTGCTGCAATAT
WH-Human, 11 Chinal 2019-Dec. Beat-CoWWham/IPG-AKS-WH-01/2019 [EPLISL, 402123 Beat-CoWWham/IPG-AKS-WH-01/2019 [EPLISL, 402129 Beat-CoWWham/IVOC-HB-01/2019 [EPLISL, 402121 BetaCoWWham/IVOC-HB-05/2019 [EPLISL, 402124 BetaCoWWham/IVOC-HB-05/2019 [EPLISL, 402124 NC_004718 [SARS coronavirus, SIRS-12, complete genome] A1593095 [SARS coronavirus] SaR21-12, complete genome] A1593095 [SARS coronavirus] SaR21-12, complete genome] B83396 [SARS coronavirus] SaR21-12, complete genome] [B83969 [SARS coronavirus] SAR21-12, complete genome] [B83969 [SARS coronavirus] NLIS-18 [SaR24, complete genome] K7352407 [Severe acute respiratory syndrome-related coronavirus strain BtKY72, compl NC_014470 [Bat coronavirus BM48-31/BGR/2008, complete genome]	C
N_Sarbeo_Oligos	CACATTGGCACCCGCAATC ACTTCCTCAAGGAACAACATTGCCA CAAGCCTCTTCTCGTTCCTC N.Sarbeco. FI N.Sarbeco. P1 N.Sarbeco. R1
WH-Human_11China]2019-Dec BetaCoVWuhan/WDCAK-9H-01/2019 [EPL/SL_402123 BetaCoVWuhan/WDCAK-9H-01/2019 [EPL/SL_402119 BetaCoVWuhan/WDC-HB-01/2019 [EPL/SL_402121 BetaCoVWuhan/WDK-2019 [EPL/SL_402124 C_007718 [KASK cornoavirus, complete genome) DQ412042 [KaSK scornoavirus, complete genome) DQ412042 [KaSK scornoavirus B1]82-12, complete genome) (XF02087 (SASK scornoavirus B1]82-12, complete genome) (XF02087 (SASK scornoavirus B1]82-12, complete genome) (XF32087 [KaSK scornoavirus SASK scornoavirus scolate longuan-140 orf1ab polyprotein, st (XF32814 [KRR-BetaCoV/VN2013, complete genome) (KF32816 [KRR-BetaCoV/VN2013, complete genome) (KF32816 [KRR-BetaCoV/VN2013, scornglete genome) (KF32816 [KRR-BetaCoV/VN2013, scornglete genome) (KF3816 [KRR-BetaCoV/VN2013, scornglete genome) (KF3816 [KRR-BetaCoV/VN2013, scornglete genome) (KF478816 [KRR-BASK-like cornoavirus Isolate Longuan-140 orf1ab polyprotein, st (KF478816 [KRR-BetaCoV/VN2013, scornglete genome) (KF47816 [KRR-BataCoV/VN2013, scornglete genome) (KF477816 [KRR-BASK-like cornoavirus Isolate Rs4081, complete genome) (K417144 [KRI SASK-like cornoavirus Isolate Rs4084, complete genome) (K417146 [KRI SASK-like cornoavirus Isolate Rs4084, complete genome) (K417146 [KRI SASK-like cornoavirus Isolate Rs4084, complete genome) (K417146 [KRI SASK-like cornoavirus Isolate Rs4084, complete genome)	C

Annex figure. Non-redundant alignments of SARS-related CoVs focused on oligonucleotide binding sites of all assays (top to bottom: RdRp, E, N). Viruses not present in these alignments have been removed because their binding sites are 100% identical to one of the members of the alignment. ("--") means sequence gaps not covered by oligonucleotides. Note that these alignments contain only one sequence of the Wuhan virus while Figure 2 above contains all presently released sequences. We will fuse this into one figure.

Annex 2: Bench Protocol

Real-time rtPCR for Betacoronavirus (Wuhan Betacoronavirus, Wu-Hu-1)

Example formulation:

Thermo Fischer /Invitrogen SuperScriptIII OneStep RT-PCR System with Platinum Taq DNA Polymerase

<u>E assay:</u>

	<u>25µl</u>	<u>Cycler:</u>	
MasterMix:	single rxn, μl		
H ₂ O (RNAse free)	2.6	55°C 10'	
2x Reaction mix*	12.5		
MgSO ₄ (50mM)	0.4	94°C 3'	
BSA (1 mg/ml)**	1	94°C 15"	
Fwd primer (10 µM)	1	58°C 30" 45x	
Rev primer (10 µM)	1		
Probe (10 µM)	0.5	40°C 30"	
SSIII/Taq EnzymeMix*	1		
	20	' = minutes; " = seconds	s
Template RNA	5		

RdRp- and N assay:

	<u>25µl</u>	Cycler:	
MasterMix:	single rxn, µl		
H ₂ O (RNAse free)	1.1	55°C 10'	
2x Reaction mix*	12.5		
MgSO ₄ (50mM)	0.4	94°C 3'	
BSA (1 mg/ml)**	1	94°C 15"	
Fwd primer (10 µM)	1.5	58°C 30" 4	5x
Rev primer (10 µM)	2	·	
Probe (10 µM)	0.5	40°C 30"	
SSIII/Taq EnzymeMix*	1		
	20	' = minutes; " = sec	conds
Template RNA	5		

* Thermo Fischer/Invitrogen: SuperScriptIII OneStep RT-PCR System with Platinum® Taq DNA Polymerase

** MgSO4 (50 mM) [Sigma], This component is not provided with the OneStep RT-PCR kit *** non-acetylated [Roche]. This component is only necessary when using glass capillaries with LightCycler. Can be replaced with water in plastic vessel machines such as ABI 7500, LC 480, etc.

Primers / probe: See table

<u>Positive Control:</u> SARS-CoV (e.g. strain Frankfurt 1) **<u>References:</u>** Corman/Drosten, unpublished

Assay/ Use	Oligonucleotide ID	Sequence (5'–3')	Comment
RdRP gene	RdRP_SARSr-F2	GTGARATGGTCATGTGTGGCGG	use 600 nM per reaction
	RdRP_SARSr-R1	CARATGTTAAASACACTATTAGCATA	use 800 nM per reaction
	RdRP_SARSr-P2	FAM-CAGGTGGAACCTCATCAGGAGATGC- BBQ	Specific for Wuhan-CoV, will not detect SARS-CoV use 100 nM per reaction and mix with P1
	RdRP_SARSr-P1	FAM- CCAGGTGGWACRTCATCMGGTGATGC- BBQ	Pan Sarbeco-Probe, will detect Wuhan virus, SARS-CoV and bat-SARS-related CoVs use 100 nM per reaction and mix with P2
E gene	E_Sarbeco_F1	ACAGGTACGTTAATAGTTAATAGCGT	use 400 nM per reaction
	E_Sarbeco_R2	ATATTGCAGCAGTACGCACACA	use 400 nM per reaction
	E_Sarbeco_P1	FAM-ACACTAGCCATCCTTACTGCGCTTCG- BBQ	use 200 nM per reaction
N gene	N_Sarbeco_F1	CACATTGGCACCCGCAATC	use 600 nM per reaction
	N_Sarbeco_R1	GAGGAACGAGAAGAGGCTTG	use 800 nM per reaction
	N_Sarbeco_P1	FAM-ACTTCCTCAAGGAACAACATTGCCA- BBQ	use 200 nM per reaction