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## The estimation of diagnostic accuracy of tests for COVID-19: A scoping review

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### SUMMARY

**Objectives:** To assess the methodologies used in the estimation of diagnostic accuracy of SARS-CoV-2 real-time reverse transcription polymerase chain reaction (rRT-PCR) and other nucleic acid amplification tests (NAATs) and to evaluate the quality and reliability of the studies employing those methods.

**Methods:** We conducted a systematic search of English-language articles published December 31, 2019–June 19, 2020. Studies of any design that performed tests on  $\geq 10$  patients and reported or inferred correlative statistics were included. Studies were evaluated using elements of the Quality Assessment of Diagnostic Accuracy Studies (QUADAS-2) guidelines.

**Results:** We conducted a narrative and tabular synthesis of studies organized by their reference standard strategy or comparative agreement method, resulting in six categorizations. Critical study details were frequently unreported, including the mechanism for patient/sample selection and researcher blinding to results, which lead to concern for bias.

**Conclusions:** Current studies estimating test performance characteristics have imperfect study design and statistical methods for the estimation of test performance characteristics of SARS-CoV-2 tests. The included studies employ heterogeneous methods and overall have an increased risk of bias. Employing standardized guidelines for study designs and statistical methods will improve the process for developing and validating rRT-PCR and NAAT for the diagnosis of COVID-19.

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### Introduction

After its emergence in December 2019, the virus now known as SARS-CoV-2 was identified and sequenced in early January 2020,<sup>1</sup> allowing for the rapid development of diagnostic testing based on the detection of viral nucleic acid (i.e., real-time reverse transcription polymerase chain reaction [rRT-PCR]).<sup>2</sup> Because infected patients can present with non-specific symptoms or be asymptomatic, the development of accurate diagnostic tests for both clinical and epidemiological purposes was a crucial step in the response to the COVID-19 pandemic.<sup>3</sup>

In the United States, the spread of SARS-CoV-2 rapidly outpaced the capacity to test for it, resulting in the Food and Drug Administration (FDA) relaxing regulatory requirements to increase

testing availability. The FDA granted the first Emergency Use Authorization (EUA) for a SARS-CoV-2 rRT-PCR diagnostic test on February 4, 2020. Consequently, hundreds of tests for SARS-CoV-2, among them rRT-PCRs, other types of nucleic acid amplification tests (NAATs), and automated and/or multiplex methods based on proprietary platforms, obtained FDA Emergency Use Authorization (EUA). As of August 4th, 2020, the FDA has granted EUAs to 203 diagnostic tests, including 166 molecular tests, 35 antibody assays, and 2 antigen tests. Although the FDA began requiring the submission of validation methods and results as part of EUA application for SARS-CoV-2 diagnostic tests, these tests were not initially required to undergo the rigorous assessment that would normally be part of the FDA approval process. Researchers also began developing alternative nucleic-acid based methodologies to detect SARS-CoV-2, including reverse-transcription loop-mediated isothermal amplification (RT-LAMP), and others.

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Concurrently with rapid test production, publications emerged reporting clinical diagnostic test performance characteristics, such as “sensitivity” and “specificity”, though some lacked the rigorous methodologies usually required to formally estimate diagnostic accuracy. Here we present a scoping review of the literature with two main objectives: 1) to assess the methodologies used in the estimation of diagnostic accuracy of SARS-CoV-2 tests and 2) to evaluate the quality and reliability of the studies employing those methods.

**Methods**

*Data sources and searches*

Searches were performed through MEDLINE (Ovid), EMBASE (Elsevier), Scopus, Web of Science, CINAHL, and PubMed following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines<sup>4</sup> between December 1, 2019 and June 19, 2020. The following search string was used: (2019-nCoV or SARS-CoV-2 or SARS-CoV2 or COVID-19 or COVID19 or COVID) and (“positive agreement” or “negative agreement” or “overall agreement” or “diagnostic accuracy” or “positive rate” or “positivity rate” or “test performance” or “reference standard” or “gold standard” or sensitivity or specificity or “percent agreement” or “concordance” or “test agreement” or “predictive value” or “false negative” or “false positive”) and (“polymerase chain reaction” or PCR or “reverse transcriptase” or “nucleic acid amplification test” or NAAT or isothermal or “RT-LAMP” or “RT-PCR” or “molecular test”). The literature hub LitCovid’s “Diagnosis” section was screened in its entirety once and then daily for relevant titles.

*Study selection*

We liberally screened articles by title and abstract for further evaluation. Articles were included if they met the following criteria on screening: 1) Peer-reviewed publication, 2) Study evaluated diagnostic test accuracy of NAAT, 3) Diagnostic test performed on ≥10 patients, 4) Diagnostic/Clinical sensitivity, specificity, other correlative statistics, or test positive rate were either identified by name or were included in the publication as a numerical value and we could reproduce the calculations. Exclusion criteria included: 1) Pre-print status, 2) Guidelines, consensus, review, opinion, and other summary articles 3) Entirely pregnant or pediatric populations, 4) Overlap of study population with another included publication.

*Data extraction and quality assessment*

Four authors independently extracted data and two authors reviewed data for accuracy. For study characteristics, we extracted: first author name, country, study design, patient population, total number of patients or samples included in test performance calculations, and number of cases according to rRT-PCR (Tables 1–5) or total number of cases based on positive result of any platform tested (Table 6). For patient characteristics, we extracted age and sex. For index test and reference standard characteristics, we extracted: test type (NAAT) or definition (clinical diagnosis, composite reference standards), specimen (NAAT), specimen dry/collection liquid status (for studies evaluating Abbott ID NOW), proprietary automated and/or multiplex systems – henceforth called “platforms” (NAAT), and target genes of primers (NAAT). For outcomes, we extracted the values of test performance characteristics with their designation according to the original authors, without our interpretation. For this reason, we indicate these outcomes as “reported” (r): reported sensitivity (rSN), specificity (rSP), positive predictive value (rPPV), negative predictive value (rNPV), accuracy

**Table 1** Studies reporting the “positive rate” of rRT-PCR testing within a population of patients suspected to have COVID-19.

Authors	Country	Study Type	No. Patients Total	rRT-PCR positive*	Demographics† Age (y)‡ % Male	Index Test Type Specimen (No.)	Primers	rRT-PCR Kit Company	Reference Standard: Case Definition/Clinical Diagnosis	Study Findings PR§ (95% CI)
Ai et al. <sup>7</sup>	China	Cross Sectional	1014	601	51 ± 15 46.0%	rRT-PCR Throat Swab (1014)	ORF1ab, N	Shanghai Huirui Biotech, Shanghai BioGerm	“patients...who were suspected of novel coronavirus infection” (p. 5)	59% (56 - 62)
Liu et al. <sup>8</sup>	China	Cross Sectional	4880	1875	50 (IQR 27) 46.13%	rRT-PCR Nasal swab, pharyngeal swab, BAL, sputum	ORF1ab, N	Shanghai Huirui Biotech	“All the cases were suspected of SARS-CoV-2 infection because of, (1) typical respiratory infection symptoms such as fever, cough and dyspnea, or (2) close contact with a COVID-19 patient.” (p.172)	38.42%
Xie et al. <sup>9</sup>	China	Cross Sectional	19	9	33 (8 - 62) 42.1%	rRT-PCR OP swab (19)	ORF1ab, N	GeneDx, Maccura, Life-river	“...suspected cases...” (p. 264)	47.4%

\* Reported instead of “cases according to reference standard” as present in other tables.

† Of cohort or cases.

‡ Format: median (range), median(IQR), or mean±SD.

§ PR: positive rate, which is the number of rRT-PCR patients out of the number of patients suspected to have COVID-19 (i.e. the reference standard). Patient population.

|| Hospitalized patients. Abbreviations- BAL: bronchoalveolar lavage, CI: confidence interval, IQR: Interquartile range, N: nucleocapsid, OP: oropharyngeal, ORF1ab: open reading frame 1ab, rRT-PCR: real-time Reverse Transcription Polymerase Chain Reaction, y: years.

**Table 2**

Studies reporting test performance characteristics of initial rRT-PCR result compared to result after repeated tests of rRT-PCR as reference standard.

Authors	Country	Study Type	Demographics <sup>†</sup>		Specimen Type (No.)	Primers/Platform No. samples (%)	No. pts with 1st rRT-PCR positive	Total No. of pts with rRT-PCR ever positive	Total No. of pts in calculations*	Interval (d) between each re-test	No. of tests performed per pt until positive	Interval (d) between initial and positive (final) rRT-PCR	Study Findings			PR	
			Age (y) <sup>‡</sup>	% Male									Correlative Statistics (95% CI)	rSN	rSP		rAcc
Bernheim et al. <sup>10</sup>	China	Cases only <sup>  </sup>	45.3 ± 15.6	50.4%	NPS, OPS, Trach Asp, BAL (nr)	ORF1ab, N rRT-PCR	90	102	102	nr	1 test: 90 >1 test: 12	nr				88%	
Fang et al. <sup>11</sup>	China	Cases only <sup>  </sup>	45 (IQR 39–55)	56.9%	Throat swab (45), sputum (6)	ORF1ab, N rRT-PCR	36	51	51	≥1	1 test: 36 2 tests: 12 3 tests: 2 4 tests: 1	nr	71% (56–83)				
Green et al. <sup>12</sup>	USA	Cohort <sup>  </sup>	53.1 ± 22.3	45.5%	NPS, OPS (nr)	RdRp, E Roche cobas 19,195 (70.1%)	10,070	17,405**	22,061	median 8 range (1 – 49)	1 test: 10,070 >1 test: 7335**	1 - 49	Lower bound estimate:** 57.9% (55.2–60.5)				
						N2, E Cepheid Xpert 6219 (22.7%) N1, N2 rRT-PCR 1884 (6.9%)											
						RdRp Abbott ID NOW 53 (0.2%) ORF1ab (x2) Hologic Panther 26 (0.1%)	10,643**	22,061	median 8 range (1 – 49)	1 test: 10,070 >1 test: 573**	1 - 49	Upper bound estimate:** 94.6% (94.2–95.0)					

(continued on next page)

Table 2 (continued)

Authors	Country	Study Type	Demographics <sup>†</sup>		Specimen Type (No.)	Primers/Plat-form No. samples (%)	No. pts with 1st rRT-PCR positive	Total No. of pts with rRT-PCR ever positive	Total No. of pts in calculations*	Interval (d) between each re-test	No. of tests performed per pt until positive	Interval (d) between initial and positive (final) rRT-PCR	Study Findings			PR
			Age (y) <sup>‡</sup>	% Male									Correlative Statistics (95% CI)	rSN	rSP	
He et al. <sup>13</sup>	Hong Kong	Case-Control <sup>  </sup>	52 (8 - 74)	50%	NPS, OPS, Trach Asp, BAL (nr)	RdRp, S rRT-PCR	27	34	82	nr	1 test: 27 >1 test: 7	1–14	79% (66–93)	100% (100)	92% (91–92)	
Lee et al. <sup>14</sup>	Singapore	Cases only <sup>  </sup>	nr	nr	NPS (70)	ORF1ab, N rRT-PCR	62	70	70	1st-2nd: 1 2nd-X <sup>th</sup> : 1–2	1 test: 62 2 tests: 5 3 tests: 1 5 tests: 1 6 tests: 1	1st-2nd: 1 2nd-Final: 2,4,7 2 - 8				88.6%
Long et al. <sup>15</sup>	China	Cases only <sup>  </sup>	44.8 ± 18.2	55.6%	OPS, NPS (nr)	ORF1ab, N rRT-PCR	30	36	36	nr	1 test: 30 2 tests: 3 3 tests: 3	2 - 8	83.3%			
Wong et al. <sup>16</sup>	Hong Kong	Cases only <sup>  </sup>	56 ± 19	40.6%	NPS, throat swab (nr)	RdRp/Hel rRT-PCR	58	64	64	nr, "not uniform"	1 test: 58 >1 test: 6	nr	91% (83–97)			
Wu et al. <sup>17</sup>	China	Cases only <sup>  </sup>	46.1 ± 15.4	48.8%	Nose swab, throat swab (nr)	ORF1ab, N rRT-PCR	41	80	80	1	1 test: 41 2 tests: 30 3 tests: 9	1 - 2				51.25%

\* Pts included in test performance calculations.

<sup>†</sup> Of cohort or cases.

<sup>‡</sup> Format: median (range), median(IQR), or mean±SD. Patient Population.

<sup>||</sup> Hospitalized patients. Abbreviations- BAL: Bronchoalveolar lavage, CI: confidence interval, d: days, E: envelope, Hel: helicase, IQR: Interquartile range, N: nucleocapsid, No.: number, NPS: nasopharyngeal swab, nr: not reported, OPS: oropharyngeal swab, ORF1ab: open reading frame 1ab, PR: positive rate, pts: patients, rRT-PCR: real-time Reverse Transcription Polymerase Chain Reaction, rAcc: reported accuracy, RdRp: RNA-dependent RNA polymerase, rSN: reported sensitivity, rSP: reported specificity, S: spike, Trach Asp: Tracheal Aspirate, y: years.

\*\* Sensitivity estimates for the first test conducted on patients were calculated based on different assumptions about true negatives. The estimate of the upper bound estimate assumes that any negative test results (whether negative on a single test or consistently negative across multiple, repeated tests) was a true negative (aka, false negative rate=0%). The estimate of the lower bound uses the proportion of repeatedly tested cases who *initially* tested negative but then tested positive in repeated tests to calculate a false negative rate (16.8%) and apply that rate to the patients who only received a single test to calculate an assumed number of false negative cases. Additional details are provided in Suppl. Fig. 1 and Green et al.

**Table 3**  
Studies that calculate test performance characteristics of rRT-PCR or automated rRT-PCR platforms compared to composite reference standards.

Authors	Country	Study Type	No. Patients		Demographics <sup>†</sup>		Specimen	Index Test			Composite Reference Standard Definition	Study Findings (95% CI)							
			Total	Cases*	Age (y) <sup>‡</sup>	% M		Type	Primers	Platform		rSN	rSP	rPPV	rNPV	rAcc	rPPA	rNPA	Cohen's $\kappa$
Cradic et al. <sup>18</sup>	USA	Cohort <sup>§</sup>	184	33	nr	nr	NPS in VTM (184)	Automated Multiplex rRT-PCR	ORF1ab, S	Diasorin Simplexa	Result obtained from at least 2 of the 3 assays is consensus result.						100% (90–100)	100% (98–100)	
								Automated Isothermal NAAT	RdRp	Abbott ID NOW							91% (79–97)	100% (98–100)	
								Automated Multiplex rRT-PCR	ORF1, E	Roche cobas 6800							100% (90–100)	100% (98–100)	
Suo et al. <sup>19,a</sup>	China	Cohort <sup>**</sup>	58	52	nr	nr	Throat swab (58)	Initial rRT-PCR (China CDC protocol)	ORF1ab, N	N/A	Positive result of repeated rRT-PCR, or serology is considered a positive result.	40% (27–55)	100% (54–100)	100% (N/A)	16% (13–19)	47% (33–60)			
Zhen & Mangi et al. <sup>20</sup>	USA	Case Control <sup>††</sup>	104	51	nr	nr	NPS (104)	rRT-PCR (US CDC protocol)	N1, N2	N/A	Result obtained by 3 out of 4 assays tested is consensus result						100% (93–100)	98% (89–99)	0.98 (0.94–1)
								Automated multiplex rRT-PCR	ORF1ab, S	Diasorin Simplexa							100% (93–100)	100% (93–100)	1.0 (0.99–1)
								Automated rRT-PCR w/sensor	N	GenMark ePlex							96% (87–99)	100% (93–100)	0.96 (0.91–1)
								Automated multiplex rRT-PCR	ORF1ab (2 targets)	Hologic Panther							100% (93–100)	96% (87–99)	0.96 (0.91–1)

\* Cases according to composite reference standard.

<sup>†</sup> Of cohort or cases.

<sup>‡</sup> Format: median (range), median(IQR), or mean $\pm$ SD. Patient Population.

<sup>§</sup> Hospitalized patients.

<sup>¶</sup> Emergency Room patients.

<sup>\*\*</sup> Outpatients, some of whom were later hospitalized.

<sup>††</sup> not reported.

<sup>a</sup> Suo et al. data is also present in Table 4. Abbreviations- E: envelope, IQR: Interquartile range,  $\kappa$ : kappa coefficient, M: male, N/A: not applicable, N: nucleocapsid, NPS: nasopharyngeal swab, nr: not reported, ORF1ab: open reading frame 1ab, rRT-PCR: real-time Reverse Transcription Polymerase Chain Reaction, rAcc: reported accuracy, RdRp: RNA-dependent RNA polymerase, rNPA: reported negative percent agreement, rNPV: reported negative predictive value, rPPA: reported positive percent agreement, rPPV: reported positive predicted value, rSN: reported sensitivity, rSP: reported specificity, S: spike, VTM: viral transport media, y: years.

**Table 4**  
Studies reporting test performance characteristics of other nucleic acid amplification test methods compared to rRT-PCR.

Authors	Country	Study Type	No. Patients		Demographics <sup>†</sup>		Specimen (No.)	Index Test Type	Primers	Platform	Ref Stnd: rRT-PCR Primers	Study Findings (95% CI or <i>p</i> -value)						
			Total	Cases*	Age (y) <sup>‡</sup>	% Male						rSN	rSP	rPPV	rNPV	rAcc	rOA	Cohen's $\kappa$
Baek et al. <sup>21</sup>	Korea	Case Control <sup>  </sup>	154	14	nr	nr	Nasal swab (154)	RT-LAMP	N	nr	ORF1ab, S	100%	98.7%					0.826
Kitagawa et al. <sup>22</sup>	Japan	Cohort <sup>  </sup>	76	30	nr	nr	NPS (76)	RT-LAMP	nr	LA-200	N	100%	95.6%				97.4%	
Lau et al. <sup>23</sup>	Malaysia	Case Control <sup>††</sup>	89 <sup>‡‡</sup>	47 <sup>‡‡</sup>	nr	nr	NPS (89)	RT-LAMP	N	tur-bidimeter	RdRp,E	100%	100%					
Lu et al. <sup>24</sup>	China	Case Control <sup>  </sup>	56	36	nr	nr	Throat swab (56)	RT-LAMP	N	nr	ORF1ab, N					92.9%		
Yan et al. <sup>25</sup>	China	Cohort <sup>††</sup>	130	58	nr	nr	Throat swab, BAL (nr)	RT-LAMP	ORF1ab, S	nr	ORF1ab, N	100% (92.3–100)	100% (93.7–100)					
Wang, Cai, & He et al. <sup>26</sup>	China	Cohort <sup>††</sup>	947	338	44 ± 17.1	60%	OPS (834), sputum (82), NPS (16), nasal swab (8), BAL (4), stool (2), blood (1)	RT-RAA	ORF1ab	RAA-F1620 fluorescent detector	ORF1ab &N, or ORF1ab	97.6% (95.2–98.9)	97.8% (96.2–98.8)	96.2% (93.4–97.8)	98.6% (97.3–99.3)			0.952 <i>p</i> < 0.05
Xue et al. <sup>27</sup>	China	Cohort <sup>††</sup>	120 <sup>‡‡</sup>	22 <sup>‡‡</sup>	nr	nr	NPS, sputum (nr)	RT-RAA	ORF1ab	RAA-1620 fluorescent detector	ORF1ab, S	100%	100%					1.0 <i>p</i> < 0.001
Perchetti et al. <sup>28</sup>	USA	Case Control <sup>††</sup>	356	186	nr	nr	NPS (356)	Triplex rRT-PCR	N1, N2	n/a	N1, N2	98.4%	100%				99.2%	
Waggoner et al. <sup>29</sup>	USA	Cohort <sup>  </sup>	27	11	nr	nr	NPS, OPS (nr)	Triplex rRT-PCR	N2, E	n/a	N2, E						100%	
Li et al. <sup>30</sup>	China	Cohort <sup>††</sup>	303 <sup>‡‡</sup>	126 <sup>‡‡</sup>	nr	nr	throat swab (267), sputum (22), nose swab (8), BAL (3), blood (3)	AIGS	ORF1ab, N, S	LifeReady 1000	ORF1ab, N	97.62% (93.2–99.5)	100%					

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Table 4 (continued)

Authors	Country	Study Type	No. Patients		Demographics <sup>†</sup>		Specimen (No.)	Index Test Type	Primers	Platform	Ref Stnd: rRT-PCR Primers	Study Findings (95% CI or <i>p</i> -value)						Cohen's $\kappa$
			Total	Cases*	Age (y) <sup>‡</sup>	% Male						rSN	rSP	rPPV	rNPV	rAcc	rOA	
Suo et al. <sup>19,a</sup>	China	Cohort**	58	52	nr	nr	Throat swab (58)	ddPCR	ORF1ab, N	QX200 System	ORF1ab, N	94% (83–99)	100% (48–100)	100% (NA)	63% (36–83)	95% (84–99)		
Bulterys et al. <sup>31</sup>	USA	Cohort <sup>††</sup>	80	30	nr	nr	NPS (80)	Isothermal amplification	ORF1ab, N	Atila iAMP kit	E	82.8% (65.0–92.9)					0.86 (0.74–0.98)	
Wang, Cai, & Zhang et al. <sup>32</sup>	China	Cohort <sup>††</sup>	181 <sup>‡‡</sup>	25 <sup>‡‡</sup>	nr	nr	Throat swab (181)	OSN-qRT-PCR	ORF1ab, N	Life Tech. 480	ORF1ab, N						0.737	

\* Cases according to reference standard.

<sup>†</sup> Of cohort or cases.<sup>‡</sup> Format: median (range), median(IQR), or mean $\pm$ SD. Patient Population.<sup>||</sup> Hospitalized patients<sup>¶</sup>Emergency Room/Immediate Care Center patients.<sup>\*\*</sup> Outpatients, who were later hospitalized.<sup>††</sup> not reported.<sup>‡‡</sup> Number of samples (when number of patients not reported).

<sup>a</sup> Suo et al. data is also present in Table 2. Abbreviations- AIGS: Automatic integrated gene detection system, BAL: Bronchoalveolar lavage, CI: confidence interval, ddPCR: digital droplet polymerase chain reaction, E: envelope, iAMP: isothermal amplification, IQR: Interquartile range,  $\kappa$ : kappa statistic, n/a: not applicable, N: nucleocapsid, No.: number, NPS: nasopharyngeal swab, nr: not reported, OPS: oropharyngeal swab, ORF1ab: open reading frame 1ab, OSN-qRT-PCR: one-step single-tube nested quantitative real-time polymerase chain reaction, rAcc: reported accuracy, RdRp: RNA-dependent RNA polymerase, Ref Stnd: reference standard, rNPA: reported negative percent agreement, rNPV: reported negative predictive value, rOA: reported overall agreement, rPPA: reported positive percent agreement, rPPV: reported positive predictive value, rRT-PCR: real-time Reverse Transcription Polymerase Chain Reaction, rSN: reported sensitivity, rSP: reported specificity, RT-LAMP: reverse transcription loop-mediated isothermal amplification, RT-RAA: reverse-transcription recombinase-aided amplification, S: spike, y: years.



**Table 5**  
Studies estimating NAAT platform test performance characteristics compared to rRT-PCR as the reference standard.

	Country	Study Type	No. Patients		Demographics <sup>†</sup>		Specimen	Index Test			Ref Stnd: rRT-PCR Primers	Study Findings (95% CI)						
			Total	Cases <sup>*</sup>	Age (y) <sup>‡</sup>	% M		Type	Primers	Platform		rSN	rSP	rPPV	rNPV	rPPA	rNPA	rOA
Mitchell et al. <sup>33</sup>	USA	Case Control <sup>††</sup>	61 <sup>‡‡</sup>	46 <sup>‡‡</sup>	nr	nr	NPS in VTM (61)	Automated Isothermal NAAT	RdRp	Abbott ID NOW	N1, N2	71.70%	100%				78.70%	
Rhoads et al. <sup>35,b</sup>	US	Cases only <sup>  </sup>	96 <sup>‡‡</sup>	96 <sup>‡‡</sup>	nr	nr	NPS (85), nasal swab (11) in NS or UTM	Automated Isothermal NAAT	RdRp	Abbott ID NOW	N1, N2			94% (87–98)				
Moore et al. <sup>34,a</sup>	USA	Cohort <sup>  §,***</sup>	200	119	50 ± 17	46%	NPS in VTM (200)	Automated Isothermal NAAT	RdRp	Abbott ID NOW				80.3% (71.9–87.1)	100% (95.4–100)			
DegliAngeli et al. <sup>36</sup>	USA	Case Control <sup>††</sup>	60 <sup>‡‡</sup>	30 <sup>‡‡</sup>	nr	nr	Nasal swab, NPS (nr)	Automated Multiplex RT-PCR	RdRp, N	Abbott RealTime	N1, N2	93%	100%					
Hou et al. <sup>37</sup>	China	Cohort <sup>  ,***</sup>	285	153	<65 y: 77.2%	55.8%	OPS (285)	Automated Multiplex RT-PCR	N2, E	Cepheid Xpert Xpress	ORF1ab, N			96.1% (91.3–98.4)	96.2% (90.9–98.6)	96.1% <sup>§</sup> (0.88–0.97)	0.92 (0.88–0.97)	
Lieberman et al. <sup>38</sup>	USA	Cohort <sup>††</sup>	26 <sup>‡‡</sup>	13 <sup>‡‡</sup>	nr	nr	NPS (26)	Automated Multiplex RT-PCR	N2, E	Cepheid Xpert Xpress	N1, N2					100% <sup>§</sup>		
Loeffelholz et al. <sup>39,a</sup>	USA, UK, FR, IT	Cohort <sup>††</sup> (enriched for positive cases)	88 <sup>‡‡</sup>	13 <sup>‡‡</sup>	nr	nr	NPS (339), NPS+OPS (97), Trach Asp (30), OPS (15)	Automated Multiplex RT-PCR	N2, E, RdRp	Cepheid Xpert Xpress	N1, N3			99.5% (97.5–99.9)	95.8% (92.6–97.6)			
			129 <sup>‡‡</sup>	60 <sup>‡‡</sup>										100% (94.0–100)	100% (94.7–100)			
			99 <sup>‡‡</sup>	74 <sup>‡‡</sup>										100% (94.2–100)	92.0% (75.0–97.8)			
			65 <sup>‡‡</sup>	30 <sup>‡‡</sup>										100% (88.7–100)	74.3% (57.9–85.8)			
			79 <sup>‡‡</sup>	35 <sup>‡‡</sup>										100% (67.6–100)	100% (92.0–100)			
Bordi et al. <sup>40</sup>	Italy	Cohort + Controls <sup>††</sup>	278 + 20	99	nr	nr	Nasal swab, NPS (nr)	Multiplex RT-PCR	ORF1ab, S	Diasorin Simplexa	RdRp, E	100%	100%				0.938 (0.89–0.98)	
Rhoads et al. <sup>35,b</sup>	US	Cases only <sup>  </sup>	96 <sup>‡‡</sup>	96 <sup>‡‡</sup>	nr	nr	NPS (85), nasal swab (11)	Multiplex RT-PCR	ORF1ab, S	Diasorin Simplexa	N1, N2			96% (90–99)				

(continued on next page)

Table 5 (continued)

	Country	Study Type	No. Patients		Demographics <sup>†</sup>		Specimen	Index Test			Ref Stnd: rRT-PCR Primers	Study Findings (95% CI)							
			Total	Cases <sup>*</sup>	Age (y) <sup>‡</sup>	% M		Type	Primers	Platform		rSN	rSP	rPPV	rNPV	rPPA	rNPA	rOA	Cohen's $\kappa$
Poljak et al. <sup>41</sup>	Slovenia	Cohort <sup>††</sup>	501	63	nr	nr	NPS (489), NPS+OPS (12)	Automated Multiplex RT-PCR	ORF1, E	Roche cobas 6800	RdRp, E					100% (92.8–100)	99.5% (98.2–99.9)	99.6% (98.4–99.9)	0.98 (0.96–1.0)
Pujadas et al. <sup>42</sup>	USA	Cohort <sup>††</sup>	963 <sup>‡‡</sup>	640 <sup>‡‡</sup>	nr	nr	NPS (963)	Automated Multiplex RT-PCR	ORF1, E	Roche cobas 6800	N1, N2, N3					94.2% (92.2–95.9)	99.6% (98.1–99.9)	95.8% (94.4–97.0)	0.904 (0.87–0.93)
Rahman et al. <sup>43</sup>	Australia	Cohort <sup>††</sup>	52 <sup>‡‡</sup>	5 <sup>‡‡</sup>	31.5 (0–84)	58%	NPS+OPS (30), NPS (16), N Asp (5), sputum (1)	Multiplex RT-PCR	ORF1	Aus Diagnostics	RdRp, E	100%	92.16%	55.56%	100%				
Hogan et al. JCV, 4–24 <sup>44</sup>	USA	Cohort <sup>††</sup>	180 <sup>‡‡</sup>	77 <sup>‡‡</sup>	nr	nr	NPS (184)	Automated Multiplex RT-PCR	ORF1ab										
(2)	Hologic Panther	E					98.7% (93.0–100)	98.1% (93.1–99.8)	98.3% (95.2–99.7)	0.97 (0.93–1.0)									
Chen et al. <sup>45</sup>	Hong Kong	Cohort <sup>  </sup>	214	91	51 (IQR 31–69)	nr	NPS (214)	Multiplex RT-PCR	ORF1ab, E, N	Luminex NxTAG CoV	RdRp/Hel, E	97.8% (92.2–99.7)	100% (97.1–100)	100% (95.9–100)	98.4% (94.3–99.8)				0.98 (0.95–1.0)
Hogan et al. JCM <sup>46</sup>	USA	Case Control <sup>  </sup>	100	50	nr	nr	NPS (100)	Automated PCR with LFA	ORF1ab (2)	Mesa BioTech Accula	E					68.0% (53.3–80.5)	100% (92.9–100)	84.0% (75.3–90.6)	0.74 (0.61–0.87)
Visseaux et al. <sup>47</sup>	France	Case Control <sup>  </sup>	69	40	nr	nr	NPS (66), BAL(1), Trach Asp (2)	Automated Multiplex RT-PCR	ORF1, E	QIAstat-Dx	RdRp, E	100%	93%					97%	

\* Cases according to reference standard.

† Of cohort or cases.

‡ Format: median (range), median(IQR), or mean±SD. Patient Population.

|| Hospitalized patients.

§ Emergency Room/Immediate Care Center patients.

\*\* Outpatients.

†† not reported.

‡‡ Number of samples (when number of patients not reported).

§ Reported as concordance.

<sup>a</sup> Loeffelholz et al. and Moore et al. also appear in Table 6.

<sup>b</sup> Rhoads et al. appears twice in Table 5 for ease of comparison of studies of the same platform. Abbreviations- BAL: Bronchoalveolar lavage, CI: confidence interval, E: envelope, Hel: helicase, IQR: Interquartile range,  $\kappa$ : kappa statistic, LFA: lateral flow assay, M: male, n/a: not applicable, N: nucleocapsid, NAAT: nucleic acid amplification test, N Asp: nasopharyngeal aspirate, No.: number, NPS: nasopharyngeal swab, nr: not reported, NS: normal saline, OPS: oropharyngeal swab, ORF1ab: open reading frame 1ab, RdRp: RNA-dependent RNA polymerase, Ref Stnd: reference standard, rNPA: reported negative percent agreement, rNPV: reported negative predictive value, rOA: reported overall agreement, rPPA: reported positive percent agreement, rPPV: reported positive predictive value, rRT-PCR: real-time Reverse Transcription Polymerase Chain Reaction, rSN: reported sensitivity, rSP: reported specificity, S: spike, Trach Asp: Tracheal Aspirate, UTM: Universal transport medium, VTM: viral transport medium, y: years.

**Table 6**  
Studies assessing agreement between NAAT platforms.

Authors	Country	Study Type	No. Patients		Demographics†		Specimen	Platform #1		Platform #2		Study Findings (95% CI)								
			Total	Cases*	Age (y)‡	% M		Type	Primers	Platform	Type	Primers	Platform	rPPA	rNPA	rOA	rPPV	rNPV	Cohen's κ	
Harrington et al. <sup>48</sup>	USA	Cohort <sup>§</sup>	524	188	nr	nr	Paired NPS in VTM (RealTime) & foam nasal swab (ID NOW) (524 pairs)	Automated Multiplex RT-PCR	RdRp, N	Abbott RealTime	Automated Isothermal NAAT	RdRp	Abbott ID NOW	75% (67.7–80.6)	99% (97.6–99.8)					
Moore et al. <sup>34,a</sup>	USA	Cohort <sup>  ,§,**,††</sup>	200	125	50 ± 17	46%	NPS in VTM (200)	Automated Multiplex RT-PCR	RdRp, N	Abbott RealTime	Automated Isothermal NAAT	RdRp	Abbott ID NOW	75.2% (66.7–82.5)	100% (95.4–100)					
Basu et al. <sup>49</sup>	USA	Cohort <sup>§</sup>	101	32	(28 - 90)	nr	NPS dry (101)	Automated Multiplex RT-PCR	N2, E	Cepheid Xpert Xpress	Automated Isothermal NAAT	RdRp	Abbott ID NOW	54.8% (37.8–70.8)	98.6% (92.3–99.7)	85.1% (76.9–90.8)	94.4% (74.3–99)	83.1% (73–89.7)		
Hogan et al. JCV, 5–1 <sup>50</sup>	USA	Cases only <sup>§</sup> Case Control <sup>††</sup>	15 100 <sup>††</sup>	15 53 <sup>††</sup>	nr	nr	NPS in VTM (15)	Automated Multiplex RT-PCR	ORF1ab (2)	Hologic Panther	Automated Isothermal NAAT	RdRp	Abbott ID NOW	80.4% (66.9–90.2)	95.9% (86.0–99.5)					
Zhen, Smith et al. <sup>51,b</sup>	USA	Cohort <sup>††</sup>	108	58	nr	nr	NPS in VTM (108)	Automated Multiplex RT-PCR	ORF1ab (2)	Hologic Panther	Automated Isothermal NAAT	RdRp	Abbott ID NOW	87.7% (76–95)	100% (93–100)					0.87 (0.78–0.96)
Smithgall et al. <sup>52</sup>	USA	Case Control <sup>  §</sup>	113	90	65 (0–101)	60.2%	NPS in VTM or UTM (113)	Automated Multiplex RT-PCR	ORF1, E	Roche cobas 6800	Automated Isothermal NAAT	RdRp	Abbott ID NOW	73.9% (63.2–82.3)	100% (83.4–100)					
Moran et al. <sup>53</sup>	USA	Cohort <sup>  ,§,**,††</sup>	103 <sup>††</sup>	42 <sup>††</sup>	nr	nr	NPS (95), nasal swab (8)	Automated Multiplex RT-PCR	ORF1, E	Roche cobas 6800	Automated Multiplex RT-PCR	N2, E	Cepheid Xpert Xpress	98.9% (92.9–100)	92% (72.4–98.6)	99%				
Craney et al. <sup>54</sup>	USA	Cohort <sup>††</sup>	389	147	nr	nr	NPS (389)	Automated Multiplex RT-PCR	ORF1ab (2)	Hologic Panther	Automated Multiplex RT-PCR	ORF1, E	Roche cobas 6800			96.4%				0.922
Zhen, Smith et al. <sup>51,b</sup>	USA	Cohort <sup>††</sup>	108	58	nr	nr	NPS in VTM (108)	Automated Multiplex RT-PCR	ORF1ab (2)	Hologic Panther	Automated Multiplex RT-PCR	N2, E	Cepheid Xpert Xpress	98.3% (91–100)	100% (93–100)					0.98 (0.95–1.0)
								Automated Multiplex RT-PCR	ORF1ab (2)	Hologic Panther	Automated RT-PCR w/sensor	N	GenMark ePlex	91.4% (81–97)	100% (93–100)					0.91 (0.83–0.99)

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Table 6 (continued)

Authors	Country	Study Type	No. Patients		Demographics <sup>†</sup>		Specimen	Platform #1			Platform #2		Study Findings (95% CI)			rPPV	rNPV	Cohen's $\kappa$
			Total	Cases*	Age (y) <sup>‡</sup>	% M		Type	Primers	Platform	Type	Primers	Platform	rPPA	rNPA			
Loeffelholz et al <sup>39,a</sup>	USA, UK, FR, IT	Cohort <sup>††</sup> (enriched for positive cases)	18 <sup>‡‡</sup>	8 <sup>‡‡</sup>	nr	nr	NPS, OPS, NPS+OPS, Trach Asp (nr)	Automated Multiplex RT-PCR	RdRp, N	Abbott RealTime	Automated Multiplex RT-PCR	N2, E	Cepheid Xpert Xpress	100% (67.6–100)	100% (77.2–100)			
Norz et al. <sup>55</sup>	Germany	Case Control <sup>††</sup>	165 <sup>‡‡</sup>	36 <sup>‡‡</sup>	nr	nr	NPS, OPS (nr)	Automated Multiplex RT-PCR	ORF1, E	Roche cobas 6800	Automated RT-PCR	E	NeuMoDx 96	100%	99.2%			

\* Case estimated as a positive result of any evaluated platform.

<sup>†</sup> Of cohort or cases.

<sup>‡</sup> Format: median (range), median(IQR), or mean $\pm$ SD. Patient Population.

<sup>||</sup> Hospitalized patients.

<sup>§</sup> Emergency Room/Immediate Care Center patients.

\*\* Outpatient.

<sup>††</sup> not reported.

<sup>‡‡</sup> Number of samples (when number of patients not reported).

<sup>a</sup> Loeffelholz et al. and Moore et al. also appear in Table 5.

<sup>b</sup> Zhen, Smith et al. appears twice in Table 6 for ease of comparison of studies of the same platform.

<sup>c</sup> Dry or VTM status not reported. Rhoads et al. appears twice in Table 5 for ease of comparison of studies of the same platform. Abbreviations- CI: confidence interval, E: envelope, IQR: Interquartile range,  $\kappa$ : kappa statistic, M: male, N: nucleocapsid, NAAT: nucleic acid amplification test, No.: number, NPS: nasopharyngeal swab, nr: not reported, OPS: oropharyngeal swab, ORF1ab: open reading frame 1ab, RdRp: RNA-dependent RNA polymerase, rNPA: reported negative percent agreement, rNPV: reported negative predictive value, rOA: reported overall agreement, rPPA: reported positive percent agreement, rPPV: reported positive predictive value, rRT-PCR: real-time Reverse Transcription Polymerase Chain Reaction, S: spike, Trach Asp: Tracheal Aspirate, UTM: universal transport medium, VTM: viral transport medium, y: years.

(rAcc), positive percent agreement (rPPA), negative percent agreement (rNPA), overall agreement (rOA), and Kappa coefficient. Additionally, we extracted “positive rate,” a non-standard term used by the included studies to refer to the number of positive NAATs in a population of patients suspected to have COVID-19 (Table 1), or to the number of positive samples in a total population of positive samples after repeat testing (Table 2). We constructed 2 × 2 contingency tables and reproduced test performance characteristic calculations to demonstrate the methods of how the original authors obtained the values (Supplementary Table 1). We report additional pertinent study data in Supplementary Table 2: enrollment dates, number of sites of enrollment, symptomatic status, and chest radiology status. No articles were excluded on the basis of quality in order to present the most comprehensive summary of the currently available evidence.

### Data synthesis and analysis

We presented the extracted data in tabular form mirrored by a descriptive synthesis<sup>5</sup> in two broad categories: diagnostic accuracy studies for rRT-PCR (Tables 1–3), and diagnostic accuracy or comparative agreement studies of two NAATs (Tables 4–6). Tables are thematically divided based on the reference standard strategy, or approach to obtaining comparative agreement measures. Diagnostic accuracy studies for rRT-PCR were arranged alphabetically in tables by first author last name (Tables 1–3). Diagnostic accuracy and comparative agreement studies for two NAATs were arranged by decreasing order of studies per methodology, then alphabetically by methodology or platform (Tables 4–6) for easy comparison. Due to significant diversity in methods and reporting of results, we reported grouped summary data for study characteristics, patient characteristics, and outcomes.

We used the framework of the Quality Assessment of Diagnostic Accuracy Studies (QUADAS-2)<sup>6</sup> to evaluate our selected articles (Supplementary Table 3). We collected data, or noted their absence, for a narrative description of risk of bias and concerns of applicability based on the QUADAS domains. For assessment of bias in patient selection, we evaluated author conflicts of interest, study design type, inclusion/exclusion criteria, method of patient enrollment, and reporting of patient demographics and characteristics (i.e. symptomatic status). For assessment of bias in reference standard and index test, we evaluated the accuracy of the reference standard, the description of duration of symptoms at time of testing, whether the threshold to determine a positive test was pre-specified, and researcher blinding to reference standard and index test results. For assessment of bias in flow and timing, we evaluated whether the reference standard was the same for all patients, the sequence and timing of the performance of the reference standard and index test, whether test performance characteristics were calculated based on sample numbers or patient numbers, and whether indeterminate or invalid results were included in test performance calculations.

## Results

### Study selection

Our search yielded 1537 articles, with 816 unique articles after deduplication. After screening title and abstract, 130 articles underwent full text evaluation. Ultimately, 49 articles were included in our review (Fig. 1).

### The performance of rRT-PCR compared to case definitions or clinical diagnoses

Three studies, with 19 to 1014 patients, report a “positive rate” as the number of positive rRT-PCR out of the number of suspected cases of COVID-19, with a range of 38.42% to 59% (Table 1).<sup>7–9</sup> The studies do not report these values as “sensitivity” directly, however these values were interpreted as reflective of the accuracy of rRT-PCR. Ai et al. compared the positive rate of rRT-PCR (59%) with the positive rate of Chest CT in order to draw conclusions about the accuracy of Chest CT for the diagnosis of COVID-19,<sup>7</sup> and Liu et al. and Xie et al. expressed concern that their low calculated positive rates (38.42% and 47.4%, respectively) were indicative of a failure of rRT-PCR to diagnose COVID-19.<sup>8,9</sup>

In terms of quality assessment, the studies lack specific details as to how patients were classified as having suspected COVID-19 infection. The accuracy of clinical diagnosis based on case definitions is unclear but is likely not ideal for diagnosis. Additionally, duration of symptoms at the time of clinical diagnosis or rRT-PCR testing was not provided (Supplementary Table 3).

### The performance of rRT-PCR compared to end result after multiple repetitions of rRT-PCR

Eight studies, with a range of 36 to 22,061 patients per study, attempted to determine the accuracy of rRT-PCR by comparing the initial rRT-PCR result to the result after multiple repeated samples from the patient submitted for rRT-PCR testing, which was called the reference standard (Table 2).<sup>10–17</sup> Three studies reported this value as a “positive rate,” ranging from 51.25% to 88%,<sup>10,14,17</sup> and five reported sensitivity, with a range of 57.9% to 94.6%.<sup>11–13,15,16</sup> Of these studies, only He et al. included an rSp of 100%, calculated from patients who remained negative for SARS-CoV-2 after repeated sample testing (Supplementary Table 1).<sup>13</sup>

Green et al. included patients in their study regardless of whether they were tested once or multiple times, using data from these subsets of patients to make assumptions for estimating clinical test characteristics. In addition, this study also conducted multiple different NAATs and rRT-PCRs on patients, whereas other studies employing this strategy used only one type of NAAT. Additionally, the authors do not clarify whether patients who had repeat SARS-CoV-2 test were consistently tested with the same NAAT/rRT-PCR test or a different one. They also calculated test performance characteristics differently from other studies: two estimates of sensitivity were calculated, one in which the rate of false negatives for single-tested patients was 0%, and one in which the “false negative” rate was the same as in repeat-tested patients in their study of approximately 16.8%.<sup>12</sup> However, the details of how they calculated test characteristics were not presented. To clarify the two assumptions made in the calculations, we reconstructed the calculation in Supplementary Figure 1 which demonstrated the range of rSN with an estimate of the lower bound of 57.9% (55.2%–60.5%) and an estimate of the upper bound of 94.6% (94.2%–95.0%).

In terms of quality assessment, most of the studies were performed with non-cohort design, and six consisted of only patients who were determined to have COVID-19 by rRT-PCR, i.e. cases only (Table 2).<sup>10,11,14–16</sup> Five of the studies had inclusion criteria which caused pertinent patients to be excluded by necessitating patients to have had a well-performed CT Chest or X-Ray (Supplementary Table 3). This excluded several patients who would otherwise have been pertinent to the study of test diagnostic accuracy.<sup>10,11,13,16,17</sup> The studies involved repeating rRT-PCR several times for a reference standard, but each patient received a different number of repeat tests over a different time period, resulting in each patient receiving a different reference standard. One study tracked negative-

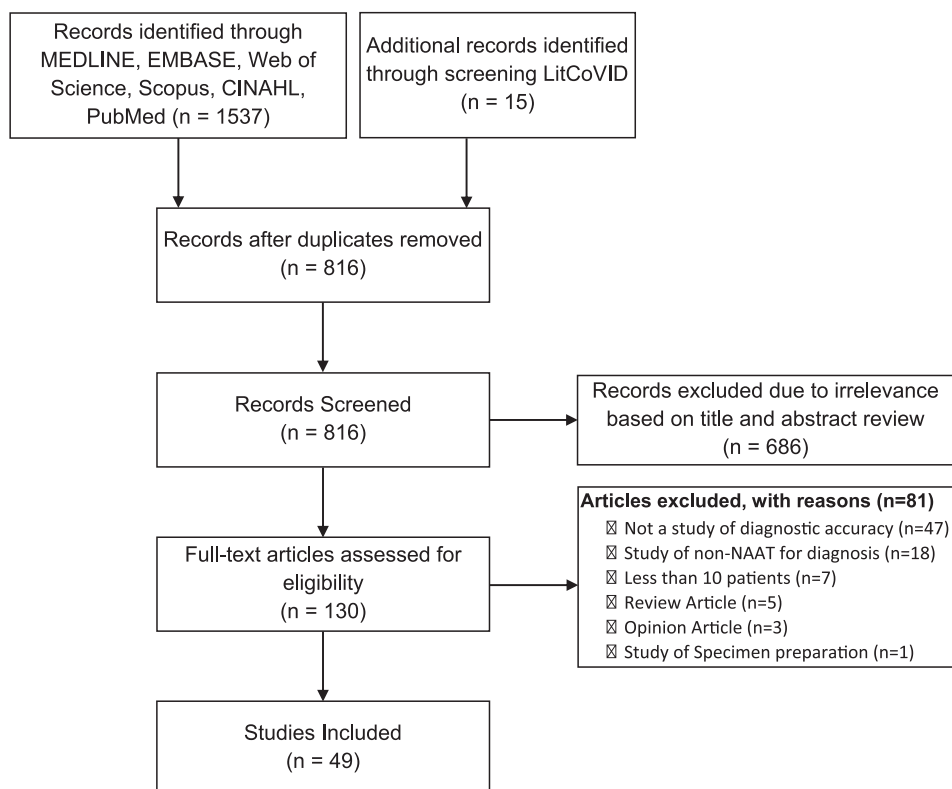


Fig. 1. PRISMA Flow diagram of studies included in the review.

to-positive conversion over 1 to 49 days,<sup>12</sup> and another tracked over 1 to 14 days,<sup>13</sup> leading to concern that potentially a patient could have been infected in the time between the initial test and the final test and confounding results. One study counted invalid results as negative results and indeterminate results as positive results when calculating test performance characteristics,<sup>12</sup> otherwise the rationale and ways invalid and indeterminate results were handled were not reported in these studies.

#### *The performance of rRT-PCR compared to various composite reference standards*

Three studies determined the accuracy or agreement of rRT-PCR or automated rRT-PCR platforms/instruments compared to a reference standard based on the results of several tests as a “composite reference standard” (Table 3).<sup>18–20</sup> There were between 58 and 184 patients per study. Suo et al. considered a positive result of either repeated measurements of rRT-PCR or serology to indicate a positive test according to the reference standard; reported sensitivity of initial rRT-PCR result was 40%, rSp 100%, rPPV 100%, and rNPV 16%.<sup>19</sup> Zhen et al. compared rRT-PCR performed according to the US CDC protocol to a composite reference standard in which the consensus result of 3 or more out of 4 molecular assays was considered the correct result. The rRT-PCR had an rPPA of 100%, an rNPA of 98%, and Cohen’s kappa coefficient of 0.98.<sup>20</sup> Cradic et al. did not study rRT-PCR but studied three automated molecular assays and used a composite reference standard of the consensus result of two or more of the three assays. While Abbott ID NOW had a rPPA of 91%, the Roche cobas 6800 and Diasorin Simplexa assays had a rPPA of 100%.<sup>18</sup>

These studies either did not report how samples were selected for evaluation (Supplementary Table 3),<sup>19,20</sup> or reported that only samples which had sufficient residual volume and had been properly stored were selected.<sup>18</sup> Cradic et al. and Zhen & Mangi et al.

had initially tested samples with one platform, and some or all samples were frozen, and subsequently thawed and tested with other platforms, leading to confounding factors in the reference standard and test performance calculations involving the various platform results.<sup>18,20</sup> Suo et al. used repeat rRT-PCR testing as part of the reference standard, with repeat tests performed 2–10 days after the initial test, after the patient had been discharged from the hospital, leading to potential exposure for initial infection or reinfection.<sup>19</sup>

#### *The performance other nucleic acid amplification test methods compared to standard rRT-PCR*

Fourteen studies compared other nucleic acid amplification test methods to detect SARS-CoV-2 to rRT-PCR (Table 4), with between 27 and 356 patients per study. Five of the studies evaluated reverse transcription loop mediated isothermal amplification (RT-LAMP),<sup>21–25</sup> four reported sensitivity of 100% and specificity of 95.6 to 100%,<sup>21–23,25</sup> and one study reported accuracy of 92.9%.<sup>24</sup> Two studies, Wang, Cai, & He et al. and Xue et al., evaluated reverse-transcription recombinase-aided amplification (RT-RAA) with Cohen’s kappa of 0.952 and 1.0.<sup>26,27</sup> Two studies, Perchetti et al. and Waggoner et al., evaluated triplex rRT-PCR, reporting overall agreement as 99.2 and 100%.<sup>28,29</sup> Li et al. evaluated an automatic integrated gene detection system (AIGS) with rSN of 97.2% and rSp 98.5%.<sup>30</sup> Suo et al. evaluated digital droplet polymerase chain reaction (ddPCR), with rSN 94%, rSp 100%, rPPV 100%, rNPV of 63%, and rAcc 95%.<sup>19</sup> Bulterys et al. study evaluated an isothermal amplification method with rSN 82.8% and Cohen’s kappa 0.86.<sup>31</sup> Wang, Cai, and Zhang et al. evaluated one-step single-tube nested quantitative polymerase chain reaction (OSN-qRT-PCR) with Cohen’s kappa of 0.737.<sup>32</sup>

Regarding evaluation of quality (Supplementary Table 3), the majority of studies did not report how patient samples were



selected for evaluation.<sup>21,22,24,25,27–30,32</sup> In the study conducted by Bulterys et al., sample selection was a convenience selection of samples with residual volume that had been stored correctly.<sup>31</sup> Most studies did not report symptomatic status of the patient<sup>21–28,30–32</sup> or patient demographics.<sup>21–25,27–32</sup> Problematically, many of the studies did not report when the reference standard was conducted on the patient samples compared to the index test, or whether actions that could potentially alter test results (such as freeze/thaw cycles) occurred between reference standard or index test.<sup>21–24,27,28,31</sup> Four studies calculated test performance characteristics based on number of samples rather than number of patients.<sup>23,27,30,32</sup> The management of indeterminate and invalid test results went largely unreported.<sup>21–25,27,30</sup>

#### *The performance of NAAT platforms compared to rRT-PCR as the reference standard*

Fifteen studies compared automated NAAT platforms to various rRT-PCR assays to determine test performance characteristics (Table 5), with between 26 and 963 patients or samples per study. Three studies evaluated Abbott ID NOW, an isothermal NAAT platform, with rPPA or rSN of 71.7% to 94%, and rNPA or rSP of 100%.<sup>33–35</sup> Two studies evaluated Abbott RealTime with rSN or rPPA of 93% to 100%, and rSP or rNPA of 92.4% to 100%.<sup>34,36</sup> Three studies evaluated Cepheid Xpert Xpress, with rPPA 96.1% to 100%, rNPA 74.3% to 100%, rOA 96.1% to 100%, and Cohen's Kappa of 0.92.<sup>37–39</sup> Two studies evaluated Diasorin Simplexa with rSN or rPPA of 96% to 100%, and rSN of 100%.<sup>35,40</sup> Two studies evaluated Roche cobas 6800 with rPPA 94.2% to 100%, rNPA 99.5% to 99.6%, and Cohen's Kappa of 0.904 to 0.98.<sup>41,42</sup> Other studies evaluated AusDiagnostics (rSN 100%, rSP 92.16%),<sup>43</sup> Hologic Panther Fusion (rPPA 98.7%, rNPA 98.1%),<sup>44</sup> Luminex NxTAG (rSN 97.8%, rSP 100%),<sup>45</sup> Mesa BioTech Accula (rPPA 68.0%, rNPA 100%),<sup>46</sup> or QIAstat-Dx (rSN 100%, rSP 93%)<sup>47</sup> compared to rRT-PCR.

With regards to quality evaluation (Supplementary Table 3), most studies did not report method of sample collection/patient recruitment,<sup>33,35,37,41–47</sup> and four studies conducted a convenience selection of samples, including enrichment for positive samples.<sup>34–36,39</sup> Eight studies conducted test performance calculations on sample numbers instead of patient numbers.<sup>33,35,36,38,39,42–44</sup> Four studies conducted calculation of test performance characteristics with indeterminate or inconclusive results as “positive,”<sup>35,38,39,42</sup> and the management of indeterminate/inconclusive as well as invalid results went unreported in an additional three studies.<sup>33,37,47</sup> No study reported the blinding of researchers to the reference standard or index test results.

#### *The agreement of NAAT platforms compared to other NAAT platforms*

Ten studies, containing between 15 and 524 patients per study, evaluated the agreement between two different types of NAAT platforms (Table 6), typically under the circumstances where one platform was the standard of care at the institution, and another was introduced. The Abbott ID NOW platform, using isothermal amplification, was the most frequently studied test, with an rPPA of 75–75.2% compared to Abbott Real Time,<sup>34,48</sup> 54.8% compared to Cepheid Xpert Xpress,<sup>49</sup> 80.4–87.7% compared to Hologic Panther Fusion,<sup>50,51</sup> and 73.9% compared to Roche cobas 6800.<sup>52</sup> Two studies evaluated Cepheid Xpert Xpress compared to Roche cobas 6800, with rPPA 98.9% and rNPA 92% in one,<sup>52</sup> and overall agreement of 99% in another.<sup>53</sup> Several platforms were compared to Hologic Panther Fusion, including Roche cobas 6800 with rOA 96.4%,<sup>54</sup> Cepheid Xpert Xpress with rPPA 98.3% and rNPA 100%, and GenMark ePLEX with rPPA 91.4% and rNPA 100%.<sup>51</sup>

In the studies, some platforms were identified as the “comparator” or “reference” platforms, including Cepheid Xpert Xpress,<sup>49</sup>

Abbott RealTime,<sup>34,48</sup> Hologic Panther Fusion,<sup>50,51</sup> and Roche cobas 6800,<sup>52,55</sup> and these were listed as “Platform #1” in Table 6. Three studies did not identify any studied platform as the “comparator” or “reference standard,” and instead only reported general, non-directional measures of agreement such as overall agreement, Cohen's Kappa, or alternatively, the calculations of PPA and NPA were identical no matter their method of calculation (Supplementary Table 1).<sup>39,53,54</sup>

Regarding quality evaluation (Supplementary Table 3), the samples used for calculating test performance characteristics were reported to be selected for enrichment of positive samples,<sup>34,39</sup> for diversity of viral load,<sup>52,54</sup> otherwise curated,<sup>50</sup> or the method of selecting samples was unreported.<sup>49,51,53,55</sup> Symptomatic status of the patients was largely unreported.<sup>39,49–55</sup> Five studies included samples where one test was conducted, then interim freezing, cooling, or other storage, before performance of the second test.<sup>39,50–52,54</sup> Two studies did not report the sequence of testing of the two platforms or interim handling or storage of the samples.<sup>49,53</sup> The status of researcher blinding to either platform result was not reported in any study.

## **Discussion**

In our scoping review of 49 articles concerning test performance characteristics of rRT-PCR and other NAAT used for the diagnosis of COVID-19, we were able to observe several overarching themes. Clinical diagnosis by the case definition for COVID-19 used in the early period of the pandemic does not correlate well with positive rates of COVID-19 rRT-PCR (Table 1). The result of the initial rRT-PCR performed on a patient, if negative, may not be reflective of the result after multiple repeated rRT-PCRs for that patient (Table 2). Several alternative NAAT methods, many of which are easier or faster to perform, may be comparable to standard rRT-PCR (Table 4). Proprietary multiplex, automated, and/or point-of-care methods are comparable to in accuracy to rRT-PCR (Table 5) and to each other (Table 6), although the Abbott ID NOW SARS-CoV-2 test appears to have lower comparative agreement to other platforms.<sup>34,48–52</sup>

These findings should be viewed cautiously as the SARS-CoV-2 tests in these studies have not undergone rigorous evaluation necessary for FDA approval due to the emergency state generated by the COVID-19 pandemic. In addition, during our scoping review, we found substantial heterogeneity among available studies in terms of test types, reference standards, metrics, and details of study design and methodology.

We categorized the included studies by four different reference standard strategies: clinical diagnosis/case definitions (Table 1), repeated index testing (Table 2), composite reference standard (Table 3), and rRT-PCR (Table 4 and 5). Additionally, we identified a fifth category, where instead of using a reference standard, comparative agreement between two NAAT platforms was calculated (Table 5 and 6).

The main limitation of the first group of studies (Table 1) was the use of a “case definition” as the reference standard to report a “positive rate” of rRT-PCR. During novel disease outbreaks, standard case definitions are often developed to assist clinicians in case identification before a diagnostic test is available. Unfortunately, the studies included in this group were unable to use a clear case definition; instead they refer to a population of “suspected cases,” for which the definition is not reported.<sup>7–9</sup> Because this group enrolled patients prior to February 15, 2020 in China, during the time in which the Chinese National Guideline for Diagnosis and Treatment of COVID-19 (NGDTC) published five different versions of the COVID-19 case definition, the case definitions in use at the time of these studies varied.<sup>56</sup> A recent study estimated that if a single guideline (specifically, version 5 of the NGDTC) had been used to

identify cases from the beginning of the outbreak to February 20, 2020, there would have been more than three times as many identified cases in Hubei province.<sup>56</sup> This is relevant to our review because the two largest studies that evaluated the rRT-PCR positive rate of patients with a clinical diagnosis of COVID-19 took place in Wuhan, Hubei province, and included patients evaluated before February 14, 2020<sup>7,8</sup> (Supplementary Table 2). This increased case estimate due to diagnosis of COVID-19 based on case definition complicates the legitimacy and reported accuracy of the “positive rate” of rRT-PCR referred to in these studies.

The second group assessed rRT-PCR test performance characteristics via repeated index rRT-PCR testing (Table 2). Most studies in this group reported “sensitivity” by dividing the number of participants with positive baseline rRT-PCRs by the total number of participants who eventually had a positive rRT-PCR after repeated measurements. While such an approach may have some advantages over the use of a case definition alone as a reference standard, this strategy is, nonetheless, an imperfect solution with its own set of inherent limitations. SARS-CoV-2 infection is transient and the associated viral loads are time-varying because of the natural pathophysiology of the infection. Therefore, the time interval between each repeated test becomes crucially important, and even relatively small time differences (and/or lack of uniformly used intervals) could complicate the interpretation of re-test results and their quality as reference standards. Furthermore, repeated use of the same test as a reference standard for itself does not eliminate the inaccuracies or limitations of the test. Such comparisons ultimately reflect the reliability of the test (assuming a short, uniform time interval between tests), rather than providing a true view of test accuracy.

The third group of three studies calculated test performance characteristics of rRT-PCR according to a composite reference standard (Table 3). Using arbitrary rules to combine multiple different and imperfect tests inevitably creates a reference standard with some degree of bias.<sup>57</sup> Furthermore, all three studies in this group included the test under evaluation as part of the composite reference standard, which leads to additional bias, described below.<sup>58</sup> Use of a biased composite standard is likely to lead to reduced sensitivity, among other errors affecting true test performance characteristics.<sup>59</sup>

The fourth group of studies evaluated SARS-CoV-2 diagnostic tests that are under development as well as proprietary testing platforms (most of which are based on standard rRT-PCR methods). These studies used traditional rRT-PCR as a reference standard; results are summarized in Tables 4 and 5, respectively. Importantly, while these studies were not designed to estimate the accuracy of rRT-PCR, their results indicate that the index tests did not identify significantly more positive samples than rRT-PCR.

Finally, the last group of studies compared SARS-CoV-2 NAAT platforms (Table 6). These comparative accuracy studies examined the agreement between two non-reference standard tests. Although most of the testing platforms evaluated in these studies were based on standard rRT-PCR, the agreement between two non-reference standard tests is not equivalent to test accuracy, as mentioned previously.

This scoping review is limited by the lack of reporting of several key study features in the majority of the articles evaluated, which is an important indicator of quality and potential bias. Based on the QUADAS-2 criteria, most of the included studies had concern for bias (Supplementary Table 3). The most prominent concerns were unclear inclusion/exclusion criteria, unclear method of enrollment/selection of patients and samples, and unclear handling of indeterminate/inconclusive and invalid results. Additionally, many of the studies were conducted in a so-called “two gate” (case-control) design, in which cases and controls were known and selected ahead of time, rather than performing the test on a

group of patients or samples with suspected COVID-19. These factors likely incorporate bias that significantly confounds the results of the studies, thus, the accuracy of the tests in other settings with different prevalences (such as asymptomatic screening, other age groups) may not be truly generalizable. Furthermore, few studies were able to evaluate both the index and reference tests simultaneously or within a short period of time, which is key to avoiding biases caused by changes in the patient’s true disease status; this bias can also affect the diagnostic accuracy of the index test.

The best approach to determining diagnostic test performance characteristics in the absence of a “gold” standard is an open question in diagnostic accuracy methodology. While many methods have been described, there are only a few well-defined statistical approaches that use a reference standard in lieu of a gold standard, reviewed elsewhere.<sup>60</sup> Latent class analysis is one commonly used approach in situations in which neither the true error rates of the reference standard nor the true prevalence of the disease are known. This approach uses the results of a set of imperfect tests to estimate parameters related to sensitivity, specificity, and prevalence often using maximum likelihood methods. However, this is not the only method available and every method has its own strengths and limitations.<sup>57</sup> Therefore, careful interpretation by studies that attempt to estimate test characteristics is warranted to account for and clarify the inherent limitations of assessing accuracy-related metrics when a gold standard is unavailable.

Evaluation of the performance characteristics of SARS-CoV-2 diagnostic tests is vital to control of the ongoing COVID-19 pandemic. While more than 200 SARS-CoV-2 molecular diagnostic tests have received FDA EUAs, we have described in this scoping review that the performance of few of these tests has been assessed appropriately. The lack of robust test performance that we noted in many studies published to date is undoubtedly due in part to the critical need for tests, which resulted in accelerated test development. However, our scoping review also uncovered imperfect methods for estimating diagnostic test performance in the absence of a gold standard and demonstrate that the accuracy of these tests should be interpreted with caution. Future studies would benefit from employing statistical methods such as latent class analysis and other methods referenced above to accurately analyze their data. Indeed, instituting national requirements for test performance analysis and reporting, perhaps based on the existing FDA guidelines on diagnostic tests,<sup>61</sup> would advance the goal of standardizing the evaluation SARS-CoV-2 diagnostic test performance. Such an initiative would lead to statistically robust conclusions regarding the accuracy of the index test, which will in turn support hospitals and clinicians as they determine the optimal test to use for COVID-19 diagnosis.

#### Declaration of Competing Interest

None

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#### Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:[10.1016/j.jinf.2020.08.043](https://doi.org/10.1016/j.jinf.2020.08.043).



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