

Evaluation of Different PCR Assay Formats for
Sensitive and Specific Detection of SARS-CoV-2 RNA

Jeremy Ratcliff¹, Dung Nyugen¹, Monique Andersson², Peter Simmonds^{1*}

¹*Nuffield Department of Medicine, Peter Medawar Building for Pathogen Research, University of Oxford, Oxford OX1 3SY, UK;* ²*Department of Infectious Diseases and Microbiology, Oxford University Hospitals NHS Foundation Trust, John Radcliffe Hospital, Oxford OX3 9DU, UK*

Running Title: *PCR assays for detecting SARS-CoV-2*

*To whom correspondence should be addressed.

Tel. +44 1865 281 233

Email: Peter.simmonds@ndm.ox.ac.uk

ABSTRACT

Accurate identification of individuals infected with SARS-CoV-2 is crucial for efforts to control the ongoing COVID-19 pandemic. Polymerase chain reaction (PCR)-based assays are the gold standard for detecting viral RNA in patient samples and are used extensively in clinical settings. Most currently used quantitative PCR (RT-qPCRs) rely upon real-time detection of PCR product using specialized laboratory equipment. To enable the application of PCR in resource-poor or non-specialist laboratories, we have developed and evaluated a nested PCR method for SARS-CoV-2 RNA using simple agarose gel electrophoresis for product detection. Using clinical samples tested by conventional qPCR methods and RNA transcripts of defined RNA copy number, the nested PCR based on the RdRP gene demonstrated high sensitivity and specificity for SARS-CoV-2 RNA detection in clinical samples, but showed variable and transcript length-dependent sensitivity for RNA transcripts. Samples and transcripts were further evaluated in an additional N protein real-time quantitative PCR assay. As determined by 50% endpoint detection, the sensitivities of three RT-qPCRs and nested PCR methods varied substantially depending on the transcript target with no method approaching single copy detection. Overall, these findings highlight the need for assay validation and optimization and demonstrate the inability to precisely compare viral quantification from different PCR methodologies without calibration.

Key words: SARS-CoV-2, Coronavirus, Polymerase Chain Reaction, Diagnostics, qPCR

Abstract Word Count: 204

Word Count: 2533

1

INTRODUCTION

2

3 SARS-Coronavirus-2 (SARS-CoV-2), a human-infective member of the *Betacoronavirus* genus
4 (family *Coronaviridae*), was first identified in the Hubei Province of China in late 2019 as the
5 causative agent behind an increased number of cases of respiratory illness occasionally leading
6 to acute respiratory distress and death.¹⁻³ The outbreak was declared a Public Health Emergency
7 of International Concern by the World Health Organization on January 30th, 2020 and the
8 associated disease was named COVID-19 on February 11th, 2020.^{4,5} The disease has since
9 spread globally and by June 23rd has infected nearly 9 million individuals in over 180 countries,
10 causing at least 465,000 deaths.^{6,7}

11 The ability to accurately identify and diagnose asymptotically and symptomatically infected
12 patients is crucial for efforts aimed at limiting person-to-person transmission and controlling the
13 outbreak.⁸⁻¹⁰ The standard method for diagnosing viral infections is through the detection of viral
14 nucleic acid in clinical samples. Reverse-transcriptase quantitative polymerase chain reaction
15 (RT-qPCR) is the gold standard used in most diagnostic laboratories.¹¹ Probe-based RT-qPCR
16 relies on the binding and amplification of three oligonucleotides (two primers and one internal
17 fluorescent probe) and the accumulation of fluorescence signal mediated by DNA polymerase
18 activity. RT-qPCR is not widely accessible as the method relies upon the use of expensive real-
19 time PCR platforms and the probe component of the assay is typically the most costly reagent.
20 An alternative, more cost-effective diagnostic method for SARS-CoV-2 RNA is nested PCR.
21 Nested PCR is based on the use of two sequential PCR amplifications wherein the secondary set
22 of primers target sequences nested within the amplicon produced by the first round amplification.
23 Compared to conventional PCR, which uses a single round of replication, nested PCR has
24 increased sensitivity and decreased risk of amplification of non-specific products.¹² Nested PCR
25 methods were developed for SARS,¹³ but no nested PCR method for SARS-CoV-2 has yet been
26 published.

27 In this study, a nested PCR assay for SARS-CoV-2 has been developed and its performance for
28 an *in vitro* transcribed partial transcript of the RNA-Dependent RNA Polymerase (RdRP) gene
29 and three full-length transcripts was compared against a corrected version of the RdRP-directed
30 oligonucleotides from the Charité Institute of Virology (Charité)¹⁴ and the United States Centers
31 for Disease Control and Prevention (CDC) N1 primer/probe set.¹⁵ Unexpectedly, the
32 performances varied substantially depending on the detection method and target assayed,

33 underpinning the need for in-house validation and optimization. The result also challenges the
34 notion that Ct values presented without context could be an informative metric for the progression
35 of disease and can be compared across different amplification techniques and laboratories.

36

37

MATERIALS AND METHODS

38

39 **Primer design**

40 1st round nested PCR primers (nF1, nR1) were designed to anneal to conserved regions flanking
41 a 1.1kb region of the RdRP gene of SARS-CoV-2. 2nd round nested primers (nF2, nR2) were
42 designed to anneal to conserved regions on the amplicon product of nF1 and nR1 amplification.
43 Both sets of nested PCR primers were acquired from Integrated DNA Technologies. The Charité
44 RT-PCR was based upon previously described primer/probes for the RdRP gene¹⁴ but with
45 modifications to the antisense primer to ensure complete sequence complementarity with SARS-
46 CoV-2 sequences (Table 1; differences underlined). All primers and probes for the Charité and
47 CDC N1 PCRs were obtained from ATDBio. All primer sequences and working concentration are
48 available in Table 1.

49 **Clinical Samples**

50 Residual RNA from clinical samples was eluted in buffer AVE (Qiagen) following extraction using
51 a Qiasymphony DSP virus/pathogen minikit. SARS-CoV-2 infection status was determined by
52 RT-qPCR operated on the Rotor-Gene Q using an RdRP gene target validated by Public Health
53 England (PHE)¹⁴ or the Altona RealStar SARS-CoV-2 RT-PCR Kit targeting the E and S genes.

54 **Nested PCR**

55 The nested PCR uses two sequential PCR amplifications for highly specific and sensitive
56 amplification of target sequences (primer sequences and concentrations listed in Table 1). For
57 the first round of amplification, a 25 µL reaction mix containing 5 µL of RNA extract, 12.5 µL of 2X
58 Quantitect Probe RT-PCR Master Mix (Qiagen), 0.5 µL of Superscript III Reverse Transcriptase
59 (ThermoFisher Scientific), 5 µL of 5X 1st-round primer mix, and 2 µL of PCR-grade water was
60 amplified on a thermal cycler using the following settings: reverse transcription at 50°C for 30 m,
61 activation at 95°C for 15 m, 40 cycles of 95°C for 15 s, 55°C for 30 s, and 68°C for 1 m, and a
62 final extension of 68°C for 5 m. For the second round amplification, a 25 µL reaction mix containing

63 1 μL of the first round product, 5 μL of 5X GoTaq Green Master Mix (Promega), 0.125 μL of 5u/ μL
64 GoTaq G2 polymerase (Promega), 2 μL of 2.5 mM dNTP mix (Strattech Scientific), 5 μL of 2nd-
65 round 5X primer mix, and 11.875 μL of PCR-grade water was amplified on a thermal cycler using
66 the following settings: 95°C for 5 m, 40 cycles of 95°C for 30 s, 55°C for 30 s, and 72°C for 1 m,
67 and a final extension of 72°C for 5 m. The presence of SARS-CoV-2 RNA was confirmed through
68 UV visualization of a PCR product of the expected length on a 2% agarose gel stained with Sybr-
69 Safe DNA Gel Stain (ThermoFisher Scientific).

70 **Real-time Reverse Transcription PCR**

71 Master Mix contents and PCR system settings were standardized for both primer/probe
72 concentrations. A 20 μL reaction mix containing 12.5 μL of 2X Quantitect Probe RT-PCR Master
73 Mix (Qiagen), 0.5 μL of RT mix from the kit, 5 μL of 5X 1st-round primer mix, and 2 μL of PCR-
74 grade water was mixed with 5 μL of RNA extract in a MicroAmp Fast Optical 96-well reaction
75 plate (ThermoFisher Scientific). Amplification was performed on an Applied Biosystems
76 StepOnePlus Real-Time PCR System (ThermoFisher Scientific) with the following settings: 50°C
77 for 30 m, 95°C for 15 m, and 40 cycles of 94°C for 15 s and 60°C for 1 m.

78 **Transcripts**

79 A 1.1kb transcript of the RdRP was synthesized from the first round nested PCR product of the
80 COV02 clinical sample. The fragment was ligated with the pJet1.2 plasmid using the CloneJET
81 PCR Cloning Kit (ThermoFisher Scientific). The recombinant plasmid was linearized using the
82 NcoI restriction enzyme (ThermoFisher Scientific) and the MEGAscript® T7 Kit (ThermoFisher
83 Scientific) was used for *in vitro* transcription of the fragment. NIBSC research reagent 19/304
84 containing packaged SARS-CoV-2 RNA was the kind gift of Giada Mattiuzzo. 19/304 was
85 extracted using the Viral RNA Mini Kit (Qiagen) and eluted into 50 μL of buffer AVE. Synthetic
86 SARS-CoV-2 RNA Control 1 - MT007544.1 and Control 2 - MN908947.3 were synthesized by
87 Twist Bioscience. The accession numbers for RNA control 1 and RNA control 2 have no
88 nucleotide differences in the binding locations of oligonucleotides used in this study.

89 **Sensitivity of PCR-based assays**

90 Experiments to determine the sensitivity of the two RT-qPCR methods and nested PCR were
91 completed using serial dilutions of each transcript (5×10^3 to 10^{-1} copies/5 μL) in a previously
92 described RNA storage buffer containing RNA storage solution (Thermo Fisher Scientific; 1 mM
93 sodium citrate, pH 6.4), herring sperm carrier RNA (50 $\mu\text{g}/\text{mL}$), and RNasin (New England

94 BioLabs UK, 100 U/mL).¹⁶ For nested PCR experiments, detection rate was assessed over five
95 replicates using the methods described above and a positive result was the presence of a PCR
96 product of the expected length. For RT-qPCR experiments, detection rate was assessed over
97 eight replicate experiments using the methods described above and a positive result was an
98 increase in signal that crossed the threshold value calculated by the machine for each experiment.
99 The CDC N1 method had a cutoff value of Ct 35.

100 **Statistical Analysis**

101 50% endpoints were estimated using the Reed-Muench method.¹⁷ Probit models were
102 constructed using function glm in R version 3.6.2.¹⁸

103

104

RESULTS

105

106 **Detection of Clinical Samples via Nested PCR**

107 To determine the ability of the nested PCR to detect and differentiate SARS-CoV-2 RNA in
108 extracts from clinical samples, 43 samples of known SARS-CoV-2 status were assessed. The
109 nested PCR successfully produced a PCR product of the expected size in 33/35 positive samples
110 (sensitivity of 94.2%) and a negative signal in 8/8 negative samples (specificity of 100%) (Figure
111 1, Figure S1). The nested PCR detected SARS-CoV-2 RNA from samples with a range of Ct
112 values – from a minimum of 17.11 to a maximum of 36.57. The two false negative samples had
113 relatively high Ct values of 29.6 and 34.66 but were not the most extreme targeted as
114 demonstrated in Figure 1.

115

116 **Comparative Sensitivity**

117 The sensitivity of the nested PCR and two RT-qPCR methods was compared by measuring the
118 50% endpoints (50EP) of detection for serial dilutions of the four transcripts described above
119 (Table 1, Figure 2). The RdRP transcript does not contain the target sequences of the CDC N1
120 method and thus was not assessed using this method. No method proved to consistently be the
121 most sensitive for all targets, and each method was the most sensitive for at least one target. The
122 RdRP transcript was detected consistently by the nested PCR with a 50EP of 5.61 copies, making
123 this the most sensitive for detecting the RdRp transcript. The Charité-RdRP RT-qPCR method

124 was the most sensitive for the 19/304 transcript with a 50EP of 50 copies. The CDC N1 method
125 was the most sensitive for both RNA Control 1 and RNA Control 2 with 50EPs of 33.4 and 15.0
126 copies, respectively. The differences between methods and a single target did not follow a clear
127 trend. Between the CDC N1 and Charité-RdRP RT-qPCR methods, the CDC N1 method showed
128 clear superiority for the RNA control 1 target (33.4 copies versus 223 copies) but only marginal
129 superiority for the RNA control 2 target (15.0 copies versus 17.1 copies). Strikingly, the nested
130 PCR was not able to detect RNA control 1 with any appreciable regularity even at the highest
131 copy number tested despite robust detection of RNA control 2.

132

133

DISCUSSION

134

135 This study establishes the first published nested PCR method for the sensitive detection of SARS-
136 CoV-2 in RNA extracted from nasopharyngeal swab samples. The nested PCR showed highly
137 concordant results with RT-qPCR performed at a hospital diagnostic laboratory, with a sensitivity
138 of 94.2%, specificity of 100%, and overall agreement of 95.3%. The use of nested PCR is a
139 practical alternative for laboratories without real-time PCR systems, but does require an RNA
140 extraction step. Whilst there has been a global shortage of necessary reagents,¹⁹ an increasing
141 number of novel extraction methods have been described in the literature.²⁰⁻²³ Furthermore,
142 nested PCR is also less amenable to automation and requires equipment for casting agarose gels
143 and amplicon visualization using UV light. The poor performance of the nested PCR for detecting
144 RNA control 1 is hypothesized to be due to the design of the transcript standard. Twist Bioscience
145 synthetic SARS-CoV-2 transcripts are six, non-overlapping 5 kb fragments. The drastically low
146 sensitivity but presence of PCR products of the correct length strongly suggests that the
147 fragmentation process separates the nF1 and nF2, nR1, and nR2 binding locations onto two
148 different fragments. This result underscores the notion that not all transcript standards are equal
149 for different PCR-based detection modalities.

150 Secondly, this study demonstrates inconsistent sensitivity for SARS-CoV-2 targets for different
151 PCR-based detection modalities. The results of this study point to remarkably different
152 performances between the three PCR methods analyzed for different SARS-CoV-2 RNA
153 transcripts. This inconsistency between different RT-qPCR methods has been noted by other
154 authors. In presenting the Charité-RdRP primer/probe set, Corman et al. found slight differences
155 between their E, N, and RdRP primer/probe sets, with the RdRP set performing the best with a

156 95% detection probability of 3.8 copies/reaction for SARS-CoV-2 genomic RNA.¹⁴ Corman et al.
157 also reported a 95% detection probability of 5.2 copies/reaction for the E gene assay – this is in
158 conflict with a 95% detection probability of 100 copies/reaction reported by Institut Pasteur using
159 the same primer/probe set.²⁴ Igloi et al. investigated the performance of 13 commercial RT-qPCR
160 kits and found variable performance between the RT-qPCR kits, with several kits having 10 fold
161 differences in sensitivity for different gene targets and one kit having a 2 log difference in
162 sensitivity between their E and RdRP/N preparations.²⁵ Using *in vitro* transcribed small transcripts
163 of five SARS-CoV-2 genes, Vogels et al. evaluated nine primer/probe sets, including the Charité-
164 RdRP and CDC N1 sets.²⁶ Vogels et al. found that eight of the nine primer/probe sets had lower
165 limits of detection of 500 copies/reaction, and that the Charité RdRP set was unable to detect any
166 replicates with 500 copies/reaction, although they did alter the primer and probe concentration
167 from those suggested by Corman et al. Lastly, Kudo et al. designed a multiplex RT-qPCR using
168 the CDC N1, N2, and RNase P primer and probe sets.²⁷ Using a full-length control, the multiple
169 RT-qPCR demonstrated a 50EP of 30.0 copies/reaction for the CDC N1 primers and 107.7
170 copies/reaction for the CDC N2 primers.

171 These results are compatible with anecdotal evidence from several labs in the United Kingdom
172 and elsewhere that consistent low level detection is unreliable and aberrant results are common.
173 This broader experience suggests that the issue may lie within the generalized approach of
174 detecting SARS-CoV-2 RNA by PCR rather than being specific to individual assays. One possible
175 factor limiting the sensitivity of PCR assays for SARS-CoV-2 may originate from the highly
176 structured nature of the genome as measured by standard thermodynamic RNA structure
177 prediction methods.²⁸ SARS-CoV-2, as well as other coronaviruses, has extensive RNA
178 secondary structure elements peppered throughout its genome – approximately 62% - 67% of
179 bases may be involved in base pairing. It is conceivable that, if a section of the target sequence
180 is embedded within highly energetically favoured RNA secondary structure, binding of the PCR
181 oligonucleotides could be competitively inhibited, delaying the initiation of the PCR reaction. This
182 effect could explain the unusual findings that no PCR based assay for SARS-CoV-2, whether in
183 this study or others, have been able to achieve single copy detection sensitivities.

184 Understanding the association between viral load and disease progression is important for the
185 selection of therapies. Tom et al.²⁹ recently proposed reporting RT-qPCR Ct values rather than
186 binary PCR test results within clinical reports as a means of estimating the viral load of patients
187 due to the inverse correlation of Ct value with copy number. While informative when tracking the
188 progression of Ct values for a single patient over time or comparing to other patients within the

189 same diagnostic pipeline, the results of this study do not support making direct comparisons
190 between data gained from different laboratories as Ct values are not readily comparable. For
191 these values to be informative, Ct values published in the literature should contain the methods
192 used as well as estimates of in-house correlation with copy. Alternatively, as also recommend by
193 Tom et al., reports could provide the Ct value as binned within ranges also seen in the laboratory
194 (i.e. in the top quartile of Ct values seen in this laboratory).

195 This current study is limited in that the standardized thermal cycler temperatures and
196 concentrations of Mg²⁺ recommended by the Quantitech RT-PCR kit were used for both the
197 Charité-RdRP and CDC N1 primers. It is feasible that slight alterations to the annealing
198 temperature and salt concentration could result in marginal increases in sensitivity, but these
199 changes would have to be assessed and calibrated on a lab by lab basis and may have different
200 values depending on the RT-qPCR master mix and PCR system used.

201

202 **ACKNOWLEDGEMENTS**

203

204 The authors acknowledge the staff of the Microbiology Department of the John Radcliffe Hospital
205 for their work in processing and testing clinical samples. The authors acknowledge Dr. Alexander
206 Mentzer for his assistance in procuring reagents for RT-qPCR and Dr. John Taylor for his gift of
207 RNA control 1 and RNA control 2. The authors want to especially thank Professor Tom Brown
208 and the team at ATDBio for rapid synthesis and delivery of oligonucleotides.

209

210 **CONFLICTS OF INTEREST**

211

212 The authors report no conflicts of interest.

REFERENCES

1. Zhu, N., et al., *A Novel Coronavirus from Patients with Pneumonia in China, 2019*. N Engl J Med, 2020. **382**(8): p. 727-733.
2. Zhou, P., et al., *A pneumonia outbreak associated with a new coronavirus of probable bat origin*. Nature, 2020. **579**(7798): p. 270-273.
3. Wu, F., et al., *A new coronavirus associated with human respiratory disease in China*. Nature, 2020. **579**(7798): p. 265-269.
4. World Health Organization (WHO), *Statement on the second meeting of the International Health Regulations (2005) Emergency Committee regarding the outbreak of novel coronavirus (2019-nCoV)*. 2020 30 January; Available from: [https://www.who.int/news-room/detail/30-01-2020-statement-on-the-second-meeting-of-the-international-health-regulations-\(2005\)-emergency-committee-regarding-the-outbreak-of-novel-coronavirus-\(2019-ncov\)](https://www.who.int/news-room/detail/30-01-2020-statement-on-the-second-meeting-of-the-international-health-regulations-(2005)-emergency-committee-regarding-the-outbreak-of-novel-coronavirus-(2019-ncov)).
5. World Health Organization (WHO), *WHO Director-General's remarks at the media briefing on 2019-nCoV on 11 February 2020*. 2020 11 February; Available from: <https://www.who.int/dg/speeches/detail/who-director-general-s-remarks-at-the-media-briefing-on-2019-ncov-on-11-february-2020>.
6. World Health Organization (WHO), *Coronavirus disease 2019 (COVID-19): situation report, 155*. 2020.
7. Dong, E., H. Du, and L. Gardner, *An interactive web-based dashboard to track COVID-19 in real time*. Lancet Infect Dis, 2020. **20**(5): p. 533-534.
8. Sun, K. and C. Viboud, *Impact of contact tracing on SARS-CoV-2 transmission*. Lancet Infect Dis, 2020.
9. Ferretti, L., et al., *Quantifying SARS-CoV-2 transmission suggests epidemic control with digital contact tracing*. Science, 2020. **368**(6491).
10. Li, R., et al., *Substantial undocumented infection facilitates the rapid dissemination of novel coronavirus (SARS-CoV-2)*. Science, 2020. **368**(6490): p. 489-493.
11. Mackay, I.M., K.E. Arden, and A. Nitsche, *Real-time PCR in virology*. Nucleic Acids Res, 2002. **30**(6): p. 1292-305.
12. van Pelt-Verkuil, E.v.B., A.; Hays, J. P., *Principles and Technical Aspects of PCR Amplification*. 2008: Springer.
13. Wu, Q., et al., *Development of Taqman RT-nested PCR system for clinical SARS-CoV detection*. J Virol Methods, 2004. **119**(1): p. 17-23.
14. Corman, V.M., et al., *Detection of 2019 novel coronavirus (2019-nCoV) by real-time RT-PCR*. Euro Surveill, 2020. **25**(3).
15. Centers for Disease Control and Prevention (CDC), *Coronavirus Disease 2019 (COVID-19)*. 2020; Available from: <https://www.cdc.gov/coronavirus/2019-ncov/lab/rt-pcr-panel-primer-probes.html>.
16. Hayes, A., et al., *A European multicentre evaluation of detection and typing methods for human enteroviruses and parechoviruses using RNA transcripts*. J Med Virol, 2019.
17. Reed, L.J.M., H., *A simple method of estimating fifty per cent endpoints*. American journal of epidemiology, 1938. **27**(3): p. 493-497.
18. R Core Team, *R: A language and environment for statistical computing*. 2013.
19. Babiker, A., et al., *SARS-CoV-2 Testing: Trials and Tribulations*. American Journal of Clinical Pathology, 2020. **153**(6): p. 706-708.
20. Merindol, N., et al., *SARS-CoV-2 detection by direct rRT-PCR without RNA extraction*. Journal of Clinical Virology, 2020. **128**: p. 104423.

11

21. Kalikiri, M.K.R., et al., *High-throughput extraction of SARS-CoV-2 RNA from nasopharyngeal swabs using solid-phase reverse immobilization beads*. medRxiv, 2020; doi: 2020.04.08.20055731.
22. Bruce, E.A., et al., *DIRECT RT-qPCR DETECTION OF SARS-CoV-2 RNA FROM PATIENT NASOPHARYNGEAL SWABS WITHOUT AN RNA EXTRACTION STEP*. bioRxiv, 2020; doi: 2020.03.20.001008.
23. Grant, P.R., et al., *Extraction-free COVID-19 (SARS-CoV-2) diagnosis by RT-PCR to increase capacity for national testing programmes during a pandemic*. bioRxiv, 2020; doi: 2020.04.06.028316.
24. Institut Pasteur, *Protocol: Real-time RT-PCR assays for the detection of SARS-CoV-2*. 2020; Available from: https://www.who.int/docs/default-source/coronaviruse/real-time-rt-pcr-assays-for-the-detection-of-sars-cov-2-institut-pasteur-paris.pdf?sfvrsn=3662fcb6_2.
25. Igloi, Z.I., M.; Abou-Nouar, Z. A-K.; Weller, B; Matheussen, V.; Coppens, J.; Koopmans, M; Molenkamp, R., *Comparison of commercial Realtime Reverse Transcription PCR Assays for the Detection of SARS-CoV-2*. J Clin Virol, 2020.
26. Vogels, C.B.F., et al., *Analytical sensitivity and efficiency comparisons of SARS-COV-2 qRT-PCR primer-probe sets*. medRxiv, 2020; doi: 2020.03.30.20048108.
27. Kudo, E., et al., *Detection of SARS-CoV-2 RNA by multiplex RT-qPCR*. bioRxiv, 2020; doi: 2020.06.16.155887.
28. Simmonds, P., *Pervasive RNA secondary structure in the genomes of SARS-CoV-2 and other coronaviruses – an endeavour to understand its biological purpose*. bioRxiv, 2020; doi: 2020.06.17.155200.
29. Tom, M.R. and M.J. Mina, *To Interpret the SARS-CoV-2 Test, Consider the Cycle Threshold Value*. Clin Infect Dis, 2020.

Table 1: Primer and Probe Sequences for Nested PCR and RT-qPCR

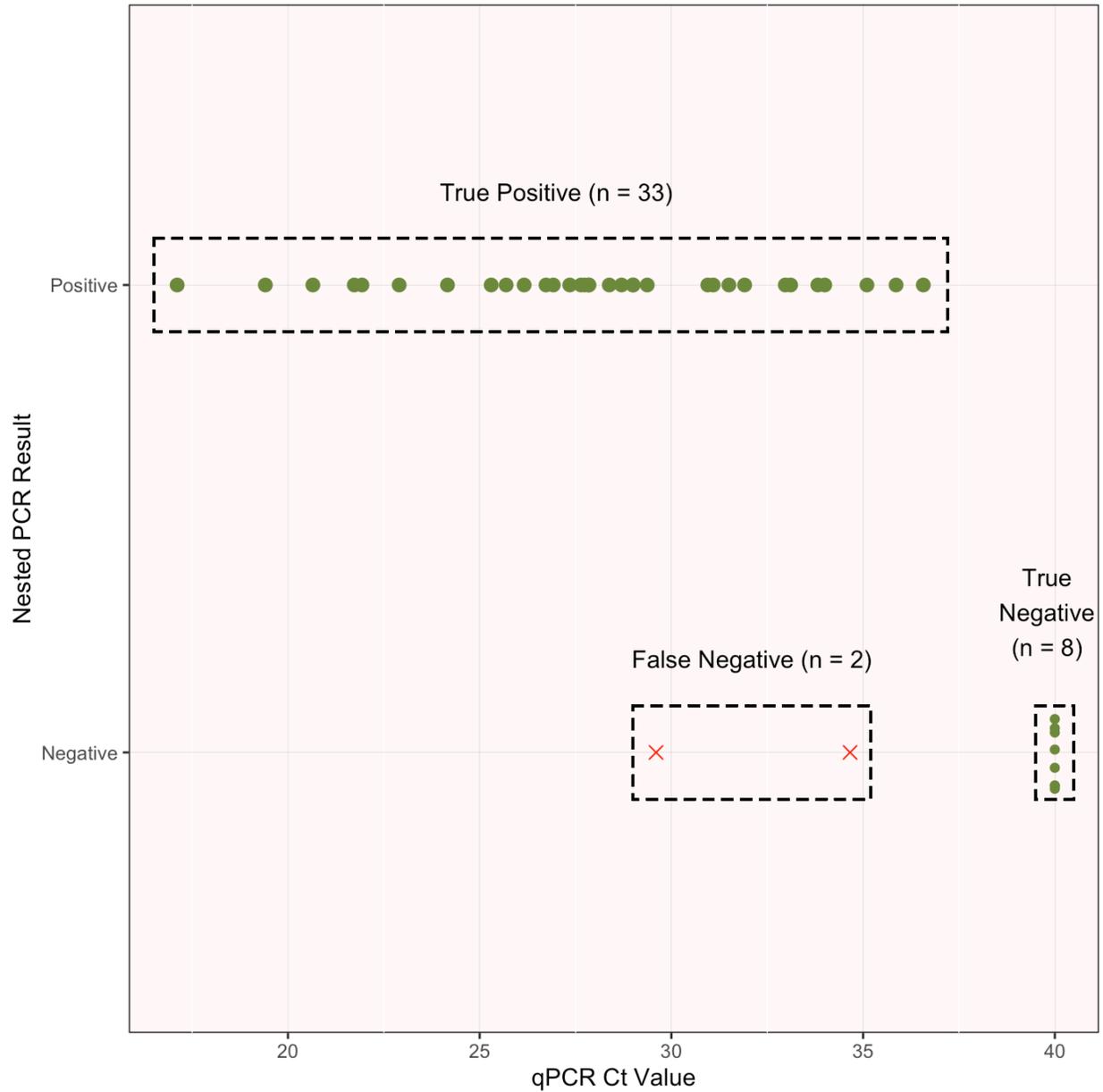
PCR Assay	Primer Name	Sequence	Reaction concentration
Nested PCR	nF1	AYTCAATGAGTTATGAGGAYCAAGATGC	400 nM
	nR1	GACATCAGCATACTCCTGATTWGGATG	400 nM
	nF2	TAGTACTATGACMAATAGACAGTTYCATC	500 nM
	nR2	CCTTTAGTAAGGTCAGTCTCAGTCC	500 nM
Charité-RdRP	RdRp_SARSr-F	GTGARATGGTCATGTGTGGCGG	600 nM
	RdRp_SARSr-P2	FAM-CAGGTGGAACCTCATCAGGAGATGC-BHQ	100 nM
	RdRp_SARSr-R	CAAATGTTAAARACACTATTAGCATA	800 nM
CDC N1	2019-nCoV_N1-F	GACCCCAAATCAGCGAAAT	500 nM
	2019-nCoV_N1-P	FAM-ACCCCGCATTACGTTTGGTGGACC-BHQ	125 nM
	2019-nCoV_N1-R	TCTGGTTACTGCCAGTTGAATCTG	500 nM

Table 2: 50% endpoints for PCR/target pairs

PCR Assay	Transcript Target (total copies/reaction)			
	19/304	RdRP Transcript	RNA Control 1	RNA Control 2
Nested	500	5.61	N.D.	70.7
Charité-RdRP	50	10	223	17.1
CDC N1	138	N/A	33.4	15.0

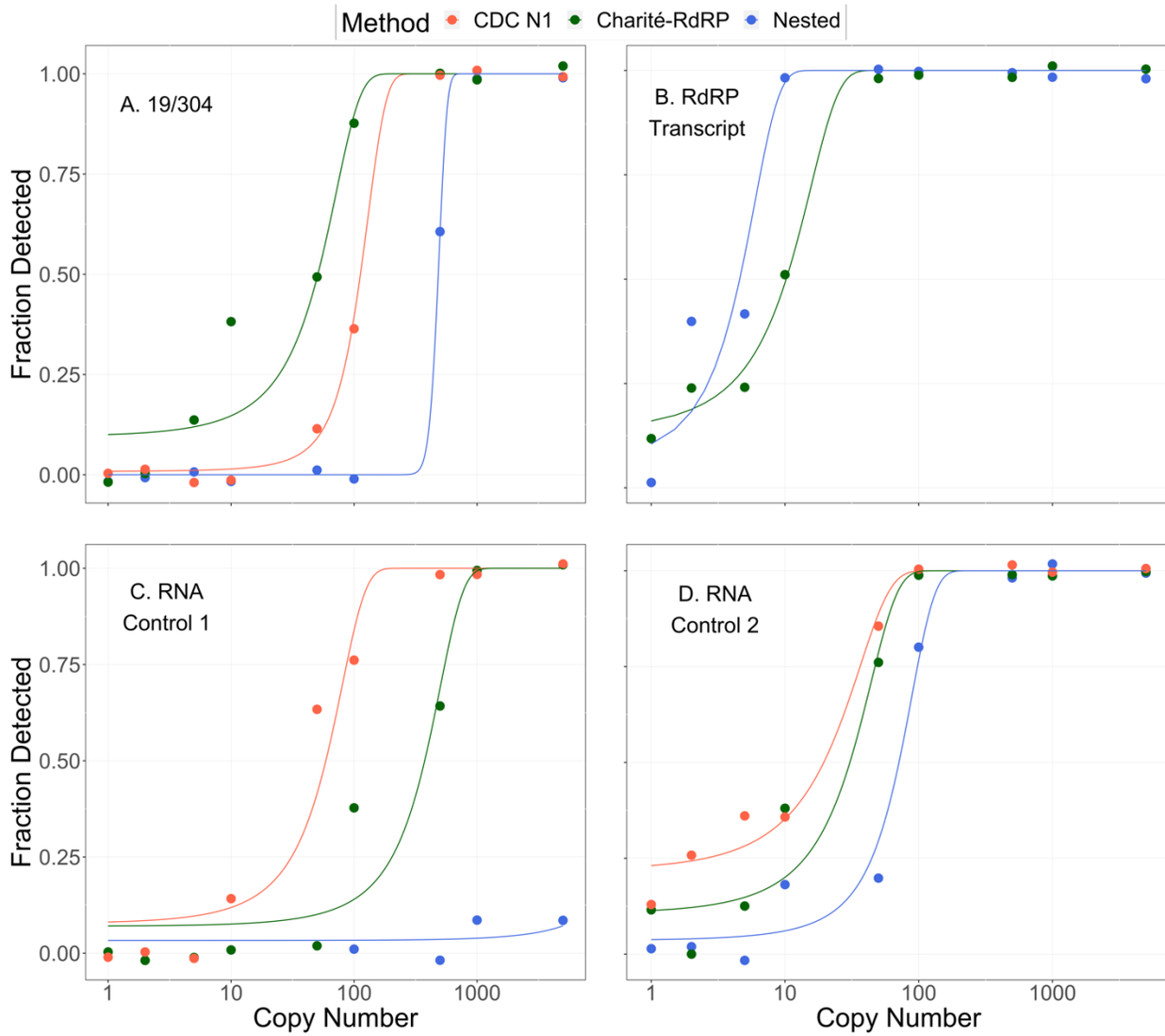
50% endpoints of detection for serial dilutions of four SARS-CoV-2 transcripts by three PCR assays. Nested PCR is detection rate over five replicates; Charité-RdRP and CDC N1 are detection rate over eight replicates. 50% endpoints were calculated by the Reed-Muench method.

Figure 1: Performance of Nested PCR for Clinical Samples



Comparison of detection of SARS-CoV-2 RNA in clinical samples by nested PCR versus clinical RT-qPCR result. Samples negative via RT-qPCR were given a representative Ct value of 40.

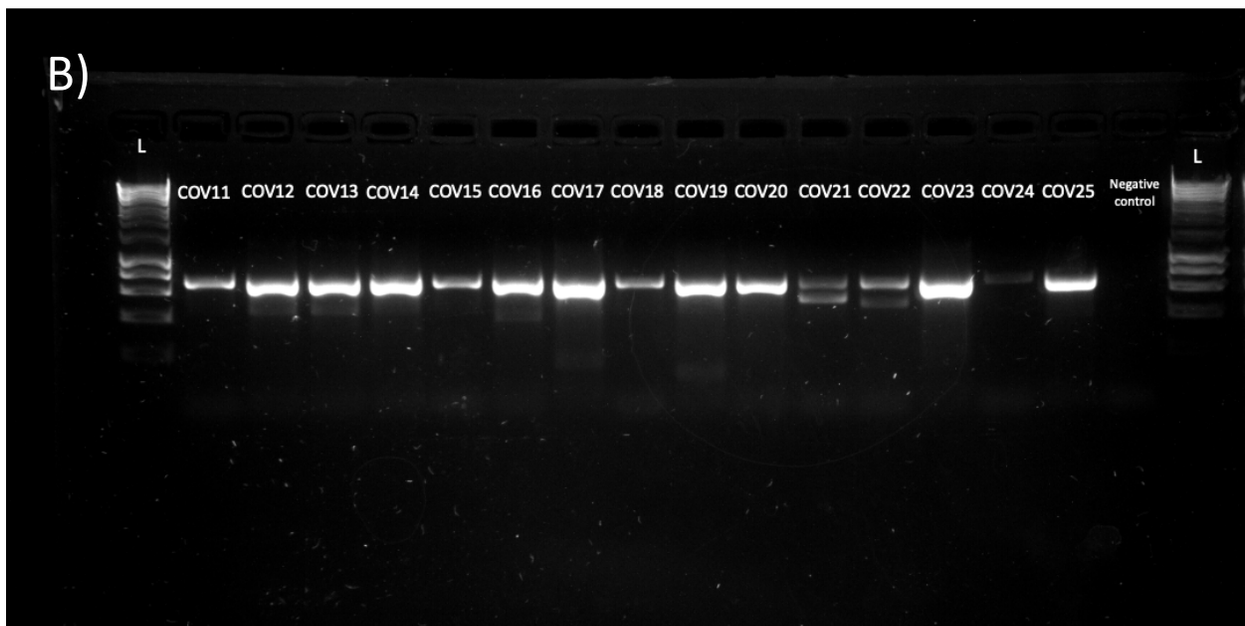
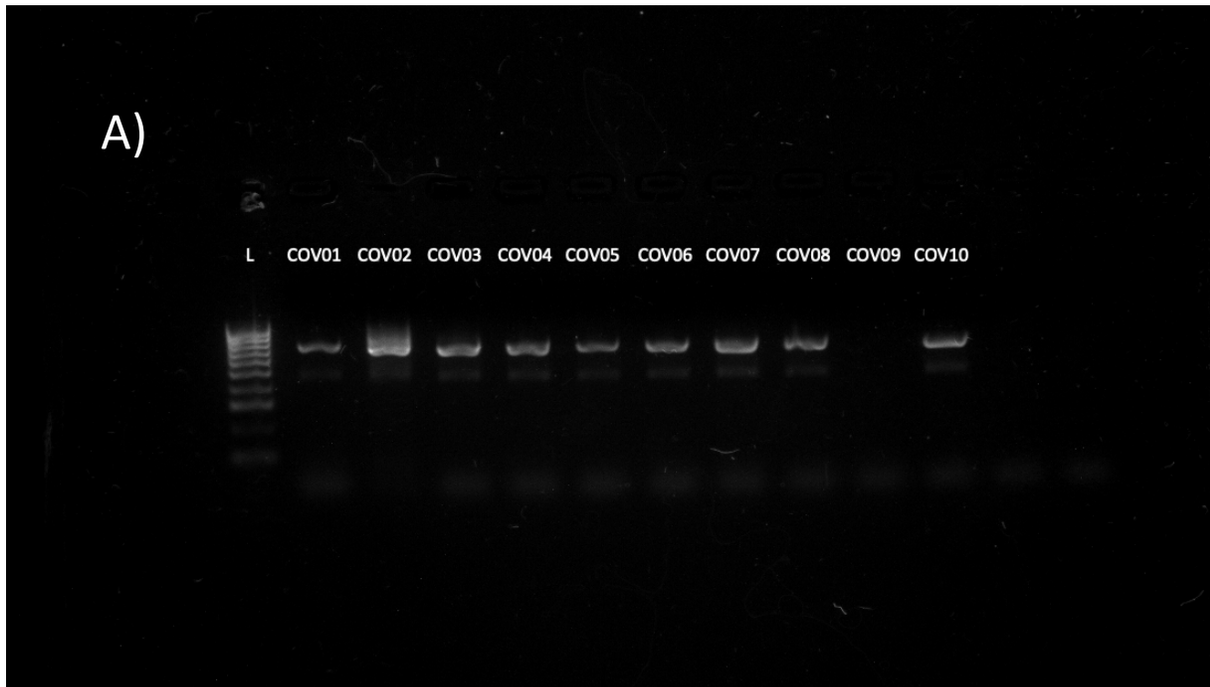
Figure 2: Comparative Detection Performance for Nested PCR and Charité-RdRP and CDC N1 RT-qPCR

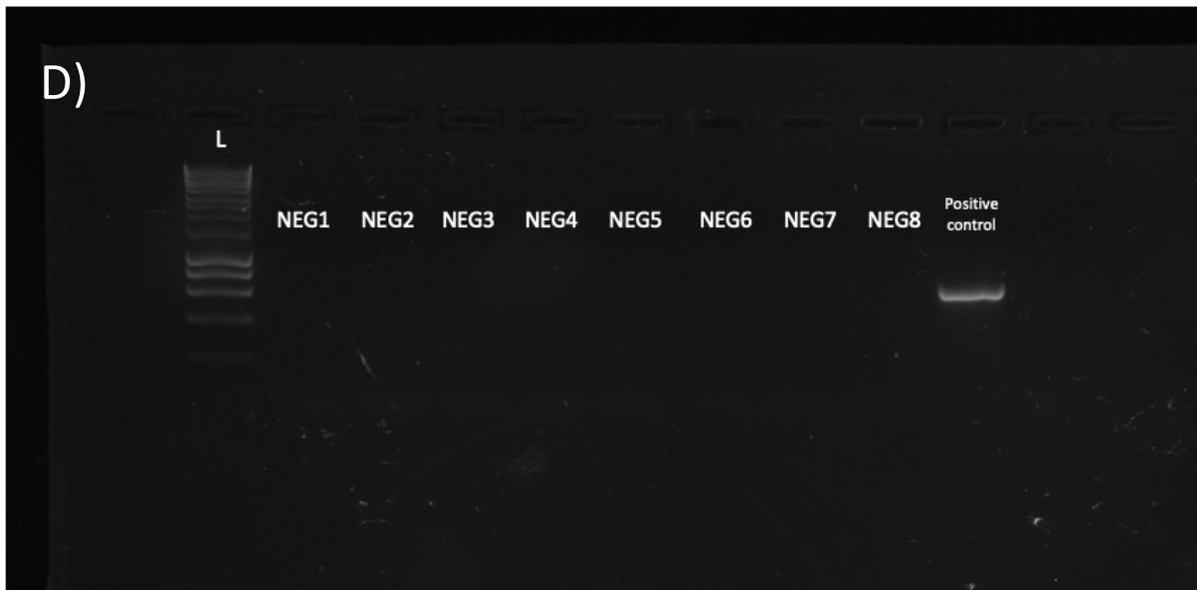
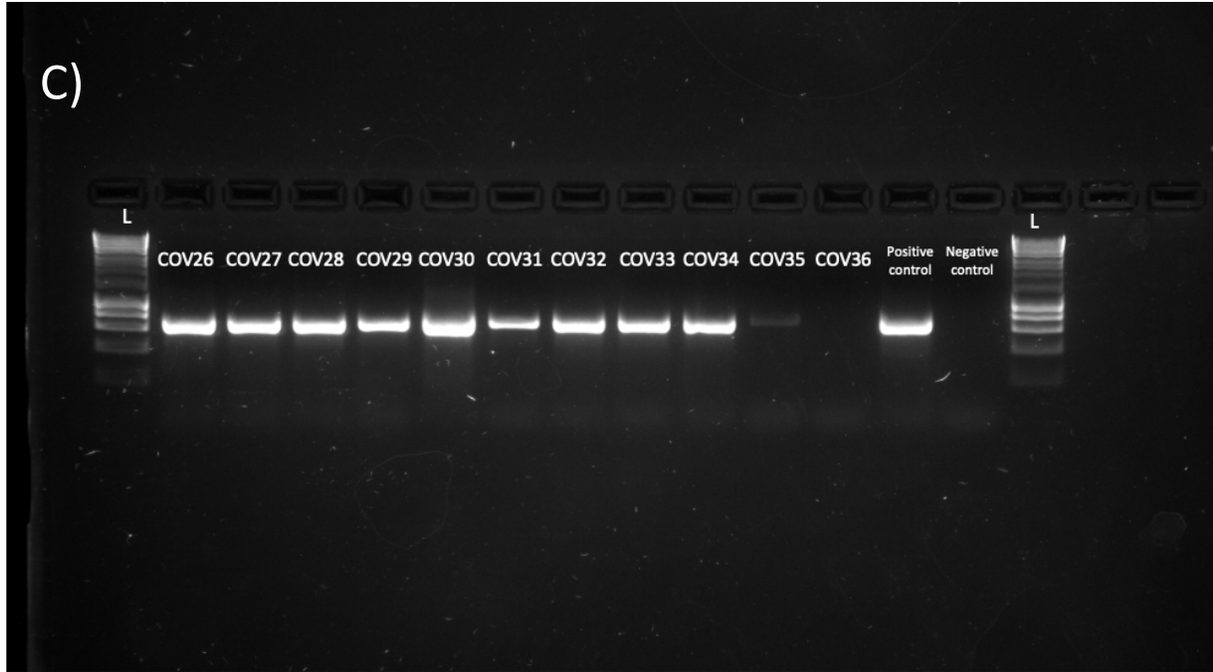


Comparative performance of detection methods for four targets. Dots for RT-qPCR experiments represent fraction detected of eight replicate reactions. Dots for nested PCR experiments represent fraction detected of five replicate reactions (15 replicates for RNA control 1). Lines represent probit regression models.

Supplementary Material

Figure S1: Nested PCR Amplification of Clinical Samples





Nested PCR detection of clinical samples assessed via RT-qPCR. COV1-COV36 (panels A-C) were positive samples, NEG1-NEG8 (Panel D) were negative samples. Panel A has a 100 bp molecular ladder (New England Biolabs); Panels B-D have 1kb molecular ladder (New England Biolabs).