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Rapid, point-of-care antigen and molecular-based tests for diagnosis of SARS-CoV-2 infection (Review)

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TABLE OF CONTENTS

HEADER	1
ABSTRACT	1
PLAIN LANGUAGE SUMMARY	3
SUMMARY OF FINDINGS	5
BACKGROUND	10
OBJECTIVES	13
METHODS	13
RESULTS	16
Figure 1	17
Figure 2	21
Figure 3	23
Figure 4	24
Figure 5	25
Figure 6	26
Figure 7	27
Figure 8	27
Figure 9	28
Figure 10	29
DISCUSSION	36
AUTHORS' CONCLUSIONS	40
ACKNOWLEDGEMENTS	41
REFERENCES	43
CHARACTERISTICS OF STUDIES	59
ADDITIONAL TABLES	313
WHAT'S NEW	323
HISTORY	323
CONTRIBUTIONS OF AUTHORS	323
DECLARATIONS OF INTEREST	323
SOURCES OF SUPPORT	324
DIFFERENCES BETWEEN PROTOCOL AND REVIEW	324
INDEX TERMS	325

i



[Diagnostic Test Accuracy Review]

Rapid, point-of-care antigen and molecular-based tests for diagnosis of SARS-CoV-2 infection

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ABSTRACT

Background

Accurate rapid diagnostic tests for SARS-CoV-2 infection could contribute to clinical and public health strategies to manage the COVID-19 pandemic. Point-of-care antigen and molecular tests to detect current infection could increase access to testing and early confirmation of cases, and expediate clinical and public health management decisions that may reduce transmission.

Objectives

To assess the diagnostic accuracy of point-of-care antigen and molecular-based tests for diagnosis of SARS-CoV-2 infection. We consider accuracy separately in symptomatic and asymptomatic population groups.

Search methods

Electronic searches of the Cochrane COVID-19 Study Register and the COVID-19 Living Evidence Database from the University of Bern (which includes daily updates from PubMed and Embase and preprints from medRxiv and bioRxiv) were undertaken on 30 Sept 2020. We checked



repositories of COVID-19 publications and included independent evaluations from national reference laboratories, the Foundation for Innovative New Diagnostics and the Diagnostics Global Health website to 16 Nov 2020. We did not apply language restrictions.

Selection criteria

We included studies of people with either suspected SARS-CoV-2 infection, known SARS-CoV-2 infection or known absence of infection, or those who were being screened for infection. We included test accuracy studies of any design that evaluated commercially produced, rapid antigen or molecular tests suitable for a point-of-care setting (minimal equipment, sample preparation, and biosafety requirements, with results within two hours of sample collection). We included all reference standards that define the presence or absence of SARS-CoV-2 (including reverse transcription polymerase chain reaction (RT-PCR) tests and established diagnostic criteria).

Data collection and analysis

Studies were screened independently in duplicate with disagreements resolved by discussion with a third author. Study characteristics were extracted by one author and checked by a second; extraction of study results and assessments of risk of bias and applicability (made using the QUADAS-2 tool) were undertaken independently in duplicate. We present sensitivity and specificity with 95% confidence intervals (CIs) for each test and pooled data using the bivariate model separately for antigen and molecular-based tests. We tabulated results by test manufacturer and compliance with manufacturer instructions for use and according to symptom status.

Main results

Seventy-eight study cohorts were included (described in 64 study reports, including 20 pre-prints), reporting results for 24,087 samples (7,415 with confirmed SARS-CoV-2). Studies were mainly from Europe (n = 39) or North America (n = 20), and evaluated 16 antigen and five molecular assays.

We considered risk of bias to be high in 29 (37%) studies because of participant selection; in 66 (85%) because of weaknesses in the reference standard for absence of infection; and in 29 (37%) for participant flow and timing. Studies of antigen tests were of a higher methodological quality compared to studies of molecular tests, particularly regarding the risk of bias for participant selection and the index test. Characteristics of participants in 35 (45%) studies differed from those in whom the test was intended to be used and the delivery of the index test in 39 (50%) studies differed from the way in which the test was intended to be used. Nearly all studies (97%) defined the presence or absence of SARS-CoV-2 based on a single RT-PCR result, and none included participants meeting case definitions for probable COVID-19.

Antigen tests

Forty-eight studies reported 58 evaluations of antigen tests. Estimates of sensitivity varied considerably between studies. There were differences between symptomatic (72.0%, 95% CI 63.7% to 79.0%; 37 evaluations; 15530 samples, 4410 cases) and asymptomatic participants (58.1%, 95% CI 40.2% to 74.1%; 12 evaluations; 1581 samples, 295 cases). Average sensitivity was higher in the first week after symptom onset (78.3%, 95% CI 71.1% to 84.1%; 26 evaluations; 5769 samples, 2320 cases) than in the second week of symptoms (51.0%, 95% CI 40.8% to 61.0%; 22 evaluations; 935 samples, 692 cases). Sensitivity was high in those with cycle threshold (Ct) values on PCR ≤25 (94.5%, 95% CI 91.0% to 96.7%; 36 evaluations; 2613 cases) compared to those with Ct values >25 (40.7%, 95% CI 31.8% to 50.3%; 36 evaluations; 2632 cases). Sensitivity varied between brands. Using data from instructions for use (IFU) compliant evaluations in symptomatic participants, summary sensitivities ranged from 34.1% (95% CI 29.7% to 38.8%; Coris Bioconcept) to 88.1% (95% CI 84.2% to 91.1%; SD Biosensor STANDARD Q). Average specificities were high in symptomatic and asymptomatic participants, and for most brands (overall summary specificity 99.6%, 95% CI 99.0% to 99.8%).

At 5% prevalence using data for the most sensitive assays in symptomatic people (SD Biosensor STANDARD Q and Abbott Panbio), positive predictive values (PPVs) of 84% to 90% mean that between 1 in 10 and 1 in 6 positive results will be a false positive, and between 1 in 4 and 1 in 8 cases will be missed. At 0.5% prevalence applying the same tests in asymptomatic people would result in PPVs of 11% to 28% meaning that between 7 in 10 and 9 in 10 positive results will be false positives, and between 1 in 2 and 1 in 3 cases will be missed.

No studies assessed the accuracy of repeated lateral flow testing or self-testing.

Rapid molecular assays

Thirty studies reported 33 evaluations of five different rapid molecular tests. Sensitivities varied according to test brand. Most of the data relate to the ID NOW and Xpert Xpress assays. Using data from evaluations following the manufacturer's instructions for use, the average sensitivity of ID NOW was 73.0% (95% CI 66.8% to 78.4%) and average specificity 99.7% (95% CI 98.7% to 99.9%; 4 evaluations; 812 samples, 222 cases). For Xpert Xpress, the average sensitivity was 100% (95% CI 88.1% to 100%) and average specificity 97.2% (95% CI 89.4% to 99.3%; 2 evaluations; 100 samples, 29 cases). Insufficient data were available to investigate the effect of symptom status or time after symptom onset.

Authors' conclusions

Antigen tests vary in sensitivity. In people with signs and symptoms of COVID-19, sensitivities are highest in the first week of illness when viral loads are higher. The assays shown to meet appropriate criteria, such as WHO's priority target product profiles for COVID-19 diagnostics ('acceptable' sensitivity \geq 80% and specificity \geq 97%), can be considered as a replacement for laboratory-based RT-PCR when immediate



decisions about patient care must be made, or where RT-PCR cannot be delivered in a timely manner. Positive predictive values suggest that confirmatory testing of those with positive results may be considered in low prevalence settings. Due to the variable sensitivity of antigen tests, people who test negative may still be infected.

Evidence for testing in asymptomatic cohorts was limited. Test accuracy studies cannot adequately assess the ability of antigen tests to differentiate those who are infectious and require isolation from those who pose no risk, as there is no reference standard for infectiousness. A small number of molecular tests showed high accuracy and may be suitable alternatives to RT-PCR. However, further evaluations of the tests in settings as they are intended to be used are required to fully establish performance in practice.

Several important studies in asymptomatic individuals have been reported since the close of our search and will be incorporated at the next update of this review. Comparative studies of antigen tests in their intended use settings and according to test operator (including self-testing) are required.

PLAIN LANGUAGE SUMMARY

How accurate are rapid tests for diagnosing COVID-19?

What are rapid point-of-care tests for COVID-19?

Rapid point-of-care tests aim to confirm or rule out COVID-19 infection in people with or without COVID-19 symptoms. They:

- are portable, so they can be used wherever the patient is (at the point of care);
- are easy to perform, with a minimum amount of extra equipment or complicated preparation steps;
- are less expensive than standard laboratory tests;
- do not require a specialist operator or setting; and
- provide results 'while you wait'.

We were interested in two types of commercially available, rapid point-of-care tests: antigen and molecular tests. Antigen tests identify proteins on the virus; they come in disposable plastic cassettes, similar to pregnancy tests. Rapid molecular tests detect the virus's genetic material in a similar way to laboratory methods, but using smaller devices that are easy to transport or to set up outside of a specialist laboratory. Both test nose or throat samples.

Why is this question important?

People with suspected COVID-19 need to know quickly whether they are infected, so that they can self-isolate, receive treatment, and inform close contacts. Currently, COVID-19 infection is confirmed by a laboratory test called RT-PCR, which uses specialist equipment and often takes at least 24 hours to produce a result.

Rapid point-of-care tests could open access to testing for many more people, with and without symptoms, potentially in locations other than healthcare settings. If they are accurate, faster diagnosis could allow people to take appropriate action more quickly, with the potential to reduce the spread of COVID-19.

What did we want to find out?

We wanted to know whether commercially available, rapid point-of-care antigen and molecular tests are accurate enough to diagnose COVID-19 infection reliably, and to find out if accuracy differs in people with and without symptoms.

What did we do?

We looked for studies that measured the accuracy of any commercially produced, rapid antigen or molecular point-of-care test, in people tested for COVID-19 using RT-PCR. People could be tested in hospital or the community. Studies could test people with or without symptoms.

Tests had to use minimal equipment, be performed safely without risking infection from the sample, and have results available within two hours of the sample being collected.

What we found

We included 64 studies in the review. They investigated a total of 24,087 nose or throat samples; COVID-19 was confirmed in 7415 of these samples. Studies investigated 16 different antigen tests and five different molecular tests. They took place mainly in Europe and North America.



Main results

Antigen tests

In people with confirmed COVID-19, antigen tests correctly identified COVID-19 infection in an average of 72% of people with symptoms, compared to 58% of people without symptoms. Tests were most accurate when used in the first week after symptoms first developed (an average of 78% of confirmed cases had positive antigen tests). This is likely to be because people have the most virus in their system in the first days after they are infected.

In people who did not have COVID-19, antigen tests correctly ruled out infection in 99.5% of people with symptoms and 98.9% of people without symptoms.

Different brands of tests varied in accuracy. Pooled results for one test (SD Biosensor STANDARD Q) met World Health Organization (WHO) standards as 'acceptable' for confirming and ruling out COVID-19 in people with signs and symptoms of COVID-19. Two more tests met the WHO acceptable standards (Abbott Panbio and BIONOTE NowCheck) in at least one study.

Using summary results for SD Biosensor STANDARD Q, if 1000 people with symptoms had the antigen test, and 50 (5%) of them really had COVID-19:

- 53 people would test positive for COVID-19. Of these, 9 people (17%) would not have COVID-19 (false positive result).
- 947 people would test negative for COVID-19. Of these, 6 people (0.6%) would actually have COVID-19 (false negative result).

In people with no symptoms of COVID-19 the number of confirmed cases is expected to be much lower than in people with symptoms. Using summary results for SD Biosensor STANDARD Q in a bigger population of 10,000 people with no symptoms, where 50 (0.5%) of them really had COVID-19:

- 125 people would test positive for COVID-19. Of these, 90 people (72%) would not have COVID-19 (false positive result).
- 9,875 people would test negative for COVID-19. Of these, 15 people (0.2%) would actually have COVID-19 (false negative result).

Molecular tests

Although overall results for diagnosing and ruling out COVID-19 were good (95.1% of infections correctly diagnosed and 99% correctly ruled out), 69% of the studies used the tests in laboratories instead of at the point-of-care and few studies followed test manufacturer instructions. Most of the data relate to the ID NOW and Xpert Xpress tests. We noted a large difference in COVID-19 detection between the two tests, but we cannot be certain about whether results will remain the same in a real world setting. We could not investigate differences in people with or without symptoms, nor time from when symptoms first showed because the studies did not provide enough information about their participants.

How reliable were the results of the studies?

In general, studies that assessed antigen tests used more rigorous methods than those that assessed molecular tests, particularly when selecting participants and performing the tests. Sometimes studies did not perform the test on the people for whom it was intended and did not follow the manufacturers' instructions for using the test. Sometimes the tests were not carried out at the point-of-care. Nearly all the studies (97%) relied on a single negative RT-PCR result as evidence of no COVID-19 infection. Results from different test brands varied, and few studies directly compared one test brand with another. Finally, not all studies gave enough information about their participants for us to judge how long they had had symptoms, or even whether or not they had symptoms.

What does this mean?

Some antigen tests are accurate enough to replace RT-PCR when used in people with symptoms. This would be most useful when quick decisions are needed about patient care, or if RT-PCR is not available. Antigen tests may be most useful to identify outbreaks, or to select people with symptoms for further testing with PCR, allowing self-isolation or contact tracing and reducing the burden on laboratory services. People who receive a negative antigen test result may still be infected.

Several point-of-care molecular tests show very high accuracy and potential for use, but more evidence of their performance when evaluated in real life settings is required.

We need more evidence on rapid testing in people without symptoms, on the accuracy of repeated testing, testing in non-healthcare settings such as schools (including self-testing), and direct comparisons of test brands, with testers following manufacturers' instructions.

How up-to-date is this review?

This review updates our previous review and includes evidence published up to 30 September 2020.

SUMMARY OF FINDINGS

Summary of findings 1. Diagnostic accuracy of point-of-care antigen and molecular-based tests for the diagnosis of SARS-CoV-2 infection

Question	What is the diagnostic accuracy of rapid point-of-care antigen and molecular-based tests for Adults or children with suspected: - current SARS-CoV-2 infection or populations undergoing screening for SARS-CoV-2 infection, including - asymptomatic contacts of confirmed COVID-19 cases - community screening
Index test	Any rapid antigen or molecular-based test for diagnosis of SARS-CoV-2 meeting the following criteria: portable or mains-powered device minimal sample preparation requirements minimal biosafety requirements no requirement for a temperature-controlled environment test results available within 2 hours of sample collection
Target condition Reference standard	Detection of current SARS-CoV-2 infection For COVID-19 cases: positive RT-PCR alone or clinical diagnosis of COVID-19 based on established guidelines or combinations of clinical features For non-COVID-19 cases: negative RT-PCR or pre-pandemic sources of samples
Action	False negative results mean missed cases of COVID-19 infection, with either delayed or no confirmed diagnosis and increased risk of community transmission due to false sense of security False positive results lead to unnecessary self-isolation or quarantine, with the potential for new infection to be acquired
Quantity of evidence	e Number studies
	Respiratory 77 24,418 Non-respira- 1 79 tory
Limitations in the evidence	vidence
Risk of bias	Participants: high (29) or unclear (27) risk in 56 studies (72%)

100 (99.0 to 100)	34.1 (29.7 to 38.8)	414	780	ω	Coris Bioconcept - COVID-19 Ag Respi-Strip
				icipants	Symptomatic participants
		cases			
	Sensitivity (95% CI)	SARS-CoV-2	Samples	Evaluations	Tests
uctions for u	Examples of pooled results for individual antigen tests using data for evaluations compliant with manufacturer instructions for use according to sympt	sts using data for eva	ividual antigen te	ed results for indi	Examples of poole
	[29% to 85%]				
	58.1 (40.2 to 74.1)		1581 (295)	12 (10)	Asymptomatic
	[15% to 95%]				onset of symp- toms) ^a
	78.3 (71.1 to 84.1)		2320 (2320)	26 (21)	Symptomatic (up
	[0% to 100%]				
	72.0 (63.7 to 79.0)		15,530 (4410)	37 (27)	Symptomatic
	[Range]			(studies)	
	Sensitivity (95% CI)	·CoV-2 cases)	Samples (SARS-CoV-2 cases)	Evaluations	
				ests	Findings: antigen tests
))	Reference standard: high concerns in 76 studies (97%)	ndard: high concer	Reference star	
	Index test (molecular tests): high concerns in 16 studies (53% of 30 studies)	າ concerns in 16 studi	lecular tests): hig	Index test (mo	studies)
	(48% of 48 studies)	Index test (antigen tests): high concerns in 23 studies (48% of 48 studies)	igen tests): high c	Index test (ant	(hasad on 78
		studies (45%)	Participants: high concerns in 35 studies (45%)	Participants: h	Concerns about
	ıdies (83%)	Flow and timing: high (29) or unclear (36) risk in 65 studies (83%)	ng: high (29) or unc	Flow and timin	
	ıdies (92%)	Reference standard: high (66) unclear (6) risk in 72 studies (92%)	ndard: high (66) un	Reference star	
	Index test (molecular tests): high (3) or unclear (22) risk in 25 studies (83% of 30 studies)	າ (3) or unclear (22) ris	lecular tests): hig	Index test (mo	ordaico/
	Index test (antigen tests): high (0) or unclear (19) risk in 19 studies (40% of 48 studies))) or unclear (19) risk i	i gen tests): high ((Index test (ant	(based on 78

tom status

Rapid, point-of-care antigen and molecular-based tests for diagnosis of SARS-CoV-2 infection (Review)

Trusted evidence. Informed decisions. Better health.

2.7% 5.9% 0.6% 1.3% 2.9%	94% 97% 84% 92% 96%	896 (888 to 898) 796 (790 to 798) 941 (929 to 946) 892 (880 to 896) 793 (782 to 797)	25 (13 to 43) 50 (26 to 85) 6 (4 to 8) 12 (9 to 16) 24 (18 to 32)	5 (2 to 12) 4 (2 to 10) 9 (4 to 21) 8 (4 to 20) 7 (3 to 18)	75 (57 to 87) 150 (115 to 174) 44 (42 to 46) 88 (84 to 91) 176 (168 to 182)	10% 20% 5% 10% 20%	SD Biosensor - STANDARD Q COVID-19 Ag
3.4% 6.8% 14.1% 1.3%	100% 100% 100% 89%	950 (941 to 950) 900 (891 to 900) 800 (792 to 800) 945 (938 to 948)	33 (31 to 35) 66 (61 to 70) 132 (122 to 141) 12 (6 to 21)	0 (0 to 10) 0 (0 to 9) 0 (0 to 8) 5 (2 to 12)	17 (15 to 19) 34 (30 to 39) 68 (59 to 78) 38 (29 to 44)	5% 10% 20% 5%	Coris Bioconcept Abbott - Panbio Covid-19 Ag
00 have COVID-1 1 – NPV	ents where 50, 100 and 2 PPV	Symptomatic participants: average sensitivity and specificity (and 95% CIs) applied to a hypothetical cohort of 1000 patients where 50, 100 and 200 have COVID-19 infection Test Prevalence TP (95% CI) FP (95% CI) FN (95% CI) TN (95% CI) PPV 1 - NPV	CIs) applied to a hypothe FN (95% CI)	ecificity (and 95% CFP (95% CI)	e sensitivity and sp TP (95% CI)	ticipants: averag	Symptomatic par infection Test
100)	99.1 (95.2 to 100)		69.2 (38.6 to 90.9)	13	127	1	SD Biosensor - STANDARD Q COVID-19 Ag
99.1)	98.1 (96.3 to 99.1)		48.9 (35.1 to 62.9)	47	474	1	Abbott - Panbio Covid-19 Ag
00)	100 (88.8 to 100)		28.6 (8.4 to 58.1)	14	45	2	Coris Bioconcept - COVID-19 Ag Respi-Strip
						rticipants	Asymptomatic participants
99.6)	99.1 (97.8 to 99.6)		88.1 (84.2 to 91.1)	336	1947	ω	SD Biosensor - STANDARD Q COVID-19 Ag
99.8)	99.5 (98.7 to 99.8)		75.1 (57.3 to 87.1)	252	1094	ω	Abbott - Panbio Covid-19 Ag

ID-19 infection			ID-19 infection				
Coris Bioconcept	0.5%	14 (4 to 29)	0 (0 to 1114)	36 (21 to 46)	9950 (8836 to 9950)	100%	0.4%
	1%	29 (8 to 58)	0 (0 to 1109)	71 (42 to 92)	9900 (8791 to 9900)	100%	0.7%
	2%	57 (17 to 116)	0 (0 to 1098)	143 (84 to 183)	9800 (8702 to 9800)	100%	1.4%
Abbott - Panbio	0.5%	24 (18 to 31)	189 (90 to 368)	26 (19 to 32)	9761 (9582 to 9860)	11%	0.3%
COVIG-13 Ag	1%	49 (35 to 63)	188 (89 to 366)	51 (37 to 65)	9712 (9534 to 9811)	21%	0.5%
	2%	98 (70 to 126)	186 (88 to 363)	102 (74 to 130)	9614 (9437 to 9712)	34%	1.0%
SD Biosensor	0.5%	35 (19 to 45)	90 (0 to 478)	15 (5 to 31)	9860 (9472 to 9950)	28%	0.2%
COVID-19 Ag	1%	69 (39 to 91)	89 (0 to 475)	31 (9 to 61)	9811 (9425 to 9900)	44%	0.3%
	2%	138 (77 to 182)	88 (0 to 470)	62 (18 to 123)	9712 (9330 to 9800)	61%	0.6%
Findings: rapid molecular tests	lecular tests						
Evaluations	Samples	SARS-CoV-2 cases		Average sensitivity (95% CI)	/ (95% CI)	Average spe	Average specificity (95% CI)
(studies)				[Range]		[Range]	
29 (26)	4351	1787		95.1 (90.5 to 97.6)		98.8 (98.3 to 99.2)	99.2)
				[57% to 100%]		[92% to 100%]	%]
Pooled results for	individual tests	using data from con	npliant with manufa	Pooled results for individual tests using data from compliant with manufacturer instructions for use	ruse		
Tests	Evaluations	Samples	SARS-CoV-2	Sensitivity (95% CI))	Specificity (95% CI)	95% CI)
			cases				
Abbott - ID NOW	4	812	222	73.0 (66.8 to 78.4)		99.7 (98.7 to 99.9)	99.9)
Cepheid - Xpert Xpress	2	100	29	100 (88.1 to 100)		97.2 (89.4 to 99.3)	99.3)
DRW - SAMBA II	1	149	33	87.9 (71.8 to 96.6)		97.4 (92.6 to 99.5)	99.5)



DNANudge COVID Nudge	н	386	71	94.4 (86.2 to 98.4)		100 (98.8 to 100)	0)
Average sensitivit	y and specificity	(and 95% CIs) appli	ed to a hypothetical	Average sensitivity and specificity (and 95% CIs) applied to a hypothetical cohort of 1000 patients where 50, 100	_	and 200 have COVID-19 infection	
Tests	Prevalence	TP (95% CI)	FP (95% CI)	FN (95% CI)	TN (95% CI)	PPV b	1 – NPV ^C
ID NOW	5%	37 (33 to 39)	3 (1 to 12)	14 (11 to 17)	947 (938 to 949)	93%	1.4%
	10%	73 (67 to 78)	3 (1 to 12)	27 (22 to 33)	897 (888 to 899)	96%	2.9%
	20%	146 (134 to 157)	2 (1 to 10)	54 (43 to 66)	798 (790 to 799)	98%	6.3%
Xpert Xpress	5%	50 (44 to 50)	27 (7 to 101)	0 (0 to 6)	923 (849 to 943)	65%	0.0%
	10%	100 (88 to 100)	25 (6 to 95)	0 (0 to 12)	875 (805 to 894)	80%	0.0%
	20%	200 (176 to 200)	22 (6 to 85)	0 (0 to 24)	778 (715 to 794)	90%	0.0%
SAMBA II	5%	44 (36 to 48)	25 (5 to 70)	6 (2 to 14)	925 (880 to 945)	64%	0.6%
	10%	88 (72 to 97)	23 (5 to 67)	12 (3 to 28)	877 (833 to 896)	79%	1.4%
	20%	176 (144 to 193)	21 (4 to 59)	24 (7 to 56)	779 (741 to 796)	89%	3.0%
COVID Nudge	5%	47 (43 to 49)	0 (0 to 11)	3 (1 to 7)	950 (939 to 950)	100%	0.3%
	10%	94 (86 to 98)	0 (0 to 11)	6 (2 to 14)	900 (889 to 900)	100%	0.6%
	20%	189 (172 to 197)	0 (0 to 10)	11 (3 to 28)	800 (790 to 800)	100%	1.4%
1 – NPV: 1 – negati positive;IFU: [manu scription polymera	ve predictive valu ufacturers'] instru se chain reaction;	1 - NPV: 1 - negative predictive value (the percentage of people with neg positive; IFU: [manufacturers'] instructions for use; PPV: positive predict scription polymerase chain reaction; TN: true negative; TP: true positive	beople with negative positive predictive va	1 – NPV: 1 – negative predictive value (the percentage of people with negative results who are infected); Ag: antigen; C positive; IFU: [manufacturers'] instructions for use; PPV: positive predictive value (the percentage of people with posit scription polymerase chain reaction; TN: true negative; TP: true positive	1 – NPV: 1 – negative predictive value (the percentage of people with negative results who are infected); Ag: antigen;CI: confidence interval; FN: false negative; FP: false positive; FU: [manufacturers'] instructions for use; PPV: positive predictive value (the percentage of people with positive results who are infected); RT-PCR: reverse transcription polymerase chain reaction; TN: true negative; TP: true positive	<pre>!: confidence interval; FN: false negative; FP: false ive results who are infected); RT-PCR: reverse tran</pre>	tive; FP: false ?: reverse tran-

 o Specificity only estimated in 8 of 26 evaluations by time after symptom onset.

c1-NPV (negative predictive value), where NPV is defined as the percentage of negative rapid test results that are truly negative according to the reference standard diagnosis. bPPV (positive predictive value) defined as the percentage of positive rapid test results that are truly positive according to the reference standard diagnosis.



BACKGROUND

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) and the resulting COVID-19 pandemic present important diagnostic evaluation challenges. These range from: understanding the value of signs and symptoms in predicting possible infection; assessing whether existing biochemical and imaging tests can identify infection or people needing critical care; and evaluating whether in vitro diagnostic tests can accurately identify and rule out current SARS-CoV-2 infection, and identify those with past infection, with or without immunity.

We are creating and maintaining a suite of living systematic reviews to cover the roles of tests and patient characteristics in the diagnosis of COVID-19. This review is the first update of a review summarising evidence of the accuracy of rapid antigen and molecular tests that are suitable for use at the point of care. In some scenarios the tests could potentially be used as alternatives to standard laboratory-based molecular assays, such as reverse transcription polymerase chain reaction (RT-PCR) assays, that are relied on for identifying current infection, in others they may be used where no testing is currently done. If sufficiently accurate, point-of-care tests have the potential to greatly expand access and speed of testing, In turn, if accurate, they may have greater impact on public health than laboratory-based molecular methods as they are less expensive, provide results more quickly and do not require the same technical expertise and laboratory capacity. These tests can be undertaken locally, avoiding the need for centralised testing facilities that rarely meet the needs of patients, caregivers, health workers and society as a whole, especially in low- and middleincome countries. As these are rapid tests, their results can be returned within the same clinical encounter, facilitating timely decisions concerning the need for isolation and contract tracing

Target condition being diagnosed

COVID-19 is the disease caused by infection with the SARS-CoV-2 virus. The key target conditions for this suite of reviews are current SARS-CoV-2 infection, current COVID-19 disease, and past SARS-CoV-2 infection. The tests included in this review concern the identification of current infection, as defined by reference standard methods of diagnosis, including molecular assays such as RT-PCR, or internationally recognised clinical guidelines for diagnosis of SARS-CoV-2. In the context of test evaluation, and throughout this review, we use the term 'reference standard' to denote the best available method (test or tests) for diagnosing the target condition, as opposed to other uses of the term in diagnostic virology (such as reference methods or reference materials).

For current infection, the severity of the disease is of ultimate importance for patient outcomes. However, rapid testing does not establish severity of disease, and for this review we consider the role of point-of-care tests for detecting SARS-CoV-2 infection of any severity, distinguishing only between symptomatic and asymptomatic infection.

COVID-19 public health interventions focus on reducing disease transmission, thus it is important to identify and isolate people who are infected before or whilst they are infectious. It is reasonably presumed that people with symptoms who meet national criteria for COVID-19 testing, or who are identified through contact tracing, have a high enough risk of being infectious to ask them to isolate.

However, assessing the risk of an individual being infectious in asymptomatic screening is more difficult, as there is no reference standard test for being 'infectious'. Using RT-PCR status as a reference standard (as is done for target condition of 'infection') will ensure that infectious people are not missed, but as RT-PCR continues to detect viral RNA days and weeks after the onset of infection will wrongly classify some people as infectious. Alternative reference standards that have been proposed for infectiousness include assessing the viability of the virus using viral culture, or using a value of the cycle threshold (Ct value) from RT-PCR results to group individuals above or below a particular value (as a proxy for viral load) as more or less likely to be infectious. Converting Ct values (also known as quantification cycle (Cq) or crossing point (Cp) values) into direct quantitative values of viral load (viral copies per cell) is possible but challenging, as the relationship between Ct values and viral load varies between machines and laboratories. Thus comparison at fixed Ct values is unlikely to be comparable across studies. Viral culture is unsuitable as a reference standard because it is technically complex and often unreliable, which leads to it being an insensitive test (the failure to culture virus potentially being a result of the culture technique and not an indicator of non-infectiousness). The suitability of RT-PCR is limited as the inverse relationship between viral load (Ct value) and risk of infection is a continuum of risk without there being a meaningful cut-point (with virus being cultured from samples with Ct values as high as 35 (Singanayagam 2020)). Similarly, those with low viral loads at the onset of infection will be missed. A preferable alternative, of tracking contacts for evidence of secondary infections, requires longitudinal follow-up and is better considered as a question about risk of transmission, which can be addressed using predictive modelling approaches (taking into account host, agent and environmental factors). This is in contrast to the diagnostic test accuracy paradigm which can only determine if individuals are infected at a single point in time.

For these reasons, this review only focuses on the target condition of 'infection' for both symptomatic and asymptomatic applications of tests. We do report results where they are presented split by an RT-PCR Ct value to report on accuracy according to groups with higher and lower viral load, but advise caution on their interpretation considering the lack of standardisation of PCR Ct values. Given the current state of the scientific knowledge we do not consider it appropriate to consider these as groups which are defined as 'infectious' and 'not infectious'.

RT-PCR carries a very small risk of false positive results for infection and a higher risk of false negative results. False positive results may result from failures in sampling or laboratory protocols (e.g. mislabelling), contamination during sampling or processing, or low-level reactions during PCR (Healy 2020; Mayers 2020). At times when SARS-CoV-2 infections have been rare, population prevalence surveys using RT-PCR have shown test positivity rates of 0.44% (95% credible interval: 0.22% to 0.76%) (August 2020; ONS 2020), and 0.077% (0.065%, 0.092%) (June to July 2020; Riley 2020 React-1 study). These values can be used to place an upper bound on the possible false positive rate of RT-PCR of less than 0.077% (as the total numbers testing positive will comprise both true positive and false positive RT-PCR results). The World Health Organization (WHO) recently issued a notice of concern regarding interpretation of specimens at or near the limit for PCR positivity (i.e. those with high cycle threshold (Ct) values), citing potential difficulties in distinguishing the presence of the target virus from these types of



background 'noise' (WHO 2020a). False negative rates have been estimated by looking at individuals with symptoms who initially test negative, but positive on a subsequent test. These rates have been estimated to be as high as 20% to 30% in the first week of symptom onset; Arevalo-Rodriguez 2020; Yang 2020a; Zhao 2020; Kucirka 2020). Including probable COVID-19 cases within the target condition, as defined by internationally recognised clinical guidelines for diagnosis of SARS-CoV-2 will partially mitigate these missed cases.

Index test(s)

The primary consideration for the eligibility of tests for inclusion in this review is that they should detect current infection and should have the capacity to be performed at the 'point of care' or in a 'nearpatient' testing role. There is an ongoing debate around the specific use and definitions of these terms, therefore for the purposes of this review, we consider 'point-of-care' and 'near patient' to be synonymous, but for consistency and avoidance of confusion, we use the term 'point-of-care' throughout.

We have adapted a definition of point-of-care testing, namely that it "refers to decentralized testing that is performed by a minimally trained healthcare professional near a patient and outside of central laboratory testing" (WHO 2018), with the additional caveat that test results must be available within a single clinical encounter (Pai 2012). Our criteria for defining a point-of-care test are therefore:

- the equipment for running and or reading the assay must be portable or easily transported, although mains power may be required;
- minimal sample preparation requirements, for example, singlestep mixing, with no requirement for additional equipment or precise sample volume transfer unless a disposable automatic fill or graduated transfer device is used;
- minimal biosafety requirements, for example, personal protective equipment (PPE) for sample collector and test operator, good ventilation and a biohazard bag for waste disposal:
- no requirement for a temperature-controlled environment; and
- test results available within two hours of sample collection.

Tests for detection of current infection that are currently suitable for use at the point of care include antigen tests and molecularbased tests. Both types of test use the same respiratory-tract samples acquired by swabbing, washing or aspiration as for laboratory-based RT-PCR. Rapid antigen tests use lateral flow immunoassays, which are disposable devices, usually in the form of plastic cassettes akin to a pregnancy test. Viral antigen is captured by dedicated antibodies that are either colloidal gold- or fluorescent-labelled. Antigen detection is indicated by visible lines appearing on the test strip (colloidal gold-based immunoassays, or CGIA), or through fluorescence, which can be detected using an immunofluorescence analyser (fluorescence immunoassays or FIA). Molecular-based tests to detect viral ribonucleic acid (RNA) have historically been laboratory-based assays using RT-PCR technology (see Alternative test(s)). In recent years, automated, single-step RT-PCR methods have been developed, as well as other nucleic acid amplification methods, such as isothermal amplification, that do not require the sophisticated thermo cycling involved in RT-PCR (Green 2020). These technological advances have allowed molecular technologies to be developed that are

suitable for use in a point-of-care context (Kozel 2017), however they still require small portable machines and many take longer to produce results than antigen tests.

Following the emergence of COVID-19 there has been prolific industry activity to develop accurate tests. The Foundation for Innovative Diagnostics (FIND) and Johns Hopkins Centre for Health Security have maintained online lists of available tests for SARS-CoV-2 (FIND 2020). At the time of writing (5 January 2021), FIND listed 129 rapid antigen tests, 118 of which are described as "commercialised" and 92 have been identified as having regulatory approval. These numbers are a substantial increase on the 48 listed, 32 commercialised and 21 with regulatory approval at the time of our original review (19 July 2020). A total of 142 molecular tests were described as automated, including both laboratory-based assays and assays suitable for use outside of a laboratory setting (i.e. near or at the point of care). Further information from FIND indicates that 53 of the 142 assays were categorised as point-of-care or near point-of-care tests, including 43 with regulatory approval. This classification was based on the information provided to FIND by the test manufacturers and does not necessarily mean that these tests meet the criteria for point-of-care tests that we have specified for this review. The numbers of tests of these types will continue to increase over time.

Given the urgent need to identify the evidence base for tests that are available for purchase, the focus of this first update of the review is on tests that are commercially produced. All commercially produced assays are supplied with a specific product code, product inserts or instructions for use (IFU) sheets that document the intended use of the test; sample storage and preparation and testing procedures; who should deliver the test and in whom; and any restrictions around the type of samples that can be used.

There are many proposals for serial testing with lateral flow tests to detect infection, rather than a single use. In this case it would be appropriate to evaluate the accuracy of the strategy rather than a single test.

Clinical pathway

Patients may be tested for SARS-CoV-2 when they present with symptoms, have had known exposure to a confirmed case, or in a screening context, with no known exposure to SARS-CoV-2. The standard approach to diagnosis of SARS-CoV-2 infection is through laboratory-based testing of swab samples taken from the upper respiratory (e.g. nasopharynx, oropharynx) or lower respiratory tract (e.g. bronchoalveolar lavage or sputum) with RT-PCR. RT-PCR is the primary method for detecting infection during the acute phase of the illness while the virus is still present. Both the WHO and the China CDC (National Health Commission of the People's Republic of China), have produced case definitions for COVID-19 that include the presence of convincing clinical evidence (some including positive serology tests) when RT-PCR is negative (Appendix 1).

Prior test(s)

Signs and symptoms are used in the initial diagnosis of suspected SARS-CoV-2 infection and to help identify those requiring tests. A number of key symptoms have been suggested as indicators of mild to moderate COVID-19, including: cough, fever greater than 37.8 °C, headache, breathlessness, muscle pain, fatigue, and loss of sense of smell and taste (Struyf 2021). However, the recently



published review of signs and symptoms found good evidence for the accuracy for these symptoms alone or in combination to be lacking (Struyf 2021).

Where people are asymptomatic but are being tested as part of screening (e.g. universal testing of students as part of a risk-reduction effort) or on the basis of epidemiological risk factors, such as exposure to someone with confirmed SARS-CoV-2 or following travel to more highly endemic countries, no prior tests will have been conducted.

Role of index test(s)

For most settings in which testing for acute SARS-CoV-2 infection in symptomatic individuals takes place, results of molecular laboratory-based RT-PCR tests are unlikely to be available within a single clinical encounter. Point-of-care tests potentially have a role either as a replacement for RT-PCR (if sufficiently accurate), or as a means of triaging and rapid management (quarantine or treatment, or both), with confirmatory RT-PCR testing for those with negative rapid test results (CDC 2020; WHO 2020b). Obtaining quick results within a healthcare visit will allow faster decisions about isolation and healthcare interventions for those with positive test results, and allow contact tracing to begin in a more timely manner. Modelling studies suggest contact tracing is most effective if it starts within 24 hours of case detection, with delays in testing (e.g. due to laboratory turnaround time for reporting PCR results) leading to reductions in the proportion of onward transmissions per index case that can be prevented by track and trace (Kretzschmar 2020).

If sufficiently accurate, negative rapid test results in symptomatic patients could allow faster return to work or school, therefore conferring important economic and educational implications. Negative results also allow immediate consideration of other causes of symptoms, which may be time-sensitive, for example bacterial pneumonia or thrombo-embolism.

For asymptomatic individuals, if accurate, rapid tests may also be considered for screening at-risk (exposed) populations, for example in hospital workers or in local outbreaks.

Rapid tests, particularly antigen tests which can be more easily delivered at scale, could also be used for mass screening purposes as recently piloted in Slovakia and in Liverpool UK (University of Liverpool 2020), or used in a more targeted fashion such as single test application at airports or for border entry, to allow entry to large public gatherings, or screening students as a risk-reduction strategy (Ferguson 2020). Preliminary data on the rollout of such a policy in the UK has highlighted the many challenges in such an approach (Deeks 2020a; Nabavi 2021), and the requirement for full and proper field trial evaluations. Frequent repeated use of antigen tests in asymptomatic individuals with no known exposure to identify COVID-19 cases has also been proposed (Larremore 2020), but field trial evaluations would be required to determine whether promising results from modelling studies can be borne out in practical settings (Crozier 2021).

Alternative test(s)

This review is one of seven that cover the range of tests and clinical characteristics being considered in the management of COVID-19 (Deeks 2020b; McInnes 2020), five of which have already been published (Deeks 2020c; Salameh 2020; Stegeman 2020; Struyf 2021), including the first iteration of this review (Dinnes 2020). Full

details of the alternative tests and evidence of their accuracy is summarised in these reviews. The SARS-CoV-2-specific biomarker tests that might be considered as alternatives to point-of-care tests are considered here.

Laboratory-based molecular tests

RT-PCR tests for SARS-CoV-2 identify viral ribonucleic acid (RNA). Reagents for RT-PCR were rapidly produced once the viral RNA sequence was published (Corman 2020). Testing is undertaken in central laboratories and can be very labour-intensive, with several points along the path of performing a single test where errors may occur, although some automation of parts of the process is possible. The amplification process requires thermal cycling equipment to allow multiple temperature changes within a cycle, with cycles repeated up to 40 times until viral DNA is detected (Carter 2020). Although the amplification process for RT-PCR can be completed in a relatively short timeframe, the stages of extraction, sample processing and data management (including reporting) mean that test results are typically only available in 24 to 48 hours. Where testing is undertaken in a centralised laboratory, transport times increase this further. The time to result for fully automated RT-PCR assays is shorter than for manual RT-PCR, however most assays still require sample preparation steps that make them unsuitable for use at the point of care. Other nucleic acid amplification methods, including loop-mediated isothermal amplification (LAMP), or CRISPR-based nucleic acid detection methods, that allow amplification at a constant temperature are now commercially available (Chen 2020). These methods have the potential to reduce the time to produce test results after extraction and sample processing to minutes, but the time for the whole process may still be significant. Laboratory-based molecular tests are most often applied to upper and lower respiratory samples although they are also being used on faecal and urine samples.

Antibody tests

Serology tests to measure antibodies to SARS-CoV-2 have been evaluated in people with active infection and in convalescent cases (Deeks 2020c). Antibodies are formed by the body's immune system in response to infections, and can be detected in whole blood, plasma or serum. Antibody tests are available for laboratory use including enzyme-linked immunosorbent assay (ELISA) methods, or more advanced chemiluminescence immunoassays (CLIA). There are also rapid lateral flow assays (LFA)s for antibody testing that use a minimal amount of whole blood, plasma or serum on a testing strip as opposed to the respiratory specimens that are used for rapid antigen tests; all assays for antibody detection are considered in Deeks 2020c.

Rationale

It is essential to understand the clinical accuracy of tests and clinical features to identify the best way they can be used in different settings to develop effective diagnostic and management pathways for SARS-CoV-2 infection and disease. The suite of Cochrane living systematic reviews summarises evidence on the clinical accuracy of different tests and diagnostic features. Estimates of accuracy from these reviews will help inform diagnosis, screening, isolation, and patient-management decisions.



Summary of the previous version of the review

The first iteration of this review (Dinnes 2020), included 22 publications reporting on a total of 18 study cohorts with 3198 unique samples, 1775 of which had confirmed SARS-CoV-2 infection. We identified data for eight commercial tests (four antigen and four molecular) and one in-house antigen test.

We did not find any studies at low risk of bias and had concerns about applicability of results across all studies. We judged patient selection to be at high risk of bias in 50% of the studies because of deliberate oversampling of samples with confirmed SARS-CoV-2 infection (sample enrichment) and unclear in 38% (7/18) because of poor reporting. Sixteen (89%) studies used only a single, negative RT-PCR to confirm the absence of SARS-CoV-2 infection, risking missing infection. There was a lack of information on blinding of index test (n = 11), and about participant exclusions from analyses (n = 10). We did not observe differences in methodological quality between antigen and molecular test evaluations.

The eight evaluations of antigen tests reported considerable variation in sensitivity across studies (from 0% to 94%) with less variation in specificities (from 90% to 100%). The average sensitivity was 56.2% (95% CI 29.5 to 79.8%) and average specificity was 99.5% (95% CI 98.1% to 99.9%) (based on 943 samples, 596 with confirmed SARS-CoV-2). Data for individual antigen tests were limited with no more than two studies for any test.

We observed less variation in sensitivities across 13 evaluations of rapid molecular assays (range 68% to 100%) with similar variation in specificities (range 92% to 100%). Average sensitivity was 95.2% (95% CI 86.7% to 98.3%) and specificity 98.9% (95% CI 97.3% to 99.5%) based on a total of 2255 samples.

We were able to calculate pooled results for only two molecular tests: ID NOW (Abbott Laboratories; 5 evaluations) and Xpert Xpress (Cepheid Inc; 6 evaluations). Summary sensitivity for the Xpert Xpress assay (99.4%, 95% CI 98.0% to 99.8%) was 22.6 (95% CI 18.8 to 26.3) percentage points higher than that of ID NOW (76.8%, (95% CI 72.9% to 80.3%), whilst the specificity of Xpert Xpress (96.8%, 95% CI 90.6% to 99.0%) was marginally lower than ID NOW (99.6%, 95% CI 98.4% to 99.9%; a difference of –2.8 percentage points (95% CI from 6.4 percentage points lower to 0.8 higher).

Changes in the evidence base since the previous version

There has been a considerable increase in the number of evaluations available of antigen tests, and a lesser rise in the number of evaluations of molecular tests. More studies report key population features such as setting, and symptom status, and there has been an increase in direct swab testing as would occur in a point-of-care setting. However, due to the nature of sampling and the use of direct swab testing, few comparative studies are available. This review considers the available evidence in relevant population groups and settings according to test brand and compliance with manufacturer IFUs. We used the WHO's priority target product profiles for COVID-19 diagnostics (i.e. acceptable performance criterion of sensitivity \geq 80% and specificity \geq 97%, or desirable criterion of \geq 80% sensitivity and \geq 99% specificity; WHO 2020c) as a benchmark against which to consider test performance.

We will update this review as often as is feasible to ensure that it provides current evidence about the accuracy of point-of-care tests.

This review follows a generic protocol that covers six of the seven Cochrane COVID-19 diagnostic test accuracy reviews (Deeks 2020b). The Background and Methods sections of this review therefore use some text that was originally published in the protocol (Deeks 2020b), and text that overlaps some of our other reviews (Deeks 2020c; Struyf 2021).

OBJECTIVES

To assess the diagnostic accuracy of rapid point-of-care antigen and molecular-based tests to determine if a person presenting in the community or in primary or secondary care has current SARS-CoV-2 infection, and to consider accuracy separately in symptomatic and asymptomatic population groups.

We estimated accuracy overall and separately according to symptom status (symptomatic and asymptomatic). Although we might expect to see differences in accuracy for testing of asymptomatic individuals with an epidemiological exposure to SARS-CoV-2 (targeted screening) compared to testing of asymptomatic individuals in a population screening setting, we did not anticipate finding sufficient numbers of studies for each testing application to allow any such difference to be explored. We will revisit this decision in subsequent iterations of this review.

Secondary objectives

Where data are available, we will investigate potential sources of heterogeneity that may influence diagnostic accuracy (either by stratified analysis or meta-regression) according to test method and index test, participant or sample characteristics (duration of symptoms and viral load), study setting, study design and reference standard used.

We investigated adherence to manufacturers' IFUs in sensitivity analyses.

METHODS

Criteria for considering studies for this review

Types of studies

We applied broad eligibility criteria to include all patient groups (that is, if patient population was unclear, we included the study) and all variations of a test.

We included studies of all designs that produce estimates of test accuracy or provide data from which we can compute estimates, including the following.

- Studies restricted to participants confirmed to either have (or to have had) the target condition (to estimate sensitivity) or confirmed not to have (or have had) the target condition (to estimate specificity). These types of studies may be excluded in future review updates.
- Single-group studies, which recruit participants before disease status has been ascertained
- Multi-group studies, where people with and without the target condition are recruited separately (often referred to as two-gate or diagnostic case-control studies)
- · Studies based on either patients or samples



We excluded studies from which we could not extract data to compute either sensitivity or specificity.

We carefully considered the limitations of different study designs in the quality assessment and analyses.

We included studies reported in published journal papers, as preprints, and publicly available reports from independent bodies.

Participants

We included studies recruiting people presenting with suspicion of current SARS-CoV-2 infection or those recruiting populations where tests were used to screen for disease (for example, contact tracing or community screening).

We also included studies that recruited people known to have SARS-CoV-2 infection and known not to have SARS-CoV-2 infection (i.e. cases only or multi-group studies).

We excluded small studies with fewer than 10 samples or participants. Although the size threshold of 10 is arbitrary, such small studies are likely to give unreliable estimates of sensitivity or specificity and may be biased.

Index tests

We included studies evaluating any rapid antigen or molecularbased test for diagnosis of SARS-CoV-2, if it met the criteria outlined in the Background, that is:

- · requiring minimal equipment;
- minimal sample preparation and biosafety considerations;
- results available within two hours of sample collection; and
- should be commercially produced (with test name and manufacturer or distributor documented).

All sample types (respiratory or non-respiratory) were eligible. Strategies based on multiple applications of a test were also eligible for inclusion.

Target conditions

The target condition was current SARS-CoV-2 infection (either symptomatic or asymptomatic). We also refer to SARS-CoV-2 infection as 'COVID-19 infection', particularly in the Plain Language Summary and Summary of findings 1.

Reference standards

We anticipated that studies would use a range of reference standards to define both the presence and absence of SARS-CoV-2 infection. For the QUADAS-2 (Quality Assessment tool for Diagnostic Accuracy Studies; Whiting 2011), assessment we categorised each method of defining the presence of SARS-CoV-2 according to the risk of bias (the chances that it would misclassify the presence or absence of infection) and whether it defined COVID-19 in an appropriate way that reflected cases encountered in practice. Likewise, we considered the risk of bias in definitions of the absence of SARS-CoV-2, and whether the definition captured all those who might be tested in practice.

Evaluations of molecular tests generally consider agreement between molecular assays, for example, agreement of a new rapid test against a more standard RT-PCR test. For the purposes of this review, we considered RT-PCR to be the 'reference standard' for SARS-CoV-2 infection, and present results as 'sensitivity' and 'specificity' as opposed to percentage agreement. The result of further RT-PCR analysis of discrepant cells (samples with results disagreeing on the rapid test and the RT-PCR) were also considered in sensitivity analyses. As discrepant analysis involves retesting only a sub-sample of patients selected according to index and reference standard results, it can introduce bias (Hadgu 1999). Retesting of all samples with a second test in a composite reference standard would be preferable when there are concerns over the accuracy of the first reference test.

Search methods for identification of studies

Electronic searches

We used two main sources for our electronic searches through 30 September 2020, which were devised with the help of an experienced Cochrane Information Specialist with diagnostic test accuracy review expertise (RSp). These searches aimed to identify all articles related to COVID-19 and SARS-CoV-2 and were not restricted to those evaluating a particular type of test. Thus, the searches used no terms that specifically focused on an index test, diagnostic accuracy or study methodology.

Cochrane COVID-19 Study Register searches

We used the Cochrane COVID-19 Study Register (covid-19.cochrane.org/), for searches conducted from inception of the Register to 28 March 2020. At that time, the register was populated by searches of PubMed, as well as trials registers at US National Institutes of Health Ongoing Trials Register ClinicalTrials.gov (clinicaltrials.gov) and the WHO International Clinical Trials Registry Platform (apps.who.int/trialsearch).

Search strategies were designed for maximum sensitivity, to retrieve all human studies on COVID-19 and with no language limits. See Appendix 2.

COVID-19 Living Evidence Database from the University of Bern

From 28 March 2020, we used the COVID-19 Living Evidence database from the Institute of Social and Preventive Medicine (ISPM) at the University of Bern (www.ispm.unibe.ch), as the primary source of records for the Cochrane COVID-19 diagnostic test accuracy reviews. This search includes PubMed, Embase, and preprints indexed in bioRxiv and medRxiv databases. The strategies as described on the ISPM website are described here (ispmbern.github.io/covid-19/). See Appendix 3. To ensure comprehensive coverage we also downloaded records from the 'Bern feed' from 1 January to 28 March 2020 and de-duplicated them against those obtained via the Cochrane COVID-19 Study Register.

Due to the increased volume of published and preprint articles, from 25 May 2020 onwards we used artificial intelligence text analysis to conduct an initial classification of documents, based on their title and abstract information, for relevant and irrelevant documents (Appendix 4).

The decision to focus primarily on the Bern feed was because of the exceptionally large numbers of COVID-19 studies available only as preprints. We are continuing to monitor the coverage of the Cochrane COVID-19 Study Register and may move back to it as the primary source of records for subsequent review updates.



Other electronic sources

Prior to 28 March 2020 (when we began using the 'Bern feed'), we identified Embase records through the Centers for Disease Control and Prevention (CDC), Stephen B Thacker CDC Library, COVID-19 Research Articles Downloadable Database (cdc.gov/library/researchguides/2019novelcoronavirus/researcharticles.html), and de-duplicated them against results from the Cochrane COVID-19 Study Register. See Appendix 5.

We also checked our search results against two additional repositories of COVID-19 publications up to 30 September 2020:

- the Evidence for Policy and Practice Information and Coordinating Centre (EPPI-Centre) 'COVID-19: Living map of the evidence' (eppi.ioe.ac.uk/COVID19_MAP/covid_map_v4.html);
- the Norwegian Institute of Public Health 'NIPH systematic and living map on COVID-19 evidence' (www.nornesk.no/ forskningskart/NIPH_diagnosisMap.html)

Both repositories allow their contents to be filtered according to studies potentially relating to diagnosis, and both have agreed to provide us with updates of new diagnosis studies added.

Searching other resources

We have also contacted or accessed the websites of independent research groups undertaking test evaluations (for example, UK Public Health England (PHE), the Société Française Microbiologie (SFM), the Dutch National Institute for Public Health and the Environment (RIVM)) and studies co-ordinated by FIND (finddx.org/covid-19/sarscov2-eval) and accessed the Diagnostics Global Health listing of manufacturer independent evaluations of antigen detecting rapid diagnostic tests (Ag-RDTs) for SARS-CoV-2 (diagnosticsglobalhealth.org). We last accessed these additional resources on 16 November 2020.

We appeal to researchers to supply details of additional published or unpublished studies at the following email address, which we will consider for inclusion in future updates (coviddta@contacts.bham.ac.uk).

Data collection and analysis

Selection of studies

A team of experienced systematic review authors from the University of Birmingham screened the titles and abstracts of all records retrieved from the literature searches following the application of artificial intelligence text analysis (described in Electronic searches). Two review authors independently screened studies in Covidence. A third, senior review author resolved any disagreements. We tagged all records selected as potentially eligible according to the Cochrane COVID-19 diagnostic test accuracy review(s) for which they might be eligible and we then exported them to separate Covidence reviews for each review title.

We obtained the full texts for all studies flagged as potentially eligible. Two review authors independently screened the full texts for one of the COVID-19 biomarker reviews (molecular, antigen or antibody tests). We resolved any disagreements on study inclusion through discussion with a third review author.

Data extraction and management

One review author extracted the characteristics of each study, which a second review author checked. Items that we extracted are listed in Appendix 6.

Both review authors independently performed data extraction of 2x2 contingency tables of the number of true positives, false positives, false negatives and true negatives. They resolved disagreements by discussion. Where possible, we separately extracted data according to symptom status (symptomatic, asymptomatic, mixed symptom status or not reported), viral load (high or low, according to Ct cut-offs defined within each study), and time post-symptom onset (week one versus week two) and for molecular assays, before and after re-analysis of samples in discrepant cells. For categorisation by symptom status, we classed studies reporting at least 75% of participants as symptomatic as 'mainly symptomatic', we considered studies with less than 75% symptomatic participants to report 'mixed' groups along with those that reported recruiting both symptomatic and asymptomatic participants but did not provide the percentages in each group. We considered studies that provided no information as to the symptom status of included participants 'not reported'. We also coded evaluations according to compliance with manufacturer IFUs. We based coding on three aspects of testing:

- sample type (use of any sample not explicitly mentioned on the IFU scored 'No', otherwise scored 'Yes'),
- provision of instructions for samples in viral transport medium ((VTM); only scored for evaluations using samples in VTM and only scored 'Yes' if specific instructions provided; scored 'Unclear' if VTM used and instructions for use of samples in VTM not documented in IFU); and
- 3. timing between sample collection and testing (scored 'Yes' only if all tests were carried out within specified time period, e.g. immediate on-site testing, or for testing in laboratories if all tests reported to have been carried out within specified time period; scored 'Unclear' if time frame for testing was not reported and 'No' if any testing was carried out beyond the maximum stipulated timeframe).

We encourage study authors to contact us regarding missing details on the included studies (coviddta@contacts.bham.ac.uk).

Assessment of methodological quality

Two review authors independently assessed risk of bias and applicability concerns using the QUADAS-2 checklist tailored to this review (Appendix 7; Whiting 2011). The two review authors resolved any disagreements by discussion.

Ideally, studies examining the use of tests in symptomatic people should prospectively recruit a representative sample of participants presenting with signs and symptoms of COVID-19, either in community or primary care settings or to a hospital setting, and they should clearly record the time of testing after the onset of symptoms. Studies in asymptomatic people at risk of infection should document time from exposure. Studies applying tests in a screening setting should document eligibility criteria for screening, particularly if a targeted approach is used and should take care to record any previous confirmed or suspected SARS-CoV-2 infection or any relevant epidemiological exposures. Studies should perform tests in their intended use setting, using appropriate samples with



or without viral transport medium and within the time period following specimen collection as indicated in the IFU document. Tests should be performed by relevant personnel (e.g. healthcare workers), and should be interpreted blinded to the final diagnosis (presence or absence of SARS-CoV-2). The reference standard diagnosis should be blinded to the result of the rapid test, and should not incorporate the result of the rapid test. If the reference standard includes clinical diagnosis of COVID-19 for RT-PCR-negative patients, then established criteria should be used. Studies including samples from participants known not to have COVID-19 should use pre-pandemic sources or if contemporaneous samples then at least two RT-PCR-negative tests were required to confirm the absence of infection. Data should be reported for all study participants, including those where the result of the rapid test was inconclusive, or participants in whom the final diagnosis of COVID-19 was uncertain. Studies should report whether results relate to participants (one sample per participant), or samples (multiple samples per participant).

Statistical analysis and data synthesis

We analysed rapid antigen and molecular tests separately. Studies often referred to 'samples' rather than 'patients', especially for the rapid molecular tests, however for many studies we do not suspect that inclusion of multiple samples per study participant was a significant issue. For consistency of terminology throughout the review, we refer to results on a per-sample basis. If studies evaluated multiple tests in the same samples, we included them multiple times. We present estimates of sensitivity and specificity per study for each test brand using paired forest plots, and summarise results using average sensitivity and specificity in tables as appropriate. As heterogeneity is apparent in many analyses, these point estimates must be interpreted as the average of a distribution of values.

We did not make any formal comparisons between antigen assay brands because of the large number of different assays and small study numbers for many of them. We did however carry out a formal comparison (based on between-study comparisons) for studies using two brands of molecular tests (ID NOW (Abbott Laboratories) and Xpert Xpress (Cepheid Inc)).

We estimated summary sensitivities and specificities with 95% confidence intervals (CI) using the bivariate model (Reitsma 2005), via the meqrlogit command of Stata/SE 16.0. When few studies were available, we simplified models by first assuming no correlation between sensitivity and specificity estimates and secondly by setting near-zero variance estimates of the random effects to zero (Takwoingi 2017). In cases where there was only one study per test, we reported individual sensitivities and specificities with 95% CI constructed using the binomial exact method.

Where studies presented only estimates of sensitivity or of specificity, we fitted univariate, random-effects, logistic regression models. In a number of instances where there was 100% sensitivity or specificity for all evaluations, we computed estimates and 95% CIs by summing the counts of TP, FP, FN and TN across 2x2 tables. These analyses are clearly marked in the tables. We present all estimates with 95% confidence intervals.

Investigations of heterogeneity

We examined heterogeneity between studies by visually inspecting the forest plots of sensitivity and specificity. Where adequate data were available, we investigated heterogeneity related to symptom status, time post-symptom onset, viral load, test brand, and test method by including indicator variables in the random-effects logistic regression models. Absolute differences between the sensitivity or specificity and the P values were reported from the model. In instances where only one study was available per test or when tests were being directly compared following summing of counts of the 2x2 tables, we performed test comparison using the two-sample test of proportions. Few studies reported specificity estimates by time after symptom onset, therefore for this variable and for analyses by viral load, we considered only effects on sensitivity.

Sensitivity analyses

We performed four sensitivity analyses.

- We estimated summary sensitivities and specificities according to test brand and symptom status using only studies that were compliant to the IFU.
- 2. We estimated sensitivity with and without studies that only evaluated samples with RT-PCR-confirmed SARS-CoV-2 (and thus did not estimate specificity).
- 3. We performed the same analysis for specificity in studies that only evaluated RT-PCR-negative control samples.
- 4. We made comparisons between analyses using the primary reference standard and analyses using results adjusted after retesting of samples with discrepant results with a second RT-PCR test (discrepant analysis).

Assessment of reporting bias

We made no formal assessment of reporting bias but have indicated where we were aware that study results were available but unpublished.

Summary of findings

We summarised key findings in a 'Summary of findings' table indicating the strength of evidence for each test and findings, and highlighted important gaps in the evidence.

Updating

We are aware of additional studies published since the electronic searches were conducted on 30 September 2020 and plan to update this review. We have already conducted the next search to 1 January 2021.

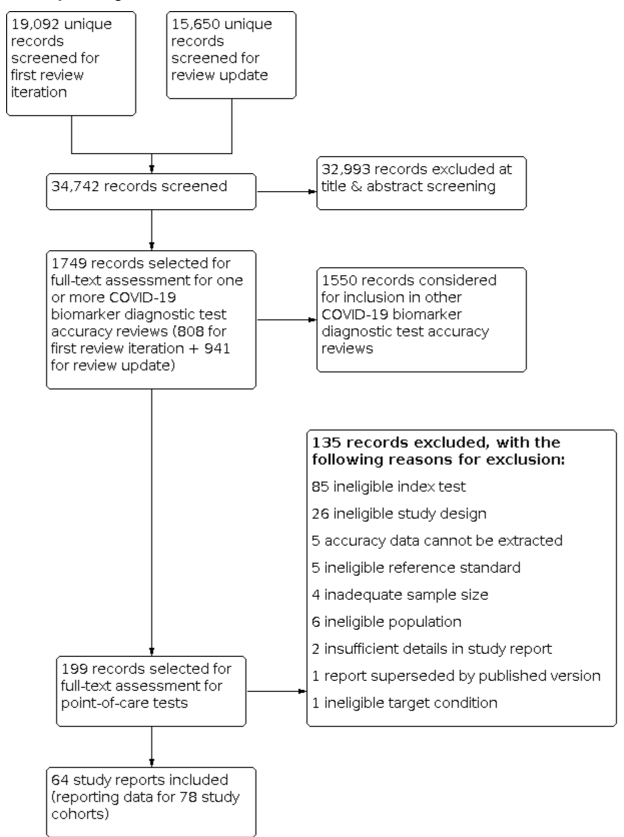
RESULTS

Results of the search

We screened 34,742 unique records (published or preprints) for inclusion in the complete suite of reviews to assist in the diagnosis of COVID-19 (Deeks 2020b; McInnes 2020). Of 1749 records selected for further assessment for inclusion in any of the four molecular, antigen or antibody test reviews, we identified 199 full-text reports requiring assessment for inclusion in this review; 90 for the first iteration of the review and 109 for this review update. See Figure 1 for the PRISMA flow diagram of search and eligibility results (McInnes 2018; Moher 2009).



Figure 1. Study flow diagram





We included 64 reports in this review, and we excluded 135 publications that did not meet our inclusion criteria. Exclusions were mainly based on index test (n = 85) or ineligible study designs (n = 26), for example, designs that did not allow estimation of test accuracy. The reasons for exclusion of all 135 publications are provided in Characteristics of excluded studies. Appendix 8 provides a list of studies evaluating eligible tests but excluded for other reasons (n = 5), and studies evaluating technologies not yet suitable for use at the point of care (n = 41).

Of the 64 study reports, 18 were available only as preprints, 38 were published papers and eight were publicly available reports either from independent reference laboratories (one from Public Health England and two identified via the SMF) or were independent evaluations co-ordinated by FIND (n = 5).

We contacted the authors of 10 study reports for further information (Blairon 2020; Courtellemont 2020; Diao 2020; Gibani 2020; Gremmels 2020(a); Linares 2020; Nash 2020; Porte 2020a; Schildgen 2020 [A]; Weitzel 2020 [A]), and received replies and the requested information with one exception (Linares 2020). We also contacted the evaluation teams at FIND and Public Health England and received additional information about study methods from FIND and some additional data from Public Health England.

The 64 included study reports relate to 78 separate studies. Please note when naming studies, we use the letters [A], [B], [C] etc. in square brackets to indicate data on different tests evaluated in the same study and (a), (b), (c) to indicate data from different participant cohorts from the same study report. For example, the five included reports from FIND correspond to eight 'studies' because three reports separately provided data from more than one evaluation centre.

Of the 78 studies, 77 reported data for respiratory samples and one (Szymczak 2020), reported data for faecal samples. The main results, Tables and Figures focus on the respiratory samples, with Szymczak 2020 reported separately.

Description of included studies

The 77 studies using respiratory samples included a total of 24,418 unique samples, with 7484 samples with RT-PCR-confirmed SARS-CoV-2 (some samples were analysed by more than one index test). Forty-eight studies evaluated antigen tests (Albert 2020; Alemany 2020; Billaud 2020; Blairon 2020; Cerutti 2020; Courtellemont 2020; Diao 2020; Fenollar 2020(a); Fenollar 2020(b); FIND 2020a; FIND 2020b; FIND 2020c (BR); FIND 2020c (CH); FIND 2020d (BR); FIND 2020d (DE); FIND 2020e (BR); FIND 2020e (DE); Fourati 2020 [A]; Gremmels 2020(a); Gremmels 2020(b); Gupta 2020; Kruger 2020(a); Kruger 2020(b); Kruger 2020(c); Lambert-Niclot 2020; Linares 2020; Liotti 2020; Mak 2020; Mertens 2020; Nagura-Ikeda 2020; Nash 2020; PHE 2020(a); PHE 2020(b); PHE 2020(c) [non-HCW tested]; PHE 2020(d) [HCW tested]; PHE 2020(d) [Lab tested]; PHE 2020(e); Porte 2020a; Porte 2020b [A]; Schildgen 2020 [A]; Scohy 2020; Shrestha 2020; Takeda 2020; Van der Moeren 2020(a); Van der Moeren 2020(b); Veyrenche 2020; Weitzel 2020 [A]; Young 2020) and 29 studies evaluated molecular tests (Assennato 2020; Broder 2020; Chen 2020a; Collier 2020; Cradic 2020(a); Cradic 2020(b); Dust 2020; Ghofrani 2020; Gibani 2020; Goldenberger 2020; Harrington 2020; Hogan 2020; Hou 2020; Jin 2020; Jokela 2020; Lephart 2020 [A]; Lieberman 2020; Loeffelholz 2020; Mitchell 2020; Moore 2020; Moran 2020; Rhoads 2020; Smithgall 2020 [A]; SoRelle 2020; Stevens 2020; Thwe 2020; Wolters 2020; Wong 2020; Zhen 2020 [A]). Summary study characteristics are presented in Table 1 with further details of study design and index test details in Appendix 9 and Appendix 10 for antigen assays and Appendix 11 and Appendix 12 for molecular assays. Full details are provided in the Characteristics of included studies table.

The median sample size of the included studies is 182 (interquartile range (IQR) 104 to 400) and median number of SARS-CoV-2 confirmed samples included is 63 (IQR 38 to 119). Sample sizes for antigen test evaluations were larger than those for molecular test evaluations (median 291.5 (IQR 155 to 502.5) compared to 104 (IQR 75 to 172)). Half of the studies (39/77, 51%) were conducted in Europe, 20 in North America, seven in South America, seven in Asia, one study included samples from more than one country and in one, the country of sample origin was unclear.

Participant characteristics

Antigen tests

Over half of the antigen test studies included samples from participants presenting in the community for COVID-19 testing at: community test centres (22/48, 46%); emergency departments (3, 6%); or as part of contact tracing or outbreak investigations (4, 8%) (Table 1). Eleven antigen test studies (23%) selected samples from those submitted to laboratories for routine RT-PCR testing with limited detail of the participants providing the samples ('laboratory-based' studies), or included multiple (8%) or unclear (2%) settings. Over half of antigen test studies were conducted in symptomatic (16, 33%) or mainly symptomatic (11, 23%) populations, with only three (6%) exclusively in asymptomatic populations (two in asymptomatic contacts of confirmed cases (Fenollar 2020(b); Shrestha 2020), and one involved staff screening, all of whom were RT-PCR-negative (PHE 2020(e)). The remaining antigen studies included samples from populations with mixed symptom status (8, 17%) or provided no information regarding symptom status (10, 21%). Of the 10 that provided no information, seven were laboratory-based studies providing no details of the settings from which the tested samples had been obtained, one included samples from a COVID-19 test centre, one was an outbreak investigation and in one the study setting could not be derived. There were no studies evaluating strategies of multiple tests.

A total of 13 studies provided accuracy data for people with no symptoms at the time of testing (3 studies exclusively in asymptomatic populations, and 10 studies providing subgroup data for people with no reported symptoms); one study provided only specificity data. Of the 12 datasets reporting both sensitivity and specificity, one (Alemany 2020), purportedly described preventive screening of the general population (although the reported prevalence of 24% is very high for such a scenario), one (Cerutti 2020), described targeted traveller screening, four (Billaud 2020; Fenollar 2020(b); Gupta 2020; Shrestha 2020), tested contacts of confirmed cases (one as part of an outbreak investigation) and the remaining six datasets were subgroups of samples from people presenting for routine testing. We identified one additional asymptomatic dataset in a report of several substudies but we did not include it as participants underwent antigen testing up to five days after a positive PCR test and it was not possible to determine the time point at which symptom status was recorded; it was also not possible to determine which 'substudy' the data related to (PHE 2020(d) [HCW tested]; PHE 2020(d) [Lab tested]).



Thirty-one of the 48 studies evaluating antigen tests reported results for SARS-CoV-2-confirmed samples above and below a Ct value from the reference standard RT-PCR. The median proportion of participants with 'high' viral load was 52% (IQR 35% to 60%). The most commonly used threshold was 24 or 25 Ct or less (n = 29 studies (or 36/58 test evaluations); 11 studies (15/58 test evaluations) reported results with at a threshold of between 31 and 33 Ct or less ; and 13 studies (13 evaluations) reported other thresholds including less than: 28 Ct (n = 3), 30 Ct (n = 5), 31 Ct (n = 3), or 35 Ct (n = 2)

Molecular tests

In contrast, studies evaluating molecular tests were mainly laboratory-based (20, 69%), with three (10%) including samples from participants presenting to emergency department or urgent care settings, two in hospital inpatients (7%), and four (14%) including samples from participants presenting in multiple settings. Twelve of the 29 studies (41%) reported included only samples from symptomatic patients, four reported mixed symptom status (10%) and 14 (48%) provided no information regarding symptom status. Of the 14 that provided no information, one was based in a hospital Accident and Emergency department, and the remaining 13 were laboratory-based studies, only three of which gave any details of the settings from which the tested samples had been obtained (three reported inclusion of samples from either inpatients and outpatients (n = 1), inpatients and ambulatory patients (n = 1) or inpatients and emergency department patients (n = 1) but did not provide the number of samples from each source). There were no studies evaluating strategies of multiple tests.

Five studies evaluating molecular assays, reported proportions with high viral load ranging from 33% to 80%, median 46%. All five studies reported results above and below a Ct value of 30.

Study design and reference standards

Table 1 shows a similar distribution of study designs between those evaluating antigen and molecular tests. Overall, 60% of studies (n = 46) used a 'single group' design to estimate both sensitivity and specificity and 22% (n = 17) used a 'two group' design with separate selection of RT-PCR-positive and RT-PCR-negative samples. In four studies (5%), the design could not be fully determined but probably deliberate separate sampling of RT-PCR-positive and RT-PCR-negative samples had been used.

Nine studies included only samples with confirmed SARS-CoV-2, thus only allowing estimation of sensitivity (six antigen and three molecular assay studies), and one study included only SARS-CoV-2-negative samples allowing estimation of specificity only. All studies defined the presence or absence of SARS-CoV-2 infection based on RT-PCR. Of the 68 studies that included SARS-CoV-2-negative samples, 63 (93%) required a single, negative PCR to confirm absence of infection and two (3%) required two negative PCR results. The remaining three studies used pre-pandemic samples (n = 2) or contemporaneous samples with other respiratory infections.

Thirty-three studies (43%), obtained paired swabs for index and reference standard, 39 (51%) used the same swab for point-of-care and RT-PCR (18 antigen and 21 molecular studies) and five studies used a mix of paired and same swabs (n = 1) or it was not possible to determine this information from the study report.

Index tests

Fifteen studies evaluated only one test, seven compared two or more tests in the same participants (four with two tests each, one with three tests and one each with four or five tests). In total the 77 studies that used respiratory samples reported on a total of 90 test evaluations. Appendix 13 provides details extracted from the manufacturer's instructions for use documents for all included tests.

Antigen tests

Forty-eight studies reported 58 evaluations of antigen tests; 41 of CGIAs, nine FIA, two alternative type of LFA using alkaline phosphatase-labelled antibodies, and six where assay type could not be determined. Studies evaluated 16 different commercially produced assays, as documented, with full assay identification details, in Appendix 13. One study reported the development of the Shenzhen Bioeasy assay (Diao 2020), but it is not clear whether the commercially available assay is identical to the one reported in the study or whether it has undergone further refinement. One study reported evaluating a Roche SARS-CoV-2 assay, which appears to be the SD Biosensor STANDARD Q (Schildgen 2020 [A]). Only 12 studies provided product codes for the tests evaluated (FIND 2020a; FIND 2020b; FIND 2020c (BR); FIND 2020c (CH); FIND 2020d (BR); FIND 2020d (DE); FIND 2020e (BR); FIND 2020e (DE); Gremmels 2020(a); Gremmels 2020(b); Porte 2020a; Weitzel 2020 [A]). The study reports or manufacturer IFUs for 11 assays reported targeting the nucleocapsid protein; this information was not reported for the Beijing Savant, Bionote, Biosynex, Liming Bio-Products, or RapiGEN Inc assays (Appendix 13). We were unable to identify any information for Beijing Savant, E25Bio or Liming Bio-Products assays online.

Multiple combinations of sample types and use of direct swab testing or swabs in viral transport medium or saline were reported across the studies (Table 1). Forty-one of 58 evaluations used nasopharyngeal (n = 30), oropharyngeal (n = 1) or nasal (n = 2) samples (type of nasal sample was not reported), or combinations of nasopharyngeal, nasal or oropharyngeal samples (n = 8; nasopharyngeal or nasal mid-turbinate in one, nasopharyngeal or combined naso- and oropharyngeal in two, naso- or oropharyngeal in two, and naso- or oropharyngeal or combined naso- and oropharyngeal samples in three. Thirteen evaluations used combined naso- and oropharyngeal samples for all participants, one used saliva samples and three evaluations (from one study) used bronchoalveolar lavage or throat wash samples. Of the six studies using nasal samples either alone (n = 2) or for at least some participants (n = 4), one reported that these were nares swabs, and the remaining five did not specify the type of nasal sample. Almost half of studies used direct swab testing (n = 28, 48%), 22 (38%) tested samples in viral transport medium, saline or other medium, and in 8 (14%) this information was not provided.

IFUs for five assays explicitly recommend against using any transport medium for swab testing (assays from Becton Dickinson, Bionote, Quidel and SD Biosensor; Appendix 13), one (Coris BioConcept) states that viral transport medium may be used, and the other nine do not mention use of transport medium, although two of the nine (from AAZ and Biosynex) imply that viral transport medium should not be used (using statements such as "use within one hour, stored in clean unused plastic tube"). We considered 29 of 58 antigen evaluations (50%) to be compliant with manufacturer



IFUs in terms of sample type, use of viral transport medium and time interval between collection and testing. Sixteen evaluations were not compliant with IFUs; nine used viral transport medium, four used freezing, four tested samples not listed on the IFUs, and in two testing was not always conducted within the one-hour time period specified in the IFU. For the remaining 13 evaluations either no IFU was available (n = 4), viral transport medium or saline was used but the IFU did not specifically address whether viral transport medium was recommended or not (n = 7), or insufficient detail was provided in the study.

Samples were collected by healthcare workers in 15 (26%) evaluations, by trained non-healthcare workers, such as firefighters or Ministry of Health employees in three (5%) evaluations, self-collected in six (10%) and the collection was not described in 34 evaluations (59%). Sample testing was conducted 'on-site' immediately or within one hour of collection in 21 (36%) evaluations by the same healthcare workers (n = 13), trained non-healthcare workers (n = 3) who collected the samples, or this information was not provided (n = 5). In the remaining 27 evaluations (47%), testing was conducted by laboratory staff (n = 12) or was inferred to be by laboratory staff (n = 15). For the latter group, the time interval between sample collection and testing was on receipt at the laboratory, some reporting delays of up to six hours.

Molecular tests

Collaboration.

Twenty-nine studies reported 32 evaluations of five different commercially available rapid molecular tests: 13 evaluating ID NOW (Abbott Laboratories), 15 evaluating Xpert Xpress (Cepheid Inc.), two of SAMBA II (Diagnostics for the Real World), and one evaluation each of Accula (Mesa Biotech Inc.) and COVID Nudge (DNANudge). None of the studies reported product codes for the tests evaluated. One study of Xpert Xpress used the 'research use only' (RUO) version of the test but reported that the RUO version contains the same reagents as the 'emergency use authorisation' (EUA) version. The RUO test allows the user to view the amplification curves for the RdRp gene as well as for the E-gene and N2 targets whereas the EUA version restricts the amplification curves to E and N2 only. ID NOW and SAMBA-II use isothermal techniques, Xpert Xpress and COVID Nudge are based on RT-PCR, and Accula is described as a PCR plus lateral flow assay.

Multiple combinations of sample types and use of direct swab testing or swabs in viral transport medium or saline were reported across the studies (Table 1). The sample types used included combined naso- and oropharyngeal samples (n = 2), nasopharyngeal samples alone (n = 16), nasal alone (n = 2), oropharyngeal samples alone (n = 1), or a combination of two or more of either nasopharyngeal or nasal or oropharyngeal samples (n = 8). One evaluation used throat saliva or lower respiratory tract specimens, one used saliva samples alone and one did not specify the sample type used. Of the six studies using nasal samples either alone (n = 2) or for at least some participants (n = 4), one reported

using nares swabs, and the remaining five did not specify the type of nasal sample used.

Eight evaluations (25%) reported direct swab testing in some (n = 1) or all (n = 7) samples, 18 (59%) used swabs in viral transport medium only (n = 12) or in viral transport medium or some other transport medium (n = 6), and six did not report whether they used any transport medium.

Sample collection was described in only three evaluations (9%) (Gibani 2020; Harrington 2020; Rhoads 2020; Table 1); the remaining studies did not describe sample collection but it is likely that samples were collected as part of routine care by healthcare workers. Sample testing was clearly described as conducted onsite by medical personnel or by laboratory personnel at local laboratories in one of the studies reporting sample collection (Harrington 2020), while a second implied testing as soon as possible after collection, possibly by the same healthcare worker (Gibani 2020). Four (12.5%) evaluations stated that laboratory staff carried out the tests. In 16 of the remaining 26 studies, testing by laboratory staff was inferred, based on delays between collection and testing of 18 hours to seven days (n = 10), or reported use of archived or frozen samples (n = 6). The remaining eight evaluations provided no useful information regarding who carried out the test (Assennato 2020; Dust 2020; Ghofrani 2020; Jin 2020; Jokela 2020; Moran 2020; Rhoads 2020; SoRelle 2020).

Two of the five manufacturers document IFU for samples stored in transport medium (Xpert Xpress and SAMBA II assays); two explicitly recommend against the use of viral transport medium (ID NOW and Accula), although at the time of the test evaluations some viral transport media were documented as acceptable for ID NOW; and one IFU does not mention the use of viral transport medium (COVID Nudge). Although immediate sample testing is preferred, all manufacturers document an acceptable period of refrigerated storage of between eight hours (COVID Nudge), and seven days with refrigeration (Xpert Xpress). See Appendix 13.

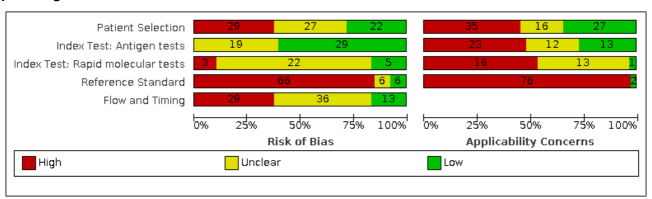
We considered only nine of 32 (28%) evaluations to be compliant with manufacturer IFUs in regard to sample type, use of viral transport medium and time interval between collection and testing. Sixteen evaluations were not compliant with IFUs; eight used viral transport medium, six used frozen samples, and two tested samples not listed on the IFUs. For the remaining seven evaluations, either the testing interval from sample collection was unclear (n = 5) or saline was used but the IFU did not specifically address whether this was recommended or not (n = 2).

Methodological quality of included studies

We report the overall methodological quality assessed using the QUADAS-2 tool for all included studies (n = 78) in Figure 2 (Whiting 2011). See Appendix 14 for separate summary plots by test method and for a plot of study-level ratings by quality domain. We explain how we reached these judgements in the Characteristics of included studies table.



Figure 2. Risk of bias and applicability concerns graph: review authors' judgements about each domain presented as percentages across included studies. Numbers in the bars indicate the number of studies



We considered whether the findings of individual studies were at risk of bias, and whether there were concerns that results might not apply to standard use of the tests. We did not judge any study at low risk of bias, although in 11 of 78 studies the only concern was that a single negative RT-PCR was used to confirm absence of COVID infection rather than the preferred two negative tests. All studies raised concerns regarding the applicability of their results, but in 13 of 78 studies the only concern was the reliance on only PCR to identify SARS-CoV-2 cases (and nine of these 13 are in common with the 11 using a single negative RT-PCR).

Participant selection

We judged 22 studies (28%) to be at low risk of bias, and 29 (37%) at high risk of bias because of deliberate sampling of participants based on the reference standard result (n = 25; 16 two-group studies and nine that only included samples with confirmed SARS-CoV-2 infection or absence of infection) or use of convenience sampling (n = 4). In 27 studies (35%) the risk of bias was unclear because of poor reporting of recruitment procedures or inclusion criteria (Figure 2).

A third (27/78) of studies were likely to have selected an appropriate patient group, recruiting participants from COVID-19 test centres, urgent care or emergency departments or identified through contact tracing. We had high concerns about the applicability of the selected participants in almost half of studies (35/78). Recruited participants were unlikely to be similar to those in whom the test would be used in clinical practice because of deliberate sampling (n = 25) or sample inclusion based on the availability of residual and sometimes frozen samples, or both (n = 22).

Index tests

Collaboration.

Poor reporting meant we could not clearly assess whether there was a risk of bias through performance of the index test in 41 (53%) studies. In general, antigen test studies were of a higher methodological standard for the index test domain compared to studies of molecular tests (Figure 2).

For antigen tests, we observed low risk of bias in 60% of studies (29/48). Risk of bias was unclear in the remaining studies because we could not judge whether interpretation of the index test was undertaken with knowledge of the reference standard result. For molecular tests, risk of bias was low in only 17% of studies (5/30). We observed high risk of bias in three studies (Moran 2020; Smithgall 2020 [A]; Wolters 2020) because they did not follow the manufacturer's prespecified threshold for the Xpert Xpress test (re-

testing of samples with presumptive positive results). Risk of bias was unclear in 73% (22/30) of studies because they did not report blinding to the reference standard (n = 22), six of these studies also did not report how they handled presumptive positive results on Xpert Xpress.

Fourteen studies (18%), including 13 antigen and one molecular test study, conducted testing as would be expected in practice (low concern regarding applicability). We had high concerns about applicability in half of all studies (39/78); 48% (23/48) of antigen and 57% (16/30) of molecular studies. Twenty-seven (11 antigen and 16 molecular) did not comply with manufacturers' IFU and a further 10 (all antigen studies), did not carry out tests as would occur in practice (i.e. trained, centralised laboratory staff carried out testing). In another two antigen studies concerns for applicability were high because tests were not available for purchase (Diao 2020; Nash 2020). Of the remaining 25 studies (12 antigen and 13 molecular) 16 conducted the test within the manufacturer IFU but none clearly described the setting for testing or personnel conducting the test.

Reference standards

Six studies were at low risk of bias for the reference standard. Although 12 used an appropriate reference standard, half (6/12) did not clearly implement blinding of the reference standard to the index test. High risk of bias (66/78) was present because studies did not use an adequate reference standard (Figure 2); they used either a single negative RT-PCR to define absence of SARS-CoV-2 infection (n = 64) or the index test formed part of a composite reference standard (n = 2).

A total of 36 studies reported blinded RT-PCR interpretation, two (with composite reference standard) did not implement blinding, and 40 (51%) provided insufficient information about blinding of the reference standard to the index test to judge risk of bias.

We judged 76 of the 78 studies to raise concerns about applicability (97%) because of defining the presence of SARS-CoV-2 infection based on a single RT-PCR-positive result. These studies will have excluded individuals who are RT-PCR-negative but have exposure and clinical features that meet the case definitions for COVID-19.

Flow and timing

Only 13 (17%) studies (all of antigen tests) were at low risk of bias for participant flow and timing (Figure 2). Twenty-nine (37%)



were at high risk of bias (19 antigen and 10 molecular) because of exclusion of samples following invalid index test results (n=23); delays between 'paired' swabs of up to three days (n=4), different reference standards used (n=3), or because they provided results on a per sample instead of per patient basis (n=2). These categories are not mutually exclusive.

We judged risk of bias unclear for 36 (46%) studies, primarily because of lack of clarity about participant inclusion and exclusion from analyses (n = 34), with no missing data or indeterminate test results reported and no Standards for Reporting Diagnostic Accuracy Studies (STARD)-style participant flow diagram and checklist (Bossuyt 2015), to fully report outcomes for all samples.

Conflicts of interest

In 27 studies all authors declared no conflicts of interest, although one study that reported the validation of a new test included a co-author affiliated to the test manufacturing company. Of these 27 studies, 19 were independent evaluations published by FIND or were from national reference laboratories. Twenty studies did not provide a conflict of interest statement, including 13 published studies and one study that reported affiliations to the test manufacturer. In the 12 remaining studies at least one author declared potential conflicts of interest in relation to the test.

Twenty-six studies provided no funding statement, 12 reported no funding sources to declare, and the remainder (n = 40) reported one or more funding sources.

Findings

Of the 78 included studies, eight reported evaluations of more than one test using the same samples and one reported evaluations of three tests using different samples (Table 1). To include all results from all tests in these analyses we have treated results from different tests of the same samples within a study as separate

data points, such that data are available on 91 test evaluations (58 evaluations of antigen tests in 48 studies and 33 evaluations of rapid molecular tests in 30 studies).

As previously stated, 77 of the 78 studies reported data for respiratory samples and one (Szymczak 2020), reported data for non-respiratory (faecal) samples. The main results, Tables and Figures focus on the respiratory samples, with Szymczak 2020 reported separately.

The results tables identify where estimates are based on multiple assessments of the same samples by including both the number of test evaluations and the number of studies. Nine datasets are from 'cases only' studies reporting only sensitivity estimates (six for antigen tests and three for molecular assays), and one antigen test evaluation is for 'non-COVID-19' cases reporting only specificity. Summary results are presented for studies providing both sensitivity and specificity data and then adding in the data from sensitivity- or specificity-only evaluations. The numbers of true positives, false positives, and total samples with and without confirmed SARS-CoV-2 infection are based on test result counts.

We present results for antigen tests overall and by subgroup in Table 2. Table 3 and Table 4 present results by test brand overall and by symptom status, and give results of sensitivity analyses restricting by compliance with manufacturer IFU. Forest plots of study data for the primary analysis are in Figure 3 and for subgroup analyses by symptom status and time after symptom onset are in Figure 4 and Figure 5. Appendix 15 provides forest plots for study data according to Ct value and study design. Individual plots by test brand are provided in Figure 6 for test brands with three or more evaluations and Figure 7 for test brands with one or two evaluations. Figure 8 shows data from studies comparing the accuracy of two or more antigen assays. Full identification details for studies of antigen-based assays are provided in Appendix 9 and Appendix 10.



Figure 3. Forest plot of studies evaluating antigen tests. BR: Brazil; CH: Switzerland; DE: Germany; HCW: healthcare worker; Lab: laboratory

Chd.	TD				C	CIti-it force oil	e	c likelin form older - lift in form old
Study	TP	FP	FN	TN		•		Sensitivity (95% CI)Specificity (95% CI)
Van der Moeren 2020(a)	16	2	1 5	332	Symptomatic	0.94 [0.71, 1.00]	0.99 [0.98, 1.00]	
Porte 2020a	77 30	0	2	45 31	Symptomatic	0.94 [0.86, 0.98]	1.00 [0.92, 1.00]	
Porte 2020b [A]	29	1	3	31	Symptomatic	0.94 [0.79, 0.99]	0.97 [0.84, 1.00]	
Porte 2020b [B] FIND 2020a	91	8	11	290	Symptomatic Symptomatic	0.91 [0.75, 0.98] 0.89 [0.82, 0.94]	0.97 [0.84, 1.00] 0.97 [0.95, 0.99]	4
FIND 2020a FIND 2020c (CH)	170	1	21	337	Symptomatic	0.89 [0.84, 0.93]	1.00 [0.98, 1.00]	
FIND 2020c (BR)	94	7	12	287	Symptomatic	0.89 [0.81, 0.94]	0.98 [0.95, 0.99]	
FIND 2020b	106	ó	18	411	Symptomatic	0.85 [0.78, 0.91]	1.00 [0.99, 1.00]	
Weitzel 2020 [D]	68	ŏ	12	31	Symptomatic	0.85 [0.75, 0.92]	1.00 [0.89, 1.00]	-
Albert 2020	43	ō	11	358	Symptomatic	0.80 [0.66, 0.89]	1.00 [0.99, 1.00]	
Fenollar 2020(a)	144	ō	38	0	Symptomatic	0.79 [0.72, 0.85]	Not estimable	
Van der Moeren 2020(b)	98	ō	27	ō	Symptomatic	0.78 [0.70, 0.85]	Not estimable	
Young 2020	29	1	9	212	Symptomatic	0.76 [0.60, 0.89]	1.00 [0.97, 1.00]	
FIND 2020e (BR)	87	4	30	355	Symptomatic	0.74 [0.65, 0.82]	0.99 [0.97, 1.00]	
Weitzel 2020 [A]	49	0	30	30	Symptomatic	0.62 [0.50, 0.73]	1.00 [0.88, 1.00]	—
Fourati 2020 [E]	182	0	113	337	Symptomatic	0.62 [0.56, 0.67]	1.00 [0.99, 1.00]	-
Fourati 2020 [B]	175	23	116	314	Symptomatic	0.60 [0.54, 0.66]	0.93 [0.90, 0.96]	
Fourati 2020 [D]	177	0	120	337	Symptomatic	0.60 [0.54, 0.65]	1.00 [0.99, 1.00]	
Fourati 2020 [C]	163	0	132	337	Symptomatic	0.55 [0.49, 0.61]	1.00 [0.99, 1.00]	
PHE 2020(a)	95	0	83	940	Symptomatic	0.53 [0.46, 0.61]	1.00 [1.00, 1.00]	
Fourati 2020 [A]	103	0	189	337	Symptomatic	0.35 [0.30, 0.41]	1.00 [0.99, 1.00]	
Veyrenche 2020	13	0	32	20	Symptomatic	0.29 [0.16, 0.44]	1.00 [0.83, 1.00]	
Weitzel 2020 [C]	13	0	65	31	Symptomatic	0.17 [0.09, 0.27]	1.00 [0.89, 1.00]	-
Weitzel 2020 [B]	0	1	9	9	Symptomatic	0.00 [0.00, 0.34]	0.90 [0.55, 1.00]	-
Gremmels 2020(b)	51	0	12	145	Not reported	0.81 [0.69, 0.90]	1.00 [0.97, 1.00]	
Tak ed a 2020	50	0	12	100	Not reported	0.81 [0.69, 0.90]	1.00 [0.96, 1.00]	
Nash 2020	80	8	20	82	Not reported	0.80 [0.71, 0.87]	0.91 [0.83, 0.96]	
Diao 2020	141	0	67	31	Not reported	0.68 [0.61, 0.74]	1.00 [0.89, 1.00]	+ -•
Mertens 2020	76	1	56	195	Not reported	0.58 [0.49, 0.66]	0.99 [0.97, 1.00]	
Lambert-Niclot 2020	47	0	47	44	Not reported	0.50 [0.40, 0.60]	1.00 [0.92, 1.00]	
Liotti 2020	49	4	55	251	Not reported	0.47 [0.37, 0.57]	0.98 [0.96, 1.00]	
Mak 2020	51	0		0	Not reported	0.32 [0.25, 0.40]	Not estimable	• •
Blairon 2020	9	0	21	26	Not reported	0.30 [0.15, 0.49]	1.00 [0.87, 1.00]	
PHE 2020(b)	13	0 5	33	105	Not reported	0.28 [0.16, 0.43]	1.00 [0.97, 1.00]	
Alemany 2020	872		79 5	450	Mixed	0.92 [0.90, 0.93]	0.99 [0.97, 1.00]	
Schildgen 2020 [C] Gupta 2020	37 6 3	25 1	14	6 252	Mixed Mixed	0.88 [0.74, 0.96]	0.19 [0.07, 0.37] 1.00 [0.98, 1.00]	
Linares 2020	44	0	16	195	Mixed	0.82 [0.71, 0.90] 0.73 [0.60, 0.84]	1.00 [0.98, 1.00]	
Cerutti 2020	77	0	32	221	Mixed	0.71 [0.61, 0.79]	1.00 [0.98, 1.00]	
Billaud 2020	53	5	46	358	Mixed	0.54 [0.43, 0.64]	0.99 [0.97, 1.00]	
FIND 2020e (DE)	13	0		1214	Mixed	0.52 [0.31, 0.72]	1.00 [1.00, 1.00]	
Schildgen 2020 [B]	21	7	21	24	Mixed	0.50 [0.34, 0.66]	0.77 [0.59, 0.90]	
Schildgen 2020 [A]	14	4	28	27	Mixed	0.33 [0.20, 0.50]	0.87 [0.70, 0.96]	
Scohy 2020	32	Ö	74	42	Mixed	0.30 [0.22, 0.40]	1.00 [0.92, 1.00]	
Courtellemont 2020	97	20	4		Mainly symptomatic	0.96 [0.90, 0.99]	0.86 [0.80, 0.91]	
PHE 2020(d) [Lab tested]	156	0	42	0	Mainly symptomatic	0.79 [0.72, 0.84]	Not estimable	
FIND 2020d (BR)	93	7	27		Mainly symptomatic	0.78 [0.69, 0.85]	0.98 [0.96, 0.99]	-
Kruger 2020(c)	36	9	11		Mainly symptomatic	0.77 [0.62, 0.88]	0.99 [0.99, 1.00]	
Gremmels 2020(a)	101	0	38		Mainly symptomatic	0.73 [0.64, 0.80]	1.00 [1.00, 1.00]	+
PHE 2020(d) [HCW tested]	156	0	67		Mainly symptomatic	0.70 [0.63, 0.76]	Not estimable	-
FIND 2020d (DE)	27	20	12	617	Mainly symptomatic	0.69 [0.52, 0.83]	0.97 [0.95, 0.98]	
Kruger 2020(a)	10	49	5		Mainly symptomatic	0.67 [0.38, 0.88]	0.93 [0.91, 0.95]	
PHE 2020(c) [non-HCW tested]	214	5				0.58 [0.52, 0.63]	1.00 [0.99, 1.00]	-
Kruger 2020(b)	4	17	4		Mainly symptomatic	0.50 [0.16, 0.84]	0.96 [0.93, 0.98]	
Nagura-Ikeda 2020	12	0	91	0	Mainly symptomatic	0.12 [0.06, 0.19]	Not estimable	
PHE 2020(e)	0	1	0	5 37	Asym pto matic	Not estimable	1.00 [0.99, 1.00]	
Shrestha 2020	40	0	7	66	Asym pto matic	0.85 [0.72, 0.94]	1.00 [0.95, 1.00]	
Fenollar 2020(b)	10	7	12	130	Asymptomatic	0.45 [0.24, 0.68]	0.95 [0.90, 0.98]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
								0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1



Figure 4. Forest plot of data for antigen tests according to symptom status. A&E: accident and emergency; BR: Brazil; CH: Switzerland; DE: Germany; HCW: healthcare worker; Lab: laboratory

Antigen tests - symptomatic TP FP Setting Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) FΝ Study TN Van der Moeren 2020(a) COVID-19 test centre 0.94 [0.71, 1.00] 0.99 [0.98, 1.00] 16 332 0.94 [0.79, 0.99] 0.91 [0.75, 0.98] 0.97 [0.84, 1.00] 0.97 [0.84, 1.00] Porte 2020b [A] 30 31 COVID-19 test centre Porte 2020b (B) 29 31 COVID-19 test centre 0.89 [0.82, 0.94] 0.97 [0.95, 0.99] FIND 2020a 91 11 290 COVID-19 test centre FIND 2020c (CH) FIND 2020c (BR) 170 21 337 COVID-19 test centre 0.89 [0.84, 0.93] 1.00 [0.98, 1.00] COVID-19 test centre 94 0.98 [0.95, 0.99] 12 287 0.89 [0.81, 0.94] FIND 2020b 0 411 COVID-19 test centre 0.85 [0.78, 0.91] 1.00 [0.99, 1.00] 106 18 Kruger 2020(c) 32 972 COVID-19 test centre 0.82 [0.66, 0.92] 0.99 [0.99, 1.00] Albert 2020 43 0.80 [0.66, 0.89] 11 358 COVID-19 test centre 1.00 [0.99, 1.00] 38 Fenollar 2020(a) 144 0 COVID-19 test centre 0.79 [0.72, 0.85] Not estimable 0 PHE 2020(d) [Lab tested] 42 0.79 [0.72, 0.84] 156 0 COVID-19 test centre Not estimable Van der Moeren 2020(b) 0.78 [0.70, 0.85] 98 27 COVID-19 test centre Not estimable 0.98 [0.96, 0.99] 0.99 [0.97, 1.00] FIND 2020d (BR) 93 27 326 COVID-19 test centre 0.78 [0.69, 0.85] FIND 2020e (BR) 30 COVID-19 test centre 0.74 [0.65, 0.82] 87 355 Gremmels 2020(a) 99 0 37 1185 COVID-19 test centre 0.73 [0.65, 0.80] 1.00 [1.00, 1.00] PHE 2020(d) [HCW tested] 156 0 67 0 COVID-19 test centre 0.70 [0.63, 0.76] Not estimable FIND 2020d (DE) 27 20 12 617 COVID-19 test centre 0.69 [0.52, 0.83] 0.97 [0.95, 0.98] Kruger 2020(a) PHE 2020(c) [non-HCW tested] 49 5 0.67 [0.38, 0.88] 0.58 [0.52, 0.63] 10 5 663 COVID-19 test centre 0.93 [0.91, 0.95] 158 214 1299 1.00 [0.99, 1.00] COVID-19 test centre Billaud 2020 40 34 69 Contacts 0.54 [0.42, 0.66] 0.95 [0.87, 0.98] Porte 2020a 77 0 5 45 Hospital A&E 0.94 [0.86, 0.98] 0.85 [0.75, 0.92] 1.00 [0.92, 1.00] 1.00 [0.89, 1.00] 12 Weitzel 2020 [D] 68 0 31 Hospital A&E Linares 2020 39 11 133 Hospital A&E 0.78 [0.64, 0.88] 1.00 [0.97, 1.00] Cerutti 2020 75 0 29 81 Hospital A&E 0.72 [0.62, 0.80] 1.00 [0.96, 1.00] Hospital A&E Weitzel 2020 [A 49 30 30 0.62 [0.50, 0.73] 1.00 [0.88, 1.00] Weitzel 2020 [C] 13 0 65 31 Hospital A&E 0.17 [0.09, 0.27] 1.00 [0.89, 1.00] 9 0.00 [0.00, 0.34] Weitzel 2020 [B] 0 Hospital A&E 0.90 [0.55, 1.00] Hospital in-patient PHE 2020(a) 95 0 83 940 0.53 [0.46, 0.61] 1.00 [1.00, 1.00] Veyrenche 2020 13 0 32 20 Hospital in-patient 0.29 [0.16, 0.44] 1.00 [0.83, 1.00] Fourati 2020 [E] 182 0 113 337 0.62 [0.56, 0.67] 1.00 [0.99, 1.00] Laboratory-based Fourati 2020 [B] Fourati 2020 [D] 175 177 Laboratory-based 23 116 314 0.60 [0.54, 0.66] 0.93 [0.90, 0.96] 0 0.60 [0.54, 0.65] 120 337 Laboratory-based 1.00 [0.99, 1.00] 0.55 [0.49, 0.61] Fourati 2020 [C] 163 132 337 Laboratory-based 1.00 [0.99, 1.00] Fourati 2020 [A] 103 Λ 189 337 Laboratory-based 0.35 (0.30, 0.41) 1.00 [0.99, 1.00] 25 ō 0.32 [0.22, 0.44] Scohy 2020 1.00 [0.66, 1.00] 52 9 Laboratory-based Courtellemont 2020 97 20 4 127 0.96 [0.90, 0.99] 0.86 [0.80, 0.91] Mixed Young 2020 Nagura-Ikeda 2020 29 1 9 212 Mixed 0.76 [0.60, 0.89] 1.00 [0.97, 1.00] ō 0.11 [0.06, 0.20] Not estimable 10 78 Mixed Schildgen 2020 [C] 10 12 0 1 Unclear 1.00 [0.69, 1.00] 0.08 [0.00, 0.36] Alemany 2020 0 31 27 0.93 [0.90, 0.95] 1.00 [0.87, 1.00] 388 Unclear Schildgen 2020 [B] 11 0.40 [0.12, 0.74] 0.85 [0.55, 0.98] Unclear Schildgen 2020 [A] 3 10 Unclear 0.30 [0.07, 0.65] 0.77 [0.46, 0.95] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Antigen tests - asymptomatic Study FP TN Setting Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Gremmels 2020(a) 0 34 COVID-19 test centre 0.67 [0.09, 0.99] 1.00 [0.90, 1.00] 1 7 1.00 [0.95, 1.00] Shrestha 2020 40 0 66 0.85 [0.72, 0.94] Contacts Gupta 2020 Contacts 0.69 [0.39, 0.91] 0.99 [0.95, 1.00] Billaud 2020 13 1 7 12 289 Contacts 0.52 (0.31, 0.72) 1.00 (0.98, 1.00) 12 0.45 [0.24, 0.68] Fenollar 2020(b) 130 0.95 [0.90, 0.98] 10 Contacts Linares 2020 0 5 Hospital A&E 0.50 [0.19, 0.81] 1.00 [0.94, 1.00] Scohy 2020 4 0 10 31 Laboratory-based 0.29 [0.08, 0.58] 1.00 [0.89, 1.00] 0.13 [0.02, 0.40] Nagura-Ikeda 2020 13 Mixed Not estimable Screening Alemany 2020 Cerutti 2020 0.99 [0.97, 1.00] 93 5 24 365 0.79 [0.71, 0.86] 0 3 140 0.40 [0.05, 0.85] Targeted screening Schildgen 2020 [C] 11 12 2 0.85 [0.55, 0.98] 0.14 [0.02, 0.43] Unclea 8 9 Schildgen 2020 [B] 4 1.0 Unclear 0.38 [0.14, 0.68] 0.71 [0.42, 0.92] Schildgen 2020 [A] 0.31 [0.09, 0.61] 0.93 [0.66, 1.00] Unclear 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Antigen tests - mixed symptoms or not reported Setting Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Study TP FP ΕN TN Gupta 2020 COVID-19 test centre 0.82 [0.71, 0.90] 1.00 [0.98, 1.00] 252 63 14 COVID-19 test centre Gremmels 2020(b) 0 145 0.81 [0.69, 0.90] 1.00 [0.97, 1.00] 51 FIND 2020e (DE) 13 0 12 1214 COVID-19 test centre 0.52 [0.31, 0.72] 1.00 [1.00, 1.00] 0.94 [0.92, 0.96] 1.00 [0.94, 1.00] Alemany 2020 391 24 Contacts 58 Billaud 2020 53 46 358 Contacts 0.54 [0.43, 0.64] 0.99 [0.97, 1.00] PHE 2020(b) 0 33 105 0.28 [0.16, 0.43] 1.00 [0.97, 1.00] 13 Contacts 0.73 [0.60, 0.84] 1.00 [0.98, 1.00] Linares 2020 16 Hospital A&E 0.81 [0.69, 0.90] 0.80 [0.71, 0.87] Takeda 2020 50 ٥ 12 100 Laboratory-based 1.00 (0.96, 1.00) Nash 2020 80 8 20 82 Laboratory-based 0.91 [0.83, 0.96] Mertens 2020 76 56 195 Laboratory-based 0.58 [0.49, 0.66] 0.99 [0.97, 1.00] Lambert-Niclot 2020 47 0 47 44 Laboratory-based 0.50 [0.40, 0.60] 1.00 [0.92, 1.00] 49 0.47 [0.37, 0.57] 0.98 [0.96, 1.00] Liotti 2020 251 Laboratory-based Mak 2020 51 ٥ 109 0 Laboratory-based 0.32 [0.25, 0.40] 0.30 [0.22, 0.40] Not estimable 1.00 [0.92, 1.00] 32 74 42 Scohy 2020 Laboratory-based 0 9 0 21 26 0.30 [0.15, 0.49] 1.00 [0.87, 1.00] Blairon 2020 Laboratory-based Cerutti 2020 77 0 32 221 Mixed 0.71 [0.61, 0.79] 1.00 [0.98, 1.00] 37 0.88 [0.74, 0.96] Schildgen 2020 [C] Unclear 0.19 [0.07, 0.37] 25 31

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Figure 4. (Continued)

Cerutti 2020	77	0	32	221	Mixed	0.71 [0.61, 0.79]	1.00 [0.98, 1.00]
Schildgen 2020 [C]	37	25	5	6	Unclear	0.88 [0.74, 0.96]	0.19 [0.07, 0.37]
Diac 2020	141	0	67	31	Unclear	0.68 [0.61, 0.74]	1.00 [0.89, 1.00]
Schildgen 2020 [B]	21	7	21	24	Unclear	0.50 [0.34, 0.66]	0.77 [0.59, 0.90]
Schildgen 2020 [A]	14	4	28	27	Unclear	0.33 [0.20, 0.50]	0.87 [0.70, 0.96]



Figure 5. Forest plot of antigen test evaluations by week post symptom onset (pso). A&E: accident and emergency; Ag: antigen; BR: Brazil; CH: Switzerland; DE: Germany

Antigen tests - week 1 after symptom onset

Study	TP	FP	FN	TN	Setting	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Porte 2020b [A]	30	1	2	31	COVID-19 test centre	0.94 [0.79, 0.99]	0.97 [0.84, 1.00]	
FIND 2020a	83	0	7	0	COVID-19 test centre	0.92 [0.85, 0.97]	Not estimable	-
FIND 2020c (BR)	88	0	9	0	COVID-19 test centre	0.91 [0.83, 0.96]	Not estimable	
Porte 2020b [B]	29	1	3	31	COVID-19 test centre	0.91 [0.75, 0.98]	0.97 [0.84, 1.00]	→ →
FIND 2020c (CH)	158	0	18	0	COVID-19 test centre	0.90 [0.84, 0.94]	Not estimable	•
Van der Moeren 2020(b)	59	0	7	0	COVID-19 test centre	0.89 [0.79, 0.96]	Not estimable	
Gu p ta 2020	49	0	8	134	COVID-19 test centre	0.86 [0.74, 0.94]	1.00 [0.97, 1.00]	
FIND 2020b	95	0	16	0	COVID-19 test centre	0.86 [0.78, 0.92]	Not estimable	-
FIND 2020d (DE)	26	0	6	0	COVID-19 test centre	0.81 [0.64, 0.93]	Not estimable	
FIND 2020d (BR)	80	0	20	0	COVID-19 test centre	0.80 [0.71, 0.87]	Not estimable	-
Kruger 2020(c)	28	7	7	907	COVID-19 test centre	0.80 [0.63, 0.92]	0.99 [0.98, 1.00]	
Albert 2020	43	0	11	358	COVID-19 test centre	0.80 [0.66, 0.89]	1.00 [0.99, 1.00]	
FIND 2020e (BR)	76	0	22	0	COVID-19 test centre	0.78 [0.68, 0.85]	Not estimable	-
FIND 2020e (DE)	10	0	3	0	COVID-19 test centre	0.77 [0.46, 0.95]	Not estimable	
Gremmels 2020(a)	75	0	26	846	COVID-19 test centre	0.74 [0.65, 0.82]	1.00 [1.00, 1.00]	-
Kruger 2020(b)	3	0	4	0	COVID-19 test centre	0.43 [0.10, 0.82]	Not estimable	
Porte 2020a	72	0	4	42	Hospital A&E	0.95 [0.87, 0.99]	1.00 [0.92, 1.00]	-
Linares 2020	32	0	5	846	Hospital A&E	0.86 [0.71, 0.95]	1.00 [1.00, 1.00]	
Veyrenche 2020	9	1	13	31	Hospital in-patient	0.41 [0.21, 0.64]	0.97 [0.84, 1.00]	─
Fourati 2020 [E]	142	0	58	0	Laboratory-based	0.71 [0.64, 0.77]	Not estimable	+
Fourati 2020 [B]	141	0	58	0	Laboratory-based	0.71 [0.64, 0.77]	Not estimable	-
Fourati 2020 [D]	137	0	63	0	Laboratory-based	0.69 [0.62, 0.75]	Not estimable	-
Fourati 2020 [C]	131	0	69	0	Laboratory-based	0.66 [0.58, 0.72]	Not estimable	-
Fourati 2020 [A]	90	0	109	0	Laboratory-based	0.45 [0.38, 0.52]	Not estimable	-
Young 2020	29	1	9	212	Mixed	0.76 [0.60, 0.89]	1.00 [0.97, 1.00]	
Nagura-Ikeda 2020	7	0	41	0	Mixed	0.15 [0.06, 0.28]	Not estimable	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Antigen tests - week 2 after symptom onset

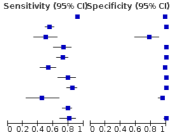
Study	TP	FP	FN	TN	Setting	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Kruger 2020(b)	1	0	0	0	COVID-19 test centre	1.00 [0.03, 1.00]	Not estimable	
Kruger 2020(c)	4	0	0	54	COVID-19 test centre	1.00 [0.40, 1.00]	1.00 [0.93, 1.00]	
FIND 2020b	11	0	2	0	COVID-19 test centre	0.85 [0.55, 0.98]	Not estimable	
FIND 2020c (CH)	12	0	3	0	COVID-19 test centre	0.80 [0.52, 0.96]	Not estimable	
Gu p ta 2020	5	0	2	5	COVID-19 test centre	0.71 [0.29, 0.96]	1.00 [0.48, 1.00]	
FIND 2020c (BR)	6	0	3	0	COVID-19 test centre	0.67 [0.30, 0.93]	Not estimable	
FIND 2020a	8	0	4	0	COVID-19 test centre	0.67 [0.35, 0.90]	Not estimable	
Van der Moeren 2020(b)	38	0	19	0	COVID-19 test centre	0.67 [0.53, 0.79]	Not estimable	
FIND 2020d (BR)	13	0	7	0	COVID-19 test centre	0.65 [0.41, 0.85]	Not estimable	
FIND 2020e (BR)	11	0	8	0	COVID-19 test centre	0.58 [0.33, 0.80]	Not estimable	
Gremmels 2020(a)	5	0	5	181	COVID-19 test centre	0.50 [0.19, 0.81]	1.00 [0.98, 1.00]	
FIND 2020e (DE)	3	0	9	0	COVID-19 test centre	0.25 [0.05, 0.57]	Not estimable	
FIND 2020d (DE)	1	0	6	0	COVID-19 test centre	0.14 [0.00, 0.58]	Not estimable	
Porte 2020a	4	0	1	3	Hospital A&E	0.80 [0.28, 0.99]	1.00 [0.29, 1.00]	
Linares 2020	7	0	6	0	Hospital A&E	0.54 [0.25, 0.81]	Not estimable	
Veyrenche 2020	4	0	10	0	Hospital in-patient	0.29 [0.08, 0.58]	Not estimable	
Fourati 2020 [D]	38	0	51	0	Laboratory-based	0.43 [0.32, 0.54]	Not estimable	
Fourati 2020 [E]	36	0	51	0	Laboratory-based	0.41 [0.31, 0.52]	Not estimable	-
Fourati 2020 [B]	32	0	53	0	Laboratory-based	0.38 [0.27, 0.49]	Not estimable	-
Fourati 2020 [C]	30	0	57	0	Laboratory-based	0.34 [0.25, 0.45]	Not estimable	
Fourati 2020 [A]	13	0	73	0	Laboratory-based	0.15 [0.08, 0.24]	Not estimable	-
Nagura-Ikeda 2020	3	0	37	0	Mixed	0.07 [0.02, 0.20]	Not estimable	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1



Figure 6. Forest plot by test brand for assays with ≥ 3 evaluations. BR: Brazil; CGIA: colloidal-gold immunoassay; CH: Switzerland; DE: Germany; FIA: fluorescent immunoassay; HCW: healthcare worker; IFU: instructions for use; Lab: laboratory; LFA: lateral flow assay

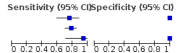
Abbott -	Panbio	Covid-19 Ag	(CGIA)
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Study	TP	FP	FN	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Alemany 2020	872	5	79	450	No	0.92 [0.90, 0.93]	0.99 [0.97, 1.00]
Fourati 2020 [C]	163	0	132	337	No	0.55 [0.49, 0.61]	1.00 [0.99, 1.00]
Schildgen 2020 [B]	21	- 7	21	24	No	0.50 [0.34, 0.66]	0.77 [0.59, 0.90]
Linares 2020	44	0	16	195	Unclear	0.73 [0.60, 0.84]	1.00 [0.98, 1.00]
Gremmels 2020(a)	101	0	38	1228	Unclear	0.73 [0.64, 0.80]	1.00 [1.00, 1.00]
Billaud 2020	53	5	46	358	Yes	0.54 [0.43, 0.64]	0.99 [0.97, 1.00]
Albert 2020	43	0	11	358	Yes	0.80 [0.66, 0.89]	1.00 [0.99, 1.00]
FIND 2020b	106	0	18	411	Y e s	0.85 [0.78, 0.91]	1.00 [0.99, 1.00]
Fenollar 2020(b)	10	7	12	130	Yes	0.45 [0.24, 0.68]	0.95 [0.90, 0.98]
Fenollar 2020(a)	144	0	38	0	Yes	0.79 [0.72, 0.85]	Not estimable
Gremmels 2020(b)	51	0	12	145	Yes	0.81 [0.69, 0.90]	1.00 [0.97, 1.00]



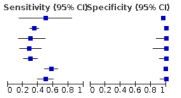
Becton Dickinson - BD Veritor (LFA - method not specified)

Study	ΤP	FΡ	FΝ	IN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Young 2020	29	1	9	212	No	0.76 [0.60, 0.89]	1.00 [0.97, 1.00]
Van der Moeren 2020(b)	98	0	27	0	No	0.78 [0.70, 0.85]	Not estimable
Van der Moeren 2020(a)	16	2	1	332	No	0.94 [0.71, 1.00]	0.99 [0.98, 1.00]



Coris Bioconcept - COVID-19 Ag Respi-Strip (CGIA)

Study	TP	FP	FΝ	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Kruger 2020(b)	4	17	4	392	Yes	0.50 [0.16, 0.84]	0.96 [0.93, 0.98]
Fourati 2020 [A]	103	0	189	337	Y e s	0.35 [0.30, 0.41]	1.00 [0.99, 1.00]
Blairon 2020	9	0	21	26	Yes	0.30 [0.15, 0.49]	1.00 [0.87, 1.00]
Veyrenche 2020	13	0	32	20	Yes	0.29 [0.16, 0.44]	1.00 [0.83, 1.00]
Scohy 2020	32	0	74	42	Y e s	0.30 [0.22, 0.40]	1.00 [0.92, 1.00]
Mertens 2020	76	1	56	195	Yes	0.58 [0.49, 0.66]	0.99 [0.97, 1.00]
Lambert-Niclot 2020	47	0	47	44	Yes	0.50 [0.40, 0.60]	1.00 [0.92, 1.00]

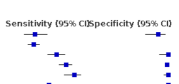


Innova Medical Group - Innova SARS-CoV-2 Ag (CGIA)

Study	TP	FP	FN	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95%
PHE 2020(b)	13	0	33	105	Unclear	0.28 [0.16, 0.43]	1.00 [0.97, 1.00]	-
PHE 2020(a)	95	0	83	940	Unclear	0.53 [0.46, 0.61]	1.00 [1.00, 1.00]	-
PHE 2020(d) [HCW tested]	156	0	67	0	Yes	0.70 [0.63, 0.76]	Not estimable	-
PHE 2020(c) [non-HCW tested]	214	5	158	1299	Y e s	0.58 [0.52, 0.63]	1.00 [0.99, 1.00]	•
PHE 2020(e)	0	1	0	537	Y e s	Not estimable	1.00 [0.99, 1.00]	
PHE 2020(d) [Lab tested]	156	0	42	0	Y e s	0.79 [0.72, 0.84]	Not estimable	
								0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8

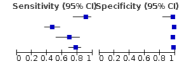
RapiGEN - BIOCREDIT COVID-19 Ag (CGIA)

Study	TP	FP	FN	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Schildgen 2020 [A]	14	4	28	27	No	0.33 [0.20, 0.50]	0.87 [0.70, 0.96]
Mak 2020	51	0	109	0	No	0.32 [0.25, 0.40]	Not estimable
Weitzel 2020 [A]	49	0	30	30	No	0.62 [0.50, 0.73]	1.00 [0.88, 1.00]
FIND 2020e (BR)	87	4	30	355	Yes	0.74 [0.65, 0.82]	0.99 [0.97, 1.00]
Shrestha 2020	40	0	7	66	Yes	0.85 [0.72, 0.94]	1.00 [0.95, 1.00]
FIND 2020e (DE)	13	0	12	1214	Yes	0.52 [0.31, 0.72]	1.00 [1.00, 1.00]



SD Biosensor - STANDARD F COVID-19 Ag (FIA)

Study	TP	FP	FN	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Porte 2020b [B]	29	1	3	31	No	0.91 [0.75, 0.98]	0.97 [0.84, 1.00]
Liotti 2020	49	4	55	251	Unclear	0.47 [0.37, 0.57]	0.98 [0.96, 1.00]
FIND 2020d (DE)	27	20	12	617	Yes	0.69 [0.52, 0.83]	0.97 [0.95, 0.98]
FIND 2020d (BR)	93	7	27	326	Yes	0.78 [0.69, 0.85]	0.98 (0.96, 0.99)



0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

SD Biosensor - STANDARD Q COVID-19 Ag (CGIA)

Study	TP	FP	FN	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Cerutti 2020	77	0	32	221	No	0.71 [0.61, 0.79]	1.00 [0.98, 1.00]
Fourati 2020 [B]	175	23	116	314	No	0.60 [0.54, 0.66]	0.93 [0.90, 0.96]
Gupta 2020	63	1	14	252	Yes	0.82 [0.71, 0.90]	1.00 [0.98, 1.00]
FIND 2020c (CH)	170	1	21	337	Yes	0.89 [0.84, 0.93]	1.00 [0.98, 1.00]
FIND 2020c (BR)	94	7	12	287	Yes	0.89 [0.81, 0.94]	0.98 [0.95, 0.99]
Kruger 2020(c)	36	9	11	1207	Y e s	0.77 [0.62, 0.88]	0.99 [0.99, 1.00]



Shenzhen Bioeasy Biotech - 2019-nCoV Ag (FIA)

Study	TP	FP	FΝ	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)
Porte 2020a	77	0	5	45	Unclear	0.94 [0.86, 0.98]	1.00 [0.92, 1.00]
Weitzel 2020 [D]	68	0	12	31	Unclear	0.85 [0.75, 0.92]	1.00 [0.89, 1.00]
Kruger 2020(a)	10	49	5	663	Yes	0.67 [0.38, 0.88]	0.93 [0.91, 0.95]



0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1



Figure 7. Forest plot by test brand for assays with < 3 evaluations; CGIA: colloidal-gold immunoassay; FIA: fluorescent immunoassay; IFU: instructions for use; LFA: lateral flow assay

AAZ - COVID-VIRO (CGIA) Study TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Fourati 2020 [E] 182 0 113 337 0.62 [0.56, 0.67] 1.00 [0.99, 1.00] Unclear Courtellemont 2020 97 20 0.96 [0.90, 0.99] 0.86 [0.80, 0.91] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 BIONOTE - NowCheck COVID-19 Ag (LFA - method not specified) Sensitivity (95% CI)Specificity (95% CI) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Study FIND 2020a 91 8 11 290 0.89 [0.82, 0.94] 0 0.2 0.4 0.6 0.8 1 E25Bio - DART (NP) (CGIA) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Study Nash 2020 80 8 20 82 Unclear 0.80 [0.71, 0.87] 0.91 [0.83, 0.96] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Fujirebio - ESPLINE SARS-CoV-2 [LFA(ALP)] TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Nagura-Ikeda 2020 0.12 [0.06, 0.19] Not estimable 12 0 91 0 No Takeda 2020 0 12 100 Unclear 0.81 [0.69, 0.90] 1.00 [0.96, 1.00] 50 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Inhouse (Bioeasy co-author) - n/a (FIA) Sensitivity (95% CI)Specificity (95% CI) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Study Diao 2020 141 0 67 31 Unclear 0.68 [0.61, 0.74] 1.00 [0.89, 1.00] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Liming Bio-Products - StrongStep® COVID-19 Ag (CGIA) Sensitivity (95% CI)Specificity (95% CI) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Weitzel 2020 [B] 0 1 9 9 Unclear 0.00 [0.00, 0.34] 0.90 [0.55, 1.00] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Quidel Corporation - SOFIA SARS Antigen (FIA) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Porte 2020b [A] 30 1 2 31 0.94 [0.79, 0.99] 0.97 [0.84, 1.00] No 0 0.2 0.4 0.6 0.8 1 Roche - SARS-CoV-2 (LFA - method not specified) Sensitivity (95% CI)Specificity (95% CI) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Schildgen 2020 [C] 37 25 5 6 No 0.88 [0.74, 0.96] 0.19 [0.07, 0.37] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Savant Biotech - Huaketai SARS-CoV-2 N Protein (LFA - method not specified) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI)Specificity (95% CI) Weitzel 2020 [C] 13 0 65 31 Unclear 0.17 [0.09, 0.27] 1.00 [0.89, 1.00] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 8. Forest plot of studies reporting comparative data. CGIA: colloidal-gold immunoassay; FIA: fluorescent immunoassay; LFA: lateral flow assay; nos: not otherwise specified

Study	TP	FP	FN	TN	Test method	Test	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Fourati 2020 [A]	103	0	189	337	CGIA	Coris Bioconcept - COVID-19 Ag Respi-Strip	0.35 [0.30, 0.41]	1.00 [0.99, 1.00]	
Fourati 2020 [B]	175	23	116	314	CGIA	SD Biosensor - STANDARD Q COVID-19 Ag	0.60 [0.54, 0.66]	0.93 [0.90, 0.96]	
Fourati 2020 [C]	163	0	132	337	CGIA	Abbott - Panbio Covid-19 Ag	0.55 [0.49, 0.61]	1.00 [0.99, 1.00]	- ·
Fourati 2020 [D]	177	0	120	337	CGIA	Biosynex - Biosynex COVID-19 Ag BSS	0.60 [0.54, 0.65]	1.00 [0.99, 1.00]	-
Fourati 2020 [E]	182	0	113	337	CGIA	AAZ - COVID-VIRO	0.62 [0.56, 0.67]	1.00 [0.99, 1.00]	
Porte 2020b [A]	30	1	2	31	FIA	Quidel Corporation - SOFIA SARS Antigen	0.94 [0.79, 0.99]	0.97 [0.84, 1.00]	→ →
Porte 2020b [B]	29	1	3	31	FIA	SD Biosensor - STANDARD F COVID-19 Ag	0.91 [0.75, 0.98]	0.97 [0.84, 1.00]	→ →
Weitzel 2020 [A]	49	0	30	30	CGIA	RapiGEN - BIOCREDIT COVID-19 Ag	0.62 [0.50, 0.73]	1.00 [0.88, 1.00]	
Weitzel 2020 [B]	0	1	9	9	CGIA	Liming Bio-Products - StrongStep	0.00 [0.00, 0.34]	0.90 [0.55, 1.00]	
Weitzel 2020 [C]	13	0	65	31	LFA (nos)	Savant Biotech - Huaketai SARS-CoV-2 N	0.17 [0.09, 0.27]	1.00 [0.89, 1.00]	-
Weitzel 2020 [D]	68	0	12	31	FIA	Shenzhen Bioeasy Biotech - 2019-nCoV Ag	0.85 [0.75, 0.92]	1.00 [0.89, 1.00]	h 0 2 0 4 0 6 0 9 1 h 0 2 0 4 0 6 0 9 1

Results for molecular tests overall and by subgroup are reported in Table 5. Forest plots of study data for the primary analysis is in Figure 9 and for subgroup analyses by Ct value, study design and sensitivity analyses by pre- and post-discrepant analysis in

Appendix 16. Individual plots by test brand are provided in Figure 10. Full identification details for studies of molecular-based assays are provided in Appendix 11 and Appendix 12. Appendix 17 provides



forest plots for study data according to Ct value and discrepant analysis.

Figure 9. Forest plot of studies evaluating rapid molecular tests. A&E: accident and emergency

Study	TP	FD	FN	TN	Symptom status	Setting	Sansitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
Chen 2020a	55			0	, .	Hospital in-patient	1.00 [0.94, 1.00]	Not estimable	
Wong 2020	118	ō	1	43	Symptomatic		0.99 [0.95, 1.00]	1.00 [0.92, 1.00]	
Assennato 2020	87	3	1		Symptomatic	Laboratory-based	0.99 [0.94, 1.00]	0.96 [0.90, 0.99]	4 4
Zhen 2020 [B]	57	0	_		Symptomatic	Laboratory-based	0.98 [0.91, 1.00]	1.00 [0.93, 1.00]	4
Gibani 2020	67	ő	4		Symptomatic	Mixed	0.94 [0.86, 0.98]	1.00 [0.99, 1.00]	-
Cradic 2020(b)	12	ŏ		169	Symptomatic	Hospital A&E	0.92 [0.64, 1.00]	1.00 [0.98, 1.00]	
Cradic 2020(a)	30	ő		151	Symptomatic	Mixed	0.91 [0.76, 0.98]	1.00 [0.98, 1.00]	-
Collier 2020	29	3		113		Hospital in-patient	0.88 [0.72, 0.97]	0.97 [0.93, 0.99]	<u> </u>
Zhen 2020 [A]	50	ō			Symptomatic	Laboratory-based	0.88 [0.76, 0.95]	1.00 [0.93, 1.00]	
SoRelle 2020	32	ŏ			Symptomatic	Laboratory-based	0.82 [0.66, 0.92]	1.00 [0.92, 1.00]	
Moore 2020	94	ŏ		79	Symptomatic	Mixed	0.79 [0.71, 0.86]	1.00 [0.95, 1.00]	
Harrington 2020	139	2			Symptomatic	Hospital A&E	0.75 [0.68, 0.81]	0.99 [0.98, 1.00]	
Thwe 2020	8	ō	6		Symptomatic	Laboratory-based	0.57 [0.29, 0.82]	1.00 [0.98, 1.00]	
Dust 2020	20	ō			Not reported	Laboratory-based	1.00 [0.83, 1.00]	1.00 [0.81, 1.00]	—
Goldenberger 2020	10	ō	ō	9	Not reported	Laboratory-based	1.00 [0.69, 1.00]	1.00 [0.66, 1.00]	
Lephart 2020 [B]	16	2	ō	_	Not reported	Hospital A&E	1.00 [0.79, 1.00]	0.97 [0.88, 1.00]	
Jokela 2020	60	0	ō	30	Not reported	Laboratory-based	1.00 [0.94, 1.00]	1.00 [0.88, 1.00]	
Lieberman 2020	13	ō	ō		Not reported	Laboratory-based	1.00 [0.75, 1.00]	1.00 [0.75, 1.00]	—
Moran 2020	42	1	0	60	Not reported	Laboratory-based	1.00 [0.92, 1.00]	0.98 [0.91, 1.00]	
Wolters 2020	58	0	0	30	Not reported	Laboratory-based	1.00 [0.94, 1.00]	1.00 [0.88, 1.00]	-
Loeffelholz 2020	219	11	1	250	Not reported	Laboratory-based	1.00 [0.97, 1.00]	0.96 [0.93, 0.98]	
Smithgall 2020 [B]	87	2	1	23	Not reported	Laboratory-based	0.99 [0.94, 1.00]	0.92 [0.74, 0.99]	• -
Broder 2020	34	0	1	0	Not reported	Laboratory-based	0.97 [0.85, 1.00]	Not estimable	-
Hou 2020	147	5	6	127	Not reported	Laboratory-based	0.96 [0.92, 0.99]	0.96 [0.91, 0.99]	
Rhoads 2020	90	0	6	0	Not reported	Laboratory-based	0.94 [0.87, 0.98]	Not estimable	-
Smithgall 2020 [A]	65	0	23	25	Not reported	Laboratory-based	0.74 [0.63, 0.83]	1.00 [0.86, 1.00]	 -
Mitchell 2020	33	0	13	15	Not reported	Laboratory-based	0.72 [0.57, 0.84]	1.00 [0.78, 1.00]	─
Lephart 2020 [A]	11	0	5	59	Not reported	Hospital A&E	0.69 [0.41, 0.89]	1.00 [0.94, 1.00]	
Hogan 2020	34	0	16	50	Not reported	Laboratory-based	0.68 [0.53, 0.80]	1.00 [0.93, 1.00]	
Stevens 2020	53	0	1	50	Mixed	Laboratory-based	0.98 [0.90, 1.00]	1.00 [0.93, 1.00]	
Ghofrani 2020	16	1	1	95	Mixed	Mixed	0.94 [0.71, 1.00]	0.99 [0.94, 1.00]	
Jin 2020	4	0	2	46	Mixed	Laboratory-based	0.67 [0.22, 0.96]	1.00 [0.92, 1.00]	
									0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1



Figure 10. Forest plot by test brand for molecular assays. A&E: accident and emergency; IFU: instructions for use

Abbott - ID NOW (Isothermal PCR)											
Study	TP	FP	FN	TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)			
Gh o frani 2020	16	1	1	95	No	0.94 [0.71, 1.00]	0.99 [0.94, 1.00]				
Smithgall 2020 [A]	65	0	23	25	No	0.74 [0.63, 0.83]	1.00 [0.86, 1.00]	-			
Rhoads 2020	90	0	6	0	No	0.94 [0.87, 0.98]	Not estimable	-			
Moore 2020	94	0	25	79	No	0.79 [0.71, 0.86]	1.00 [0.95, 1.00]				
Mitchell 2020	33	0	13	15	No	0.72 [0.57, 0.84]	1.00 [0.78, 1.00]	-			
Zhen 2020 [A]	50	0	7	50	No	0.88 [0.76, 0.95]	1.00 [0.93, 1.00]				
SoRelle 2020	32	0	7	44	No	0.82 [0.66, 0.92]	1.00 [0.92, 1.00]				
Cradic 2020(a)	30	0	3	151	No	0.91 [0.76, 0.98]	1.00 [0.98, 1.00]	→			
Cradic 2020(b)	12	0	1	169	Unclear	0.92 [0.64, 1.00]	1.00 [0.98, 1.00]				
Lephart 2020 [A]	11	0	5	59	Yes	0.69 [0.41, 0.89]	1.00 [0.94, 1.00]	─			
Jin 2020	4	0	2	46	Yes	0.67 [0.22, 0.96]	1.00 [0.92, 1.00]				
Harrington 2020	139	2	47	336	Yes	0.75 [0.68, 0.81]	0.99 [0.98, 1.00]	-			
Thwe 2020	8	0	6	147	Y e s	0.57 [0.29, 0.82]	1.00 [0.98, 1.00]				
								0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			
Cepheid - Xpert X	press	(Au	tom	ated	RT-PCR)						
Study	TP	FP	FN	I TN	IFU compliant	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)			
Goldenberger 2020	10	0	0	9	No	1.00 [0.69, 1.00]	1.00 [0.66, 1.00]				
Chen 2020a	55	0	0	0	No	1.00 [0.94, 1.00]	Not estimable	-			
Hou 2020	147	5	6	127	No	0.96 [0.92, 0.99]	0.96 [0.91, 0.99]				
Stevens 2020	53	0) 1	50	No	0.98 [0.90, 1.00]	1.00 [0.93, 1.00]				
Zhen 2020 [B]	57	0) 1	50	No	0.98 [0.91, 1.00]	1.00 [0.93, 1.00]				
Wong 2020	118	0) 1	. 43	No	0.99 [0.95, 1.00]	1.00 [0.92, 1.00]	4 -4			
Wolters 2020	58	0) 0	30	No	1.00 [0.94, 1.00]	1.00 [0.88, 1.00]				
Dust 2020	20	0) (18	Unclear	1.00 [0.83, 1.00]	1.00 [0.81, 1.00]	-			
Jokela 2020	60	0	0	30	Unclear	1.00 [0.94, 1.00]	1.00 [0.88, 1.00]				
Smithgall 2020 [B]	87	2	1	. 23	Unclear	0.99 [0.94, 1.00]	0.92 [0.74, 0.99]	• -			
Moran 2020	42	1	. 0	60	Unclear	1.00 [0.92, 1.00]	0.98 [0.91, 1.00]				
Loeffelholz 2020	219	11	. 1	250	Unclear	1.00 [0.97, 1.00]	0.96 [0.93, 0.98]				
Broder 2020	34	- 0) 1	. 0	Yes	0.97 [0.85, 1.00]	Not estimable	· —			
Lieberman 2020	13	0	0	13	Yes	1.00 [0.75, 1.00]	1.00 [0.75, 1.00]	-			
Lephart 2020 [B]	16	2	2 0	56	Yes	1.00 [0.79, 1.00]	0.97 [0.88, 1.00]				
DNANudge - COVI	D Nud	ge (Auto	omate	ed RT-PCR)			0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			
Study TP	FP FN	. т	KI IE	III so	mpliant Cancit	ivity (95% CI) Specif	isitu (OEW CI)	Sensitivity (95% CI)Specificity (95% CI)			
,	0 4			0 0	•		•				
Gibani 2020 67	0 4	31	.5		Yes 0.9	94 [0.86, 0.98] 1.0	00 [0.99, 1.00]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			
DRW - SAMBA II (A	utoma	ated	RT-	PCR)				0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1			
Study	TP FP	FN	ı T	N IFI	Lonmoliant Se	ensitivity (95% CI) Sp	ecificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)			
•	87 3			1	Unclear	0.99 [0.94, 1.00]	0.96 [0.90, 0.99]				
	29 3	_	1 11		Yes	0.88 [0.72, 0.97]	0.97 [0.93, 0.99]				
Comer 2020	25 3	, -	+ 11		160	0.00 [0.72, 0.97]	0.57 [0.55, 0.58]				

Collier 2020	29	3	4 113	Yes

Mesa Biotech - Accula (other molecular) TP FP FN TN IFU compliant Sensitivity (95% CI) Specificity (95% CI) Study Nο

0 0 2 0 4 0 6 0 8 1 Sensitivity (95% CI)Specificity (95% CI)

0 0.2 0.4 0.6 0.8 1

Accuracy of antigen tests overall and by subgroup

Hogan 2020 34 0 16 50

Results showed high levels of heterogeneity in sensitivity. Average sensitivity was 68.9% (95% CI 61.8% to 75.1%) and average specificity was 99.6% (95% CI 99.0% to 99.8%) across the 51 evaluations of antigen tests reporting both sensitivity and specificity (based on 21,614 samples, including 6136 samples with confirmed SARS-CoV-2; Table 2; Figure 3). Adding the six 'sensitivity only' datasets and single 'specificity only' datasets had a negligible impact on results (Table 2). In the sections below we show that there are substantial differences between subgroups of studies according to symptom status, timing, test method and brand, therefore this average value is unlikely to accurately predict the performance of the test in a given setting and should not be used for this purpose.

Subgroup analysis by symptom status

1.00 [0.93, 1.00]

Subgroup analysis by symptom status suggests that average test sensitivity to detect infection is 13.8 percentage points lower in asymptomatic (58.1%, 95% CI 40.2% to 74.1%; based on 12 evaluations, 1581 samples and 295 cases) compared to symptomatic (72.0%, 95% CI 63.7% to 79.0%; based on 37 evaluations, 15,530 samples and 4410 cases) participants (95% CI for the difference in sensitivity: 33.1 percentage points lower to 5.4 percentage points higher; Table 2; Figure 4). Restricting the comparison by symptom status to the nine evaluations reporting data for both symptomatic and asymptomatic subgroups (thus ensuring the comparison is made between the same tests used in the same way) showed a similar difference in sensitivity (14.4 percentage points lower in asymptomatic participants, 95% CI 38.8 lower to 10.0 percentage points higher; Table 2). Average results for the 19 evaluations in participants with mixed symptom status

0.68 [0.53, 0.80]



(n = 10) or symptom status not reported (n = 9) were between those observed for the symptomatic and asymptomatic subgroups: sensitivity 63.0% (95% CI 52.2% to 72.6%) and specificity 98.4% (95% CI 98.0% to 98.8%) (6220 samples; 2392 cases).

We did not observe any important differences in specificity according to symptom status (Table 2).

Subgroup analysis by time from symptom onset

We pooled data by time from symptom onset separately for sensitivity and specificity because the majority of evaluations did not report these data for people without SARS-CoV-2 (Table 2; Figure 5). Sensitivity was 78.3% (95% CI 71.1% to 84.1%) (26 evaluations; 5769 samples, 2320 cases) in the first seven days after symptom onset compared to 51.0% (40.8% to 61.0%) (22 evaluations; 935 samples, 692 cases) in the second week of symptoms (a decrease of 27.3 percentage points, 95% CI -32.8 to -21.9 percentage points decrease). This difference remained on restriction to the 22 evaluations reporting data for people in both week one and week two of symptoms (removing other between-study differences; Table 2).

We did not observe any differences in specificity according to time after symptom onset (Table 2).

Subgroup analysis by Ct value

A total of 36 evaluations reported sensitivity according to Ct value using a threshold of 24 (n = 18) or 25 (n = 18) Ct or less to define higher viral load (Table 2; Appendix 15). Summary sensitivity in those with higher viral load was 94.5% (95% CI 91.0% to 96.7%) (based on 2613 cases), compared to 40.7% in those with lower viral load (95% CI 31.8% to 50.3%) (based on 2632 cases) (i.e. sensitivity was 53.8 percentage points lower for those with lower viral load; 95% CI 63.6 to 44.1 percentage points lower)). Applying a Ct threshold of \leq 33 (n = 13) or < 32 (n = 2) led to a bigger difference in sensitivity although the number of samples in the lower viral load subgroup was considerably smaller: sensitivity associated with higher viral load was 82.5% (95% CI 74.0% to 88.6%) (based on 2127 samples) and for lower viral load was 8.9% (3.3% to 21.7%) (based on 346 samples), a difference of 73.5 percentage points (95% CI 84.7 to 62.4 percentage points lower).

Subgroup analysis by study design

We did not observe any clear differences in average sensitivity or specificity when studies were grouped by study design (15,336 samples and 3536 cases in 29 single group studies and 5729 samples and 2396 cases in 20 two-group studies; Table 2; Appendix 15). Average sensitivity was lower in two-group studies (64.1%, 95% CI 48.5% to 77.2%) compared to single-group studies (72.1%, 95% CI 64.8% to 78.3%), however confidence intervals overlapped and the difference was within that which may be expected by chance (8.0 percentage points lower, 95% CI from 24.2 percentage points lower to 8.2 higher). Average specificities were 2.3 percentage points lower in the two-group studies (95% CI from 2.9 to 1.6 percentage points lower), at 97.3% (95% CI 96.7% to 97.8%) compared to 99.6% (95% CI 99.1% to 99.8%) in single-group studies.

Subgroup analysis by test method

We observed differences in accuracy according to test method (Table 2). The majority of evaluations (n = 36; 17,448 samples,

5085 cases) reported using a CGIA, average sensitivity was lower (64.0%, 95% CI 55.7% to 71.6%) than for FIAs (79.6%, 95% CI 67.5% to 88.0%; n = 9; 2820 samples, 712 cases; absolute difference of 15.6 percentage points, 95% CI 2.6 to 28.5 percentage points). We also observed marginal differences in specificity, with estimates of 99.0% (95% CI 98.8% to 99.2%) for CGIA and 97.7% (95% CI 95.3% to 98.8%) for FIA, a difference of 1.3 percentage points (95% from 3.0 percentage points lower to 0.3 higher). Results for lateral flow assays where the method could not be determined (n = 5) and for the single evaluation of an alkaline phosphatase (ALP)-labelled assay were heterogeneous but largely in the realms of those observed for the other assay types (Table 2).

Results by test brand according to symptom status and IFU compliance

Results by test brand overall and sensitivity analyses by IFU compliance (based on sample type, use of viral transport medium, and time period between sample collection and test procedure) are reported in Table 3. Results by test brand for symptomatic and asymptomatic subgroups overall and by IFU compliance are in Table 4. Given the mixed settings in which asymptomatic individuals were tested (Results of the search), the data for asymptomatic subgroups cannot be considered applicable to any particular scenario for asymptomatic testing. Only three studies reported direct comparisons of tests, two using nasopharyngeal or oropharyngeal samples (Fourati 2020 [A]; Weitzel 2020 [A]).

We observed considerable heterogeneity in sensitivities for all assays.

AAZ - COVID-VIRO

Two evaluations of the COVID-VIRO assay included 880 samples and 396 SARS-CoV2-positive samples (Figure 7). We did not pool the studies due to the heterogeneity in both sensitivity and specificity, although both were conducted in symptomatic or mainly symptomatic participants using nasopharyngeal samples.

In one study that compared antigen assays using nasopharyngeal samples in viral transport medium, sensitivity was 61.7% (95% CI 55.9% to 67.3%) and specificity (in pre-pandemic samples) 100% (95% CI 98.9% to 100%; 632 samples, 295 cases; 'Fourati 2020 [E]).

The second study used direct swab testing in compliance with the manufacturer's IFU. Twenty participants in the study who previously tested positive on PCR retested negative with PCR at the time of the antigen test. All twenty samples showed weak lines on antigen testing. We considered these as false positives in the review (based on the negative result of the concurrent PCR test) whereas the study authors considered them to be true positives. With our re-calculation, the test demonstrated sensitivity of 96.0% (95% CI 90.2% to 98.9%) and specificity of 86.4% (95% CI 79.8% to 91.5%; Courtellemont 2020). Sensitivity in this study may have been inflated by the inclusion of hospitalised, confirmed SARS-CoV-2-positive participants.

Abbott - Panbio Covid-19 Ag

We identified 11 evaluations of the Panbio assay, including 5691 unique samples, with 2031 SARS-CoV-2-positive cases (Figure 6). One of the 11 evaluations included only SARS-CoV-2-positive cases (n = 182 samples). Studies were conducted in community COVID-19 test centres or emergency departments (n = 6), in contacts of confirmed cases (n = 2), and laboratory-based evaluations (n = 2).



The setting was not clear in one study. Participants were reportedly symptomatic (n = 5), asymptomatic (n = 1), with mixed symptom status (n = 4), or symptom status was not reported (n = 1). Nine evaluations used nasopharyngeal samples (Albert 2020; Billaud 2020; Fenollar 2020(b); FIND 2020b; Fourati 2020 [C]; Gremmels 2020(a); Gremmels 2020(b); Linares 2020), one (Alemany 2020), tested nasopharyngeal or nasal samples and one (Schildgen 2020 [A]), used bronchoalveolar lavage or throat wash samples. Only three of the 11 evaluations reported product codes for the assays used, one of which was for the assay for use with nasopharyngeal swabs (41FK10) and two (from the same study report) were for the assay for use with nasal swabs (41FK11), although the study reports using nasopharyngeal samples (Gremmels 2020(a); Gremmels 2020(b)).

Five of the 11 evaluations complied with manufacturer IFU for the test. Reasons for non-compliance included use of viral transport medium, frozen storage, type of swab tested, or lack of clear reporting of test procedures used.

The average sensitivity and specificity of the Panbio assay were:

- 72.0% (95% CI 60.6% to 81.1%) and 99.3% (95% CI 99.0% to 99.6%) overall (n = 10; 5509 samples; 1849 cases; Table 3);
- 74.1% (95% CI 60.8% to 84.0%) and 99.8% (95% CI 99.5% to 99.9%) in symptomatic people (n = 8; 3699 samples, 1162 cases); and
- 58.1% (95% CI 41.7% to 72.9%) and 98.4% (95% CI 92.2% to 99.7%) in asymptomatic people (n = 6; 1097 samples, 190 cases; Table 4).

Restricting to IFU-compliant evaluations, average sensitivities and specificities were:

- 72.0% (95% CI 56.5% to 83.5%) and 99.2% (95% CI 98.5% to 99.5%) overall (n = 5; 1776 samples, 362 cases; Table 3);
- 75.1% (95% CI 57.3% to 87.1%) and 99.5% (95% CI 98.7% to 99.8%) in symptomatic people (n = 3; 1094 samples, 252 cases); and
- 48.9% (95% CI 35.1% to 62.9%) and 98.1% (95% CI 96.3% to 99.1%) in asymptomatic people (n = 2; 474 samples, 47 cases; Table 4).

The addition of one evaluation that reported sensitivity only in symptomatic participants led to only marginal differences in average sensitivity (Fenollar 2020(a); Table 4).

Becton Dickinson - BD Veritor

We identified three evaluations of the BD Veritor assay, including 727 unique samples, with 180 SARS-CoV-2-positive cases (Figure 6). One of the three evaluations included only SARS-CoV-2-positive cases (n=125 samples). Studies were conducted in community COVID-19 test centres (n=2), or in multiple settings (n=1). All participants were symptomatic. Two evaluations used combined naso- and oropharyngeal samples and one tested nasal samples.

None of the evaluations complied with manufacturer IFU for the test because the interval between sample collection and testing was greater than the maximum of one hour.

Average sensitivity and specificity of the BD Veritor assay were:

82.3% (95% CI 62.1% to 93.0%) and 99.5% (95% CI 98.3%, 99.8%) in symptomatic people (n = 2; 602 samples, 55 cases; Van der Moeren 2020(a); Young 2020; Table 3; Table 4).

Adding the 'cases only' evaluation reduced average sensitivity to 79.4% (95% CI 72.9% to 84.7%) (n = 3; 180 cases; Van der Moeren 2020(b)).

The BD Veritor assay requires interpretation using a Veritor analyzer device, but Van der Moeren 2020(a) found that visual inspection of the test device resulted in the same sensitivity as with the Analyzer device, and similar specificity (100% compared to 99% using the Analyzer device).

BIONOTE - NowCheck COVID-19 Ag

We identified a single IFU-compliant evaluation of the NowCheck assay in symptomatic participants (FIND 2020a; Figure 7). The study included 400 samples with 102 SARS-CoV-2-positive cases, from participants presenting at a community-based COVID-19 test centre.

The sensitivity and specificity in this study were 89.2% (95% CI 81.5% to 94.5%) and 97.3% (95% CI 94.8% to 98.8%; Table 3; Table 4).

Biosynex - Biosynex COVID-19 Ag BSS

We identified a single evaluation of the Biosynex assay in symptomatic participants (Fourati 2020 [D]), including 634 samples with 297 with confirmed SARS-CoV-2 (Figure 7). The evaluation was not in compliance with the manufacturer's IFU because samples were stored in viral transport medium and frozen prior to testing. The setting in which participants presented for testing was not reported.

Observed sensitivity was 59.6% (95% CI 53.8% to 65.2%) and specificity 100% (95% CI 98.9% to 100%; Table 3; Table 4).

Coris Bioconcept - COVID-19 Ag Respi-Strip

The seven evaluations of the Coris Bioconcept assay included 1781 samples, with 707 SARS-CoV-2-positive cases (Blairon 2020; Fourati 2020 [A]; Kruger 2020(b); Lambert-Niclot 2020; Mertens 2020; Scohy 2020; Veyrenche 2020; Figure 6). Five of the seven were laboratory-based evaluations with limited detail regarding study participants. One study recruited from community-based COVID-19 test centres and one included samples from hospital inpatients. Three studies included only or mainly symptomatic participants, one was in a mixed group and three did not report symptom status.

All evaluations tested naso- or oropharyngeal swabs and were compliant with the manufacturer IFU, however, it may be worth noting that the IFU for this assay permits the use of viral transport medium and freezing of samples, although immediate testing is recommended.

The average sensitivity and specificity of the COVID-19 Ag Respi-Strip were:

- 39.7% (95% CI 31.3% to 48.7%) and 98.3% (95% CI 97.4% to 98.9%) overall (n = 7; 1781 samples, 707 cases; Table 3);
- 34.1% (95% CI 29.7% to 38.8%) and 100% (95% CI 99.0% to 100%) in symptomatic people (n = 3; 780 samples, 414 cases); and



28.6% (95% CI 8.4% to 58.1%) and 100% (95% CI 88.8% to 100%) in asymptomatic people (n = 1; 45 samples, 14 cases; Scohy 2020; Table 4).

E25Bio - DART (nasopharyngeal)

We identified a single evaluation of the E25Bio DART assay that included 190 samples, 100 with SARS-CoV-2 (Nash 2020; Figure 7). The symptom status of included participants was not reported and the manufacturer IFU is not yet available as the assay has been submitted for Emergency Use Authorisation (EUA) approval with the US Food and Drug Administration (FDA).

Sensitivity was 80.0% (95% CI 70.8% to 87.3%) and specificity 91.1% (95% CI 83.2% to 96.1%; Table 3).

Fujirebio - ESPLINE SARS-CoV-2

We included two eligible evaluations were included, with a total of 265 samples, 165 were SARS-COV-2-positive (Nagura-Ikeda 2020; Takeda 2020; Figure 7). One study reported only sensitivity data (Nagura-Ikeda 2020).

Takeda 2020 reported sensitivity of 80.6% (95% CI 68.6% to 89.6%) and specificity of 100% (95% CI 96.4% to 100%) in nasopharyngeal samples (162 samples, 62 cases; Table 3). They did not report symptom status of participants and provided insufficient detail to allow us to judge IFU compliance.

Nagura-Ikeda 2020 evaluated the assay using saliva samples in symptomatic participants (not within IFU specifications), the ESPLINE assay correctly identified 12 of 103 PCR-positive samples (sensitivity 11.6%, 95% CI 6.2% to 19.5%; Table 3; Table 4).

Innova Medical Group - Innova SARS-CoV-2 Ag

We included one report that evaluated the Innova study as six separate substudies; three reported both sensitivity and specificity (PHE 2020(a); PHE 2020(b); PHE 2020(c) [non-HCW tested]), two reported sensitivity alone (PHE 2020(d) [HCW tested]; PHE 2020(d) [Lab tested]), and one reported specificity alone (PHE 2020(e); Figure 6). The studies reported a total of 3904 participants, including 1017 SARS-CoV-2-positive cases. Detail regarding symptom status, was limited, however the study populations were coded as: symptomatic (samples from hospital inpatients in PHE 2020(a)), mainly symptomatic for samples from COVID-19 testing centres (PHE 2020(c) [non-HCW tested]; PHE 2020(d) [HCW tested]; PHE 2020(d) [Lab tested]), although data on symptom status were reported for only two of these studies (PHE 2020(d) [HCW tested]; PHE 2020(d) [Lab tested]), not reported for the outbreak investigation in PHE 2020(b) and asymptomatic staff screening for PHE 2020(e). The study authors for the outbreak evaluation study did not report the sensitivity value of 28.3% (95% CI 16.0% to 43.5%) in the publications but provided it to us on request.

All evaluations used naso- or oropharyngeal samples, two in viral transport medium (PHE 2020(a); PHE 2020(b)), and four using direct swab testing in compliance with manufacturer IFU (PHE 2020(c) [non-HCW tested]; PHE 2020(d) [HCW tested]; PHE 2020(d) [Lab tested]; PHE 2020(e)).

For studies reporting both sensitivity and specificity, average sensitivity and specificity were:

- 47.9% (95% CI 34.3% to 61.8%) and 99.8% (95% CI 99.5% to 99.9%) overall (n = 3; 2945 samples, 596 cases; Table 3); and
- 56.2% (95% CI 52.0% to 60.3%) and 99.8% (95% CI 99.5% to 99.9%) in symptomatic people (n = 2; 2794 samples, 550 cases; Table 4).

Only one of the three studies that reported both sensitivity and specificity was compliant with manufacturer IFU, the sensitivity and specificity were:

• 57.5% (95% CI 52.3% to 62.6%) and 99.6 (95% CI 99.1%, 99.9%) overall (n = 1; 1676 samples, 372 cases).

Summary results from the four IFU-compliant evaluations were calculated as follows:

- average sensitivity across three evaluations of mainly symptomatic participants 69.1% (95% CI 58.3% to 78.2%; n = 3; 793 cases; Table 3; Table 4);
- average specificity from two evaluations of 99.7% (95% CI 99.3% to 99.9%; n = 2; 1842 samples with no SARS-CoV-2; Table 3).

Adding data from single-group evaluations in either RT-PCR-positive or RT-PCR-negative participants:

- average sensitivity was 59.0% (43.4%, 73.0%) (n = 5; 1015 cases)
- average specificity was 99.8% (99.5%, 99.9%) (n = 4; 2887 RT-PCR negative samples) (Table 4).

Results for each of the three IFU-compliant evaluations by test operator were (Figure 6):

- sensitivity of 57.5% (95% CI 52.3% to 62.6%) and specificity 99.6% (95% CI 99.1% to 99.9%), when the test was used by selftrained, non-healthcare workers (n = 1; 1676 samples, 372 cases; PHE 2020(c) [non-HCW tested]);
- sensitivity of 70.0% (95% CI 63.5% to 75.9%) when the test was used by healthcare workers (n = 1; 223 cases; PHE 2020(d) [HCW tested]);
- sensitivity of 78.8% (95% CI 72.4% to 84.3%) when the test was used by laboratory scientists (n = 1; 198 cases; PHE 2020(d) [Lab tested]).

Liming Bio-Products - StrongStep® COVID-19 Ag

We identified a single evaluation of the StrongStep assay in 19 symptomatic participants with nine SARS-CoV-2 positive samples ((Weitzel 2020 [B]; Figure 7). We could not identify the manufacturer's IFU for this assay. The study authors terminated the evaluation early following poor early results for this assay.

Sensitivity was 0% (95% CI 0% to 33.6%) and specificity 90.0% (95% CI 55.5% to 99.7%; 19 samples, 9 cases; Table 3; Table 4).

Quidel Corporation - SOFIA SARS Antigen

We identified a single evaluation of the SOFIA assay in symptomatic participants, including 64 samples with 32 SARS-CoV-2-positive cases (Porte 2020b [A]; Figure 7). The study used combined naso- and oropharyngeal swab samples in viral transport medium, therefore the evaluation was not compliant with the manufacturer IFU.

Sensitivity was 93.8% (95% CI 79.2% to 99.2%) and specificity was 96.9% (95% CI 83.8% to 99.9%; Table 3; Table 4).



RapiGEN - BIOCREDIT COVID-19 Ag

We identified six evaluations of the RapiGen BIOCREDIT assay; these reported data for 2170 samples, with 470 confirmed SARS-COV-2-positive cases (FIND 2020e (BR); FIND 2020e (DE); Mak 2020; Schildgen 2020 [A]; Shrestha 2020; Weitzel 2020 [A]; Figure 6). One laboratory-based study included cases only (n = 160). The other evaluations included participants from community-based COVID-19 test centres (n = 2), emergency departments (n = 1), contact tracing (n = 1) or did not clearly report the setting (n = 1). Two studies included only symptomatic participants, two reported including both symptomatic and asymptomatic participants (mixed group) and one did not report symptom status. All evaluations apart from one (Schildgen 2020 [A]), tested nasopharyngeal or combined naso- or oropharyngeal samples.

Only three of the six evaluations complied with manufacturer IFU, with non-compliance because of the use of viral transport medium, or the type of swab tested.

The average sensitivity and specificity of the BIOCREDIT assay were:

- 63.3% (95% CI 45.7% to 78.0%) and 99.5% (95% CI 99.1 to 99.8) overall (n = 5; 2010 samples, 310 cases; Table 3);
- 58.4% (95% CI 36.3% to 77.5%) and 96.4% (95% CI 82.8% to 99.3%) in symptomatic people (n = 3; 608 samples, 206 cases);
- 63.2% (95% CI 21.7% to 91.4%) and 98.9% (95% CI 82.9% to 99.9%) in asymptomatic people (n = 2; 140 samples, 60 cases) (Table 4).

Restricting to IFU-compliant evaluations, average sensitivities and specificities were:

- 73.0% (95% CI 57.4% to 84.4%) and 99.8% (95% CI 99.4% to 99.9%) overall (n = 3; 1828 samples, 189 cases; Table 3);
- 74.4% (95% CI 65.5% to 82.0%) and 98.9% (95% CI 97.2% to 99.7%) in symptomatic people (n = 1; 476 samples, 117 cases);
- 85.1% (95% CI 71.7% to 93.8%) and 100% (95% CI 94.6% to 100%) in asymptomatic people (n = 1; 113 samples, 47 cases; Shrestha 2020; Table 4).

The addition of one evaluation that reported sensitivity only led to a decrease in overall average sensitivity of 5.6 percentage points (Mak 2020; Table 4).

Roche - SARS-CoV-2

According to the manufacturer IFU, the Roche SARS-CoV-2 assay is available under a partnership with SD Biosensor.

There was a single evaluation of the Roche assay using 73 bronchoalveolar lavage or throat wash samples (not covered by the IFU) in participants with mixed symptom status (Figure 7); 42 of the 73 samples were RT-PCR-positive (Schildgen 2020 [A]).

Overall, using bronchoalveolar lavage or throat wash samples, the sensitivity and specificity were 88.1% (95% CI 74.4% to 96.0%) and 19.4% (95% CI 7.5% to 37.5%) (73 samples, 42 cases; Table 3). Only the results for the subgroup of 50 throat wash samples could be separated by symptom status:

in symptomatic participants, sensitivity was 100% (95% CI 69.2% to 100%) and specificity was 7.7% (95% CI 0.2% to 36.0%) with 23 throat wash samples and 10 cases;

• in asymptomatic participants, sensitivity was 84.6% (95% CI 54.6% to 98.1%) and specificity was 14.3% (95% CI 1.8% to 42.8%), with 27 throat wash samples, 13 cases; Table 4).

Savant Biotech - Huaketai SARS-CoV-2 N Protein

We identified a single evaluation of the Huaketai assay in 109 symptomatic participants, using combined naso- or oropharyngeal swabs in viral transport medium (Weitzel 2020 [C]; Figure 7). We could not obtain the manufacturer IFU.

Sensitivity was 16.7% (95% CI 9.2% to 26.8%) and specificity was 100% (95% CI 88.8% to 100%; 109 samples, 78 cases; Table 3; Table 4).

SD Biosensor - STANDARD F COVID-19 Ag

We identified four evaluations of the STANDARD F assay; these reported data for 1552 samples, with 295 confirmed SARS-COV-2-positive cases (FIND 2020d (BR); FIND 2020d (DE); Liotti 2020; Porte 2020b [B]; Figure 6). Three evaluations included all or mainly symptomatic participants from community-based COVID-19 test centres and one was a laboratory-based study that did not provide details regarding symptom status.

All evaluations tested nasopharyngeal or combined nasoor oropharyngeal samples, however only two complied with manufacturer IFU. Reasons for non-compliance were the use of viral transport medium, or lack of information concerning viral transport medium.

The average sensitivity and specificity of the STANDARD F COVID-19 Ag assay were:

- 72.6% (95% CI 54.0% to 85.7%) and 97.5% (95% CI 96.4% to 98.2%) overall (n = 4; 1552 samples, 295 cases; Table 3);
- 78.0% (95% CI 71.6% to 83.3%) and 97.2% (95% CI 96.0% to 98.1%) in symptomatic people (n = 3; 1193 samples, 191 cases; Table 4).

No data for asymptomatic people were available.

Restricting to IFU-compliant evaluations, average sensitivity and specificity were:

 75.5% (95% CI 68.2% to 81.5%) and 97.2% (95% CI 96.0 to 98.1%), both studies in symptomatic people (n = 2; 1129 samples, 159 cases; Table 4).

SD Biosensor - STANDARD Q COVID-19 Ag

We identified six evaluations of the STANDARD Q assay; these reported data for 3480 samples, with 821 confirmed SARS-CoV-2-positive cases (Figure 6). Four evaluations included participants from community-based COVID-19 test centres, one was a laboratory-based study, and one included multiple settings. Four evaluations included symptomatic or mainly symptomatic participants, and two included mixed symptomatic and asymptomatic participants.

All evaluations tested nasopharyngeal or combined naso- or oropharyngeal samples, four of which were compliant with manufacturer's IFUs, the other two used samples in viral transport medium.



The average sensitivity and specificity of the STANDARD Q COVID-19 Ag assay were:

- 79.3% (95% CI 69.6% to 86.6%) and 98.5% (95% CI 97.9% to 98.9%) overall (n = 6; 3480 samples, 821 cases; Table 3);
- 80.1% (95% CI 68.5% to 88.1%) and 98.1% (95% CI 97.4% to 98.6%) in symptomatic people (n = 5; 2760 samples, 731 cases); and
- 61.1% (95% CI 37.9% to 80.2%) and 99.6% (95% CI 97.3% to 99.9%) in asymptomatic people (n = 2; 272 samples, 18 cases; Table 4).

Restricting to IFU-compliant evaluations, average sensitivities and specificities were:

- 85.8% (95% CI 80.5% to 89.8%) and 99.2% (95% CI 98.2% to 99.6%) overall (n = 4; 2522 samples, 421 cases; Table 3);
- 88.1% (95% CI 84.2% to 91.1%) and 99.1% (95% CI 97.8% to 99.6%) in symptomatic people (n = 3; 1947 samples, 336 cases);
- 69.2% (95% CI 38.6% to 90.9%) and 99.1% (95% CI 95.2% to 100%) in asymptomatic people (n = 1; 127 samples, 13 cases; Table 4).

Shenzhen Bioeasy Biotech - 2019-nCoV Ag

We included three evaluations of the Bioeasy FIA; these included 965 samples with 177 SARS-CoV-2-positive cases ((Kruger 2020(a); Porte 2020a; Weitzel 2020 [D]; Figure 6). Studies were conducted in hospital emergency departments (n = 2) or a community COVID-19 test centre (n = 1). Participants in studies were all symptomatic or mainly symptomatic.

Two evaluations used combined naso- or oropharyngeal swabs and one tested either nasopharyngeal or oropharyngeal swabs. Two evaluations used swabs in viral transport medium, which was not documented as suitable for use on the manufacturer IFU.

The average sensitivity and specificity of the Shenzhen Bioeasy assay were:

86.2% (95% CI 72.4% to 93.7%) and 93.8 (95% CI 91.9% to 95.3%) overall (all symptomatic; n = 3; 965 samples, 177 cases; Table 3; Table 4).

The single IFU-compliant evaluation Kruger 2020(a) reported sensitivity of 66.7% (95% CI 38.4% to 88.2%) and specificity of 93.1% (95% CI 91.0% to 94.9%; 727 samples, 15 cases).

We also included an additional study that reported the development of this assay but we did not pool data with the other evaluations as it was a development and not a validation study (Diao 2020; Figure 7). Sensitivity was 67.8% (95% CI 61.0% to 74.1%) and specificity was 100% (95% CI 88.8% to 100%; 239 samples, 208 cases).

Direct test comparisons

Three studies reported direct comparisons of different antigen assays in naso- or oropharyngeal samples; however none of the studies had any assay comparisons in common. All three studies utilised swabs in viral transport medium and all were conducted in symptomatic participants. We cannot derive any clear conclusions about comparative performance of tests from these studies.

Figure 8 shows variable diagnostic performance between and to some extent within studies. Four of the five assays in Fourati 2020 [A] demonstrated sensitivities in the range of 55% to 62% (SD Biosensor STANDARD Q, Abbott Panbio Covid-19 Ag, Biosynex COVID-19 Ag, AAZ – COVID-VIRO), with one outlier (Coris Bioconcept – Covid-19 Ag) at 35% (maximum of 297 cases). Specificity was 100% for all assays apart from SD Biosensor SDQ (specificity 93%; 337 pre-pandemic samples).

In Porte 2020b [A] (32 cases) both assays had sensitivities over 90% (SD Biosensor STANDARD F and Quidel Sofia SARS Antigen), with specificities 97% (32 non-COVID-19 samples)

Weitzel 2020 [A] observed a range in assay sensitivities from 0% for the Liming Bio-Products assay (based on only nine cases), to 17% (for Savant Biotech – Huaketai SARS-CoV-2 N), 62% (RapiGEN – BIOCREDIT COVID-19 Ag) and 85% for Shenzhen Bioeasy Biotech – 2019 nCov Ag (78 to 80 cases for the latter three assays). Specificities were 100% for all assays (based on 30 to 31 samples) apart from the one from Liming Bio-Products (specificity 90% based on 10 samples).

Accuracy of rapid molecular tests overall and by subgroup

Average sensitivity and specificity for the 29 rapid molecular test evaluations that included samples with and without SARS-CoV-2, were 95.1% (95% CI 90.5% to 97.6%) and 98.8% (95% CI 98.3% to 99.2%; 4351 samples, 1781 with confirmed SARS-CoV-2; Table 5). Adding the three 'cases only' studies made little difference to the average sensitivity (95.5%, 95% CI 91.5% to 97.7%; 1973 cases).

Figure 9 demonstrates heterogeneity in sensitivity estimates (ranging from 57% to 100%), with consistently high specificities (92% to 100%, but with upper limits of 95% CIs of 99% or 100% in every study).

Subgroup analyses by viral load

We extracted sensitivity data according to viral load from 10 evaluations of molecular tests, six of which reported data at a Ct threshold for higher viral load of 30 or less (Jokela 2020; Lieberman 2020; Mitchell 2020; Smithgall 2020 [A]; Smithgall 2020 [B]; Wolters 2020), four using Xpert Xpress and two using ID NOW. (Appendix 16)

All sensitivity estimates for the higher viral load subgroups were 100% (based on 204 samples with confirmed SARS-CoV-2), with a 95% CI for the average of 98.2% to 100%. For the lower viral load group, average sensitivity was 95.6% (95% CI 55.7% to 99.7%) (149 samples with confirmed SARS-CoV-2; Table 5).

We observed a similar pattern for the studies using alternative Ct thresholds to define higher and lower viral load (Appendix 17).

Subgroup analysis by study design

We did not observe any clear differences in average sensitivity or specificity when studies were separated by study design (2899 samples and 976 cases in 18 single-group studies and 1265 samples and 718 cases in nine two-group studies; Table 5; Appendix 17). Average sensitivity was higher in two-group studies (97.2%, 95% CI 90.7% to 99.2%) compared to single-group studies (93.2%, 95% CI 85.5% to 97.0%); a difference of 4.0 percentage points (95% CI from 2.2 percentage points lower to 10.1 higher). Average specificities had almost identical point estimates at 99.4% (95% CI 98.4 to 99.8%) and 99.3% (95% CI 96.5% to 99.8%) respectively (Table 5).



Abbott - ID NOW

Thirteen studies evaluated the ID NOW assay, with 1949 samples and 730 confirmed SARS-CoV-2 cases; one study included only SARS-CoV-2-positive cases (n = 36; Figure 10). Seven evaluations were laboratory-based, three recruited participants from emergency department settings and three were conducted in multiple settings. Seven studies included only symptomatic participants, two included both symptomatic and asymptomatic people, and four did not report symptom status.

Eleven evaluations used nasopharyngeal or nasal swab samples, one was conducted using saliva samples and one did not specify the sample type. Only four evaluations were compliant with manufacturer IFUs; lack of compliance was based on the use of viral transport medium, sample type, and interval between sample collection and testing.

Pooled analyses demonstrated average sensitivity and specificity of:

- 78.6% (95% CI 73.7% to 82.8%) and 99.8% (95% CI 99.2% to 99.9%) overall (n = 12; 1853 samples, 634 cases); and
- 73.0% (95% CI 66.8% to 78.4%) and 99.7% (95% CI 98.7% to 99.9%), restricted to evaluations that were compliant with the manufacturer's IFU (n = 4; 812 samples, 222 cases; Table 5).

Average sensitivity increased to 81.5% (95% CI 75.2% to 86.5%), with the addition of the cases only study (730 cases; Rhoads 2020).

Cepheid Inc - Xpert Xpress

The Xpert Xpress assay was evaluated in 15 studies using respiratory specimens, with 1781 samples and 1001 confirmed SARS-CoV-2 cases; two of the studies included only SARS-CoV-2-positive cases (n = 90; Figure 10). Thirteen evaluations were laboratory-based, one recruited participants from emergency department settings and one included samples from hospital inpatients. Three studies included only symptomatic participants, one included both symptomatic and asymptomatic people (mixed symptom status), and 11 did not report symptom status.

Fourteen evaluations used nasopharyngeal, oropharyngeal or nasal swab samples, and one was conducted using throat saliva or lower respiratory samples. Only three evaluations were compliant with manufacturer IFUs. Lack of compliance with the IFU was because of the use of frozen samples (n=8), or sample type (n=1) or concerns about the timing between sample collection and testing (n=3).

Pooled analyses demonstrated average sensitivity and specificity of:

- 99.1% (95% CI 97.7% to 99.7%) and 97.9% (95% CI 94.6 % to 99.2%) overall (n = 13; 1691 samples, 911 with confirmed SARS-CoV-2);
- 100% (95% CI 88.1% to 100%) and 97.2% (95% CI 89.4%, 99.3%), restricted to evaluations that were compliant with the manufacturer's IFU (n = 2; 100 samples, 29 cases; Table 5)

Average sensitivity did not change with addition of two cases-only studies (99.1%, 95% CI 97.8% to 99.6%; n = 15; 730 cases; Broder 2020; Chen 2020a).

One additional study considered accuracy in non-respiratory samples using Xpert Xpress (Szymczak 2020). Sensitivity in stool samples obtained up to 33 days after symptom onset was 93.1% (95% CI 77.2% to 99.1%) and specificity was 96.0% (95% CI 86.3% to 99.5%; 79 samples, 29 cases).

Comparison of ID NOW with Xpert Xpress

Comparing the overall pooled results between ID NOW and Xpert Xpress, the average sensitivity of Xpert Xpress was 19.8 (95% CI 14.9 to 24.7) percentage points higher than that of ID NOW (P < 0.0001; Table 5).

The average specificity of Xpert Xpress was marginally lower than that of ID NOW, a difference of -1.9 percentage points (95% CI -3.8 to -0.1).

DNAnudge - COVID Nudge

We included one evaluation of COVID Nudge with a total of 386 participants and 71 SARS-CoV-2-positive cases (Gibani 2020; Figure 10). Participants were recruited from multiple settings including hospital inpatients (n = 88), accident and emergency (n = 15) and healthcare workers and their families (n = 280). All participants were symptomatic and direct testing of nasopharyngeal samples was used (within manufacturer IFU).

The sensitivity of the COVID Nudge assay was 94.4% (95% CI 86.2 to 98.4%) and specificity was 100% (95% CI 98.8% to 100%; 386 samples and 71 cases; Table 5).

Diagnostics for the Real World (DRW) - SAMBA II

We included two evaluations of SAMBA II with 321 samples (121 with confirmed SARS-CoV-2; Figure 10). All participants were symptomatic. One study conducted direct testing of combined naso- or oropharyngeal samples from hospital inpatients and the other obtained combined naso- or oropharyngeal samples in viral transport medium from Public Health England. It was not reported whether the PHE samples were stored or frozen prior to testing so we could not determine whether they complied with the IFU for the assay.

The average sensitivity and specificity of SAMBA-II were 96.0% (95% CI 81.1% to 99.3%) and 97.0% (95% CI 93.5% to 98.6%; 2 studies; 321 samples, 121 with confirmed SARS-CoV-2; Table 5).

In the IFU-compliant evaluation, sensitivity was 87.9% (95% CI 71.8% to 96.6%) and specificity was 97.4% (95% CI 92.6% to 99.5%; 149 samples, 33 cases; Collier 2020; Table 5).

Mesa Biotech – Accula

We included one evaluation of the Accula assay with a total of 100 samples (50 SARS-CoV-2 positive; Hogan 2020; Figure 10). The study was laboratory-based and symptom status was not reported.

The study used nasopharyngeal samples in viral transport medium or saline, therefore the evaluation was not compliant with IFU requirements.

The sensitivity and specificity of the Accula test were 68.0% (95% CI 53.3% to 80.5%) and 100% (95% CI 92.9% to 100%; 100 samples, 50 cases; Table 5).



Sensitivity analysis of the impact of discrepant analysis

Six evaluations of molecular tests (in 1533 samples) reported results before and after discrepant analysis where selected samples were re-tested with either the same (Collier 2020; Harrington 2020; Moran 2020; Stevens 2020), or an alternative RT-PCR assay (Assennato 2020; Loeffelholz 2020). Four studies also reported retesting of samples with the index test (Assennato 2020; Collier 2020; Harrington 2020; Moran 2020; Appendix 16; Appendix 17).

Discrepant analysis reduces the number of samples deemed to be false negative or false positive errors. Discrepant analysis reduced the false negative proportion (1-sensitivity) from 2.1% to 0.8% and the false positive rate (1-specificity) from 2.2% to 0.4%. Three of the five studies reporting initially false positive results reported zero false positives after sample re-testing and one reported a drop in false positives from 11 to 3 (Loeffelholz 2020; Appendix 16). Three of the four studies that reported re-testing of initially false negative results reported reclassification as true negative on re-testing, and in the other the single false negative remained as a false negative. Given the bias inherent in choosing the reference test dependent on the observed results, we caution against these findings.

An additional study tested all samples with two different RT-PCR assays, and hence used a more accurate reference standard in all samples, not just samples with discrepant results (Moore 2020). Six initial true negatives were reclassified as false negatives after the second RT-PCR. Had discrepant analysis been undertaken these misclassifications would have been missed, further underlining the methodological flaws inherent to discrepant analysis.

Other sources of heterogeneity

We also planned to evaluate the effect of sample type and reference standard.

For sample type, the use of variable combinations of sample types with or without viral transport media created numerous sparse subgroups by sample type (Appendix 18). Instead we considered study compliance with manufacturer IFU requirements which is a more pragmatic classification.

All studies used RT-PCR alone as the reference standard for diagnosing SARS-CoV-2 infection.

Publication bias

We did not formally test for publication bias evident in the pattern of results, but did note that the identity of tests not meeting the PHE assessment criteria were not reported due to confidentiality agreements (PHE 2020(a)).

DISCUSSION

This is the second iteration of a Cochrane living review summarising the accuracy of point-of-care antigen and molecular tests for detecting current SARS-CoV-2 infection. This version of the review is based on published journal articles or studies available as preprints from 1 January 2020 up until 30 September 2020. In addition, we also included evaluations of antigen assays that were available as independent national reference laboratory publications or that were co-ordinated and published by FIND, and journal articles that were listed on the Diagnostics Global Health website to 16 November 2020.

Summary of main results

We included data from 77 studies using respiratory specimens, including 24,418 samples (7484 samples with confirmed SARS-CoV-2), and one study of faecal specimens (79 samples, 29 with confirmed SARS-CoV-2). Forty-eight studies (reporting 58 test evaluations) considered antigen tests; 30 studies (reporting 33 test evaluations) considered rapid molecular tests, including the single study (evaluation) in faecal samples. Key findings are presented in the Summary of findings 1.

We summarise six key findings from this review:

- 1. Despite a considerable increase in the number of studies evaluating point-of-care tests, particularly antigen tests, there are still no published or preprint reports of accuracy for a significant number of commercially produced point-of-care tests. This review located evaluations for 16 antigen tests (three of which we could not identify as available for purchase) and five molecular assays. These represent a small proportion of assays currently on the market (118 commercialised antigen tests and 53 molecular assays).
- 2. The new studies have more robust and appropriate study designs compared to those in the first version of this review. Particularly for antigen tests where there are now studies recruiting participants from community-based COVID-19 testing clinics. Reporting of key details, such as settings and symptom status have improved, and studies are now evaluating direct swab testing as would occur in a point-of-care setting. However, concerns about risk of bias and applicability of results remain, and further improvements in study methods and reporting are needed before strong conclusions can be drawn about the accuracy of many antigen and molecular tests reviewed here. As it is not known whether these limitations will lead to over- or underestimates of test accuracy, estimates should be cautiously interpreted in context of their methodological limitations and the settings in which they were conducted. More direct comparisons of test brands are needed, with evaluations undertaken in the intended use settings for these tests.

Particular methodological concerns include the use of deliberate sampling according to known presence or absence of SARS-CoV-2 infection; use of anonymised samples submitted to laboratories for routine RT-PCR testing (with no setting or participant details); and no information on symptoms or time from symptom onset. Differences in case-mix related to symptomatic status, time post-symptom onset and distribution of viral load are likely to have contributed to the observed variation in accuracy.

RT-PCR was the reference standard in all studies - no study defined the presence of COVID-19 using clinical or radiological features in the absence of a negative RT-PCR result.

3. Studies frequently did not follow the manufacturer's instructions or did not use the test at the point of care. Fewer than half conducted the tests according to the manufacturers' IFU (41% (37/91); 29/58 antigen test evaluations and 8/33 molecular test evaluations). Reasons for non-compliance included use of frozen samples, use of viral transport media, or lengthy intervals between sample collection and testing. Almost a third of studies (23/78) undertook on-site, direct swab testing immediately or within an hour of sample collection; trained laboratory staff conducted tests in 16 (21%) studies, and 31 (40%) studies did not clearly describe the test operator and setting for the test procedure but we inferred



that tests were carried out in a centralised laboratory setting, for example based on reported delays between collection and testing or reported use of archived or frozen samples.

4. For antigen test evaluations in symptomatic participants, we observed considerable heterogeneity in sensitivities (and to a lesser extent the specificities). Whilst the average sensitivity was 72.0% (95% CI 63.7% to 79.0%) and specificity was 99.5% (95% CI 98.5% to 99.8%), average sensitivity decreased with time since onset of symptoms, being higher in the first week (78.3%, 95% CI 71.1% to 84.1%) than when done later (51.095% CI 40.8% to 61.0%). Sensitivity was high in those with higher viral loads defined by Ct values ≤ 25 (94.5% 95% CI 91.0% to 96.7%) compared to those with lower viral loads (40.7%, 95% CI 31.8% to 50.3%). Focusing on studies that used the test in accordance with the manufacturer's instructions, sensitivities for different brands varied from 34% to 96% (either based on pooled results or single studies). WHO have set a minimum 'acceptable' sensitivity requirement of 80%, and acceptable and ideal (or 'desirable') specificity requirements of 97% and 99% respectively (WHO 2020c). Only one assay (SD Biosensor STANDARD Q) met the WHO acceptable criterion for sensitivity based on pooled results of several studies. One further test (BIONOTE NowCheck) also met the acceptable sensitivity criterion, but only one study evaluated it. Abbott Panbio met the sensitivity criterion in individual studies but not overall. The acceptable performance criterion of 97% specificity was also met for all three tests, and two tests met the desirable criterion of more than 99% specificity (Abbott Panbio and SD Biosensor STANDARD Q).

Considerable heterogeneity in sensitivities remained after restricting analyses by test brand and symptom status, suggesting an effect not only from participant characteristics but from setting, sample type and collection method, sample storage and preparation, and testing procedures that cannot be easily unpicked. The PHE studies included in this review allow some consideration of the effect of test operator experience on the accuracy of the Innova test although different samples were tested by each test operator such that only an indirect comparison of sensitivity can be made. Sensitivity increased from 57.5% (95% CI 52.3%, 62.6%; 372 samples) when testing was conducted on-site by trained non-healthcare workers (PHE 2020(c) [non-HCW tested]), to 70.0% (95% CI 63.5% to 75.9%; 223 samples) in samples tested onsite by healthcare workers ((PHE 2020(d) [HCW tested]), to 78.8% (95% CI 72.4% to 84.3%; 198 samples) for those tested by laboratory scientists (PHE 2020(d) [Lab tested]). The effect of test operator on accuracy has been observed for rapid diagnostic tests for other infectious diseases such as malaria (Boyce 2018; Landier 2018), and is worthy of further investigation for diagnosis of SARS-CoV-2.

5. Twelve studies evaluated the accuracy of antigen tests in asymptomatic people for detection of SARS-CoV-2 infection defined by PCR status. As discussed, this does not address the issue of whether the test is identifying those who are infectious (as there is no reference standard that can be used). The average sensitivity for detecting infection in asymptomatic participants was 58.1% (95% CI 40.2% to 74.1%) with specificity of 98.9% (95% CI 93.6% to 99.8%), both lower than in symptomatic people. Only half of studies reported clearly defined asymptomatic cohorts (e.g. preventive screening in the general population (n = 1), in returning travellers (n = 1), or in contacts of confirmed cases (n = 4)), the other six reported asymptomatic subgroups from mixed symptom

cohorts. Only one of the 12 studies provided data by viral load (Fenollar 2020(b)); 5% (1/22) of RT-PCR-positive samples had a Ct value of 25 or less, but 50% (11/22) had Ct values of 30 or less. No information on time after exposure to infection was reported.

6. For rapid molecular assays there were differences between test brands. Most data were for ID NOW and Xpert Xpress assays; average sensitivity for ID NOW was 78.6% (95% CI 73.7% to 82.8%) and Xpert Xpress 99.1% (95% CI 97.7% to 99.7%). Specificity for ID NOW was 99.8% (95% CI 99.23%, 99.9%) and Xpert Xpress 97.9% (95% CI 94.6% to 99.2%). These differences are beyond those expected by chance (P < 0.0001).

We were not able to investigate the effects of symptomatic status, or time from symptom onset: 12/29 were from symptomatic populations, three from 'mixed' symptomatic and asymptomatic populations (percentage from each group not reported), and the remaining 14 evaluations provided no information on symptom status (2/14 recruited from A&E and 12 were laboratory-based). These and other methodological limitations in the studies mean that we do not know how the assays would perform in any specific clinical setting when used in people suspected of having SARS-CoV-2 infection on the basis of symptoms, or of exposure to a confirmed case in the absence of symptoms. It is likely however that some difference in sensitivity between ID NOW and Xpert Xpress would be maintained in the absence of bias. The difference in specificity between the tests is small (ID NOW being 1.9% more specific compared to Xpert Xpress), but potentially important especially if used in a low-prevalence setting. However, this difference in specificity would not be an issue should testpositives be confirmed by a laboratory-based RT-PCR assay.

7. There are proposals for repeated use of antigen tests in different asymptomatic groups, such as school children and staff, hospital and care home workers, and even the general public, with a variety of different testing strategies. We found no data or studies evaluating the accuracy of any of these serial screening strategies.

We did not formally compare antigen with molecular assays because there were no head-to-head comparisons of the two test types. Instead, we illustrate predicted numbers of true positives, false positives, false negatives and true negatives, applying summary estimates of test accuracy to a hypothetical cohort of people suspected of SARS-CoV-2 infection across a range in prevalence of SARS-CoV-2 infection (Summary of findings 1). For both antigen and molecular assays, we only use summary data from evaluations conducted in accordance with manufacturers' IFUs, and for antigen tests we used separate results from symptomatic and asymptomatic participants.

Illustration of predicted effect of antigen testing by symptom status

For antigen test evaluations in symptomatic people, we selected three assays representing the range in observed average sensitivities: Coris Bioconcept COVID-19 Ag Respi-Strip (34.1% to 95% CI 29.7% to 38.8%), Abbott - Panbio Covid-19 Ag (75.1% to 95% CI 57.3% to 87.1%); and SD Biosensor - STANDARD Q COVID-19 Ag (88.1% to 95% CI 84.2% to 91.1%). Average specificities for the same three assays were 100% (95% CI 99.0% to 100%) to 99.5% (95% CI 98.7% to 99.8%) and 99.1% (95% CI 97.8% to 99.6%) respectively. Applied to a cohort of 1000 people with signs and symptoms of



COVID-19, in whom 50 people had confirmed infection (prevalence of 5%), for the three assays above we predicted that:

- 17, 43 or 53 people would have a positive test result, of which 0, 5 and 9 would be false positives (positive predictive values (PPV) 100%, 88.4% and 83.0%, respectively), and
- 33, 12 and 6 people with negative test results would be falsely negative (negative predictive values (NPV) 96.6%, 98.7%, and 99.4%).

Increasing the prevalence to 10% or 20%, increases PPV and decreases NPV. As there is considerable heterogeneity in the estimates of sensitivity, the values observed in practice could vary considerably from these figures as shown by the estimates derived from the confidence intervals (Summary of findings 1).

For antigen test evaluations in asymptomatic participants there was considerably less available data from IFU-compliant evaluations. We selected the same three exemplars, average sensitivities for identification of any infection (whether infectious or not) were lower than for symptomatic populations: 28.6% (95% CI 8.4% to 58.1%) for the Coris Bioconcept assay; 48.9% (95% CI 35.1% to 62.9%) for the Abbott assay; and 69.2% (95% CI 38.6% to 90.9%) for the SD Biosensor assay. Average specificities for the same three assays were: 100% (95% CI 88.8% to 100%), 98.1% (95% CI 96.3% to 99.1%), and 99.1% (95% CI 95.2% to 100%).

Applying the average values to a larger cohort of 10,000 people asymptomatic for COVID-19 and with a lower prevalence of 0.5% in whom 50 people had confirmed infection (infectious or not):

- 14, 213 or 125 individuals would have a positive test result of which 0, 189 and 90 would be false positives (PPVs of 100%, 11% and 28%, respectively), and
- 36, 26 and 15 people with negative test results would be falsely negative (NPVs 99.6%, 99.7%, and 99.8%).

We derived the summary estimates used in these calculations from asymptomatic participants identified for testing in a number of scenarios and they cannot be directly translated to a particular setting, such as mass screening, for example. The confidence intervals for the average estimates used in these calculations are also extremely wide for both sensitivities and specificities, such that the numbers of false positives and false negatives observed in practice could differ substantially from these figures. Increasing the prevalence of confirmed SARS-CoV-2 infection to 1% or 2% makes little difference to the absolute number of false positive results for these assays, but has a large relative effect when considered in relation to the number of positive test results (PPVs for the Abbott and SD Biosensor assays increasing to 40% and 61% at 2% prevalence).

Illustration of predicted effect of rapid molecular tests for symptomatic testing

For molecular assays, data from IFU-compliant evaluations were available for four of the five assays: ID NOW (Abbott Laboratories), Xpert Xpress (Cepheid Inc), SAMBA II (Diagnostics for the Real World) and COVID Nudge (DNAnudge). Average sensitivities were derived as 73.0% (95% CI 66.8% to 78.4%), 100% (95% CI 88.1% to 100%), 87.9% (95% CI 71.8% to 96.6%) and 94.4% (95% CI 86.2% to 98.4%). Average specificities were 99.7% (95% CI 98.7% to 99.9%),

97.2% (95% CI 99.4% to 99.3%), 97.4% (95% CI 92.6% to 99.5%) and 100% (95% CI 98.8% to 100%), respectively (Summary of findings 1).

Data by symptom status for these assays were very limited, therefore we assumed that the intended use is most likely to be for diagnosis of acute infection in symptomatic individuals and have applied the average estimates of accuracy to a hypothetical cohort of 1000 people, at prevalences of 5%, 10% and 20% (Summary of findings 1). If 50 of 1000 people had confirmed infection (5% prevalence):

- 40, 77, 69 and 47 individuals would have a positive test result of which 3, 27, 25 or 0 would be false positive (PPVs of 93.0%, 64.9%, 63.8%, and 100% respectively).
- 14, 0, 6 and 3 people with negative test results would be falsely negative (NPVs 98.6%, 100%, 99.4% and 99.7%).

Increasing the prevalence of confirmed SARS-CoV-2 infection to 10% or 20% has a large relative effect when considered in relation to the number of positive test results for both Xpert Xpress and SAMBA II (PPVs were 64.9% and 63.8% at 5% prevalence compared to 90.1% and 89.3% at 20% prevalence). Less variation in PPV was observed for ID NOW and COVID-Nudge because of the higher observed specificities. The NPV for the molecular assays is not affected to the same degree by these prevalence changes because of their relatively high sensitivities and the relatively low-prevalence scenarios being considered.

Across all exemplar assays in the Summary of findings 1, we observed the widest variation in NPV for the Coris Bioconcept antigen assay in symptomatic participants (86% to 97%), demonstrating that even in a low-prevalence setting, tests with poor sensitivity can have a considerable impact on the level of confidence that can be had in a negative test result.

Strengths and weaknesses of the review

Our review used a broad search screening all articles concerning COVID-19 or SARS-CoV-2. We undertook all screening and eligibility assessments, QUADAS-2 assessments (Whiting 2011), and data extraction of study findings independently and in duplicate. Although it is possible that the use of artificial intelligence text analysis to identify studies most relevant to diagnostic questions may have led to some eligible studies being missed, we believe that the multi-stranded search strategy used will have identified most if not all relevant literature. Whilst we have reasonable confidence in the completeness and accuracy of the findings up until the search date, should errors be noted please inform us at coviddta@contacts.bham.ac.uk so that we can verify and correct in our next update.

We undertook a careful assessment of sample preparation and biosafety requirements as well as time to test result, to ensure that included tests were suitable for use at the point of care. The application of these index test criteria led to the exclusion of 39 of the 85 studies that we excluded on the basis of the index tests evaluated. Evaluations of alternative laboratory-based molecular technologies are under consideration for inclusion in another review in our series of Cochrane COVID-19 diagnostic test accuracy reviews. Furthermore, for this iteration of the review, we explicitly considered whether the test evaluations were conducted in accordance with the manufacturer IFU, regarding the sample



types used, the use of viral transport medium and the permitted time between sample collection and testing.

We did not consider any manufacturer statements on the intended use of the tests by population, but we are aware that some IFUs recommend testing only in symptomatic people and within certain time frames after symptom onset (e.g. the Innova assay). Where possible, however, we did provide data separately for symptomatic and asymptomatic participants and identified clear trends towards lower sensitivities in asymptomatic individuals for detection of infection. We were unable to assess the accuracy of antigen tests for identification of infectious individuals, as there is no established reference standard for infectiousness (and it seems unlikely that one will ever be established). We have presented results by Ct value where it has been reported by the individual studies. We recognise the limitations from this approach, and given the extent to which RT-PCR Ct values vary between assays (Vogels 2020), and between laboratories, we strongly caution against the direct application of our results in high and low Ct value subgroups to any particular clinical context. There is no 'step change' in 'infectiousness' according to any fixed Ct value; increasing numbers of studies demonstrate successful viral culture in individuals considered to have 'low' viral load (Jaafar 2020; Singanayagam 2020), and, more importantly, that transmission of infection does occur from index cases with low RT-PCR Ct values (Lee 2021; Marks 2021). Ultimately, viral load on its own is only one factor influencing an individual's ability to transmit infection, 'infectiousness' being modified by host factors such as the health of an individual's immune system or presence of comorbidities, and environmental risk factors including closeness and length of contact with others.

Weaknesses of the review primarily reflect the weaknesses in the primary studies and their reporting. Although study quality improved in comparison to the first iteration of this review, many studies continue to omit descriptions of participants, and key aspects of study design and execution. In order to include data for all tests in pooled analyses we had to include some samples multiple times. We have been explicit about these issues where they arose. It is possible that eligible studies have been missed by our search strategy however we believe the risk to be very low considering our broad approach to identification of literature. Despite our best efforts to be as comprehensive as possible, new evaluations are continuously becoming available and it is impossible for any published and peer-reviewed systematic review to be fully up to date.

Around a quarter (18/78) of the studies we have included are currently only available as preprints, and as yet, have not undergone peer review. As published versions of these studies are identified in the future, we will double-check study descriptions, methods and findings, and update the review as required.

Applicability of findings to the review question

There are an increasing number of roles and testing strategies for which antigen and rapid molecular assays are considered, and it is likely that the performance of these tests needs to be considered separately for each of the use cases.

Our review shows that antigen tests do not appear to perform as well in asymptomatic populations compared to symptomatic populations for detecting infection. The amount of available data for asymptomatic populations is less than that from symptomatic

populations and is also based on asymptomatic individuals tested in a range of scenarios, from preventive or targeted screening, to contact tracing or testing at dedicated COVID-19 test centres, which may explain some of the observed variability. It is also not clear whether individuals in these studies were truly cases of asymptomatic infection as opposed to pre- or postsymptomatic, or were even mildly symptomatic and mislabelled as asymptomatic. Incomplete symptom assessment and lack of adequate follow-up to identify subsequent development of symptoms or previous history of symptoms can all contribute to inappropriate classification of individuals as asymptomatic infection (Meyerowitz 2020). As the studies in our review did not systematically attempt to identify pre- or post-symptomatic individuals, it may be more appropriate to consider the estimates for test accuracy for asymptomatic populations as primarily representing accuracy in those without clearly defined symptoms at the time of testing.

We are aware that several important studies in asymptomatic individuals have been reported since the close of our search. In mass screening in Liverpool, Innova was positive in 28 of 70 PCRdetected cases (sensitivity for infection 40.0%, 95% CI 28.5% to 52.4%) and 26 of 39 with Ct values less than 25 (sensitivity 66.7%, 95% CI 49.8% to 80.9%). Screening University of Birmingham students found 2 of 7185 students positive with Innova, and estimated sensitivity of 3.2% (95% CI 0.6% to 15.6%) for detecting any infection, 9.1% (95% CI 1.0% to 49.1%) for Ct values less than 30 and 100% (95% CI 15.8% to 100%) for Ct less than 25 (Ferguson 2020). BinaxNOW (which uses the same test strip as PanBio) has been tested in asymptomatic groups: in San Francisco the test detected 7 of 11 PCR-positive cases (sensitivity 63.6%, 95% CI 30.8% to 89.1%), and 6 of 6 with Ct values less than 30 (100%, 95% CI 54.1% to 100%; Pilarowski 2021); in a drive-through centre in Massachusetts it detected the virus in 70 of 107 in adults (sensitivity 65.4%, 95% CI 55.6 to 74.4) and 40 of 57 in children (70.2%, 95% CI 56.6% to 81.6%)); no breakdown by viral load is available (Pollock 2020). The specificity of the tests in all studies has remained high (above 99%). This selection of results is not based on a systematic search (this will occur in the next update) but these results suggest that emerging evidence is illustrating a range of sensitivity values for the ability of the tests to detect infection, with high detection rates only in groups with very high viral loads.

Given the superior test performance characteristics for symptomatic populations in the first week of symptoms and in those with higher viral loads, the observed poorer performance in those without symptoms is perhaps not surprising. Evidence suggests that higher viral loads are observed in the first week of illness, beginning two days prior to the development of symptoms (Cevik 2021). Viral load patterns in asymptomatic people are less clear but similarly high titers of SARS-CoV-2 have been observed at the onset of infection with a suggestion of faster clearance (Cevik 2021). However, variation in viral trajectories means that even if an asymptomatic person can identify a clear contact with a confirmed case of SARS-CoV-2 infection, it is not possible to pinpoint when (or even if) that individual will have a sufficient viral load to be detected on antigen testing. A serial testing policy would be likely to identify at least some infected asymptomatic contacts, but comes at the cost of increased numbers of false positives, especially in lowprevalence settings. There were no evaluations of serial testing in any of the studies.



For molecular tests, we observed a lack of studies undertaken in intended use settings, with most data being from laboratory testing. Although more evidence is available for accuracy in symptomatic people, applicability issues regarding the way in which the tests are carried out and in how cases of SARS-CoV-2 infection are defined remain, and it is not yet possible to determine how tests will perform in practice.

We recommend caution in applying the results outside of the individual study (or closely related) contexts and use case scenarios.

AUTHORS' CONCLUSIONS

Implications for practice

We consider the implications for practice for this review separately for symptomatic and for asymptomatic testing.

In the Role of index test(s) section, we suggested that for symptomatic individuals, and if sufficiently accurate, point-of-care testing could be used either to replace laboratory-based RT-PCR or as a triage to RT-PCR. As point-of-care tests are more accessible and provide a result more quickly than RT-PCR, theoretically their use may increase detection and speed up isolation and contact-tracing, leading to reduction in disease spread and reduce the burden on laboratory services.

The evidence included to date suggests that:

1. For diagnosis in symptomatic individuals in the first few days of symptoms, the most accurate rapid antigen tests are a useful alternative to laboratory-based RT-PCR where immediate results are required for timely patient management or where there are significant logistical or financial challenges in delivering RT-PCR in a timely manner. Rapid antigen tests are only sufficiently sensitive in the first week since onset of symptoms.

Antigen tests vary in sensitivity, and only those shown to meet appropriate criteria, such as WHO's priority target product profiles for COVID-19 diagnostics (i.e. sensitivity \geq 80% and specificity \geq 97%; WHO 2020c), could be considered as a rational substitute for RT-PCR.

Tests had high specificity, thus in symptomatic populations (where prevalence is likely to be high) the risk of false positives is low. At 80% sensitivity compared to RT-PCR, the probability that infected individuals are missed is 20% higher than for RT-PCR. Thus the possibility of false negative results should be considered in those with a high clinical suspicion of COVID-19, particularly if tested several days after onset of symptoms when viral load levels may have fallen.

2. Rapid antigen tests may be used simultaneously in combination with RT-PCR for symptomatic people, particularly where RT-PCR turn-around times are slow, to exploit the benefits of earlier results and consequent contact-tracing and isolation. Given the risk of false-negative results, isolation may be required until RT-PCR-negative results are obtained. Similarly, for investigation of local outbreaks, rapid antigen testing in a clearly defined population may establish cases and contacts that require isolation whilst awaiting results from RT-PCR.

In other circumstances rapid antigen tests may be used to triage to follow-on RT-PCR tests (rather than all receiving PCR tests) dependent on prevalence and the consideration of the consequences of false positive and false negative results.

Where prevalence is low, *positive* rapid test results require confirmatory testing to avoid unnecessary quarantine measures (PPVs around 85% to 90% for antigen assays mean that between 1 in 10 and 1 in 7 positive results will be falsely positive). If unverified, negative rapid test results should be delivered with appropriate advice on self-isolation procedures for the duration of symptoms in order to minimise the effect on transmission of infection from missed cases. RT-PCR tests should still be considered for people with a high clinical suspicion of COVID-19 and negative rapid test..

Where prevalence is higher (i.e. 20% or higher), false positives are less of a concern (PPVs are 96% to 100%) but the impact from false negative results becomes increasingly important and all test negatives may be considered for verification. At 20% prevalence, and using data for the more sensitive of our three exemplar assays, between 3% and 6% of those with negative rapid test results are missed cases of SARS-CoV-2 (24 to 50 cases missed out of a total of 200 cases). The lower the NPV the greater the potential effect on transmission of infection from missed cases and greater the impact from delays in commencement of contact tracing. For scenarios in which positive results do not have confirmatory testing, it is important that assays with high specificities (in the range of 99% to 100%) are selected in order to minimise the impact from false positive results at higher prevalences of disease.

3. We identified virtually no evidence for mass screening of asymptomatic individuals using rapid antigen tests in people with no known exposure. A small study screening travellers returning from high-risk countries (Cerutti 2020), identified only five SARS-CoV-2 infections (prevalence of 3%) with a reported sensitivity of antigen testing for detecting infection of 40%. However, important larger studies have been published since the end of our search, as mentioned above.

The key focus in mass screening is identification of individuals who are or will become infectious. PCR-positives define those who had detectable viral particles on their swab, which will include most of those who are or will become infectious, but also include individuals post-infection with residual viral particles. Without a reference standard for infectiousness, test accuracy studies cannot assess the ability of the test to detect the infectious subgroup of infections, and cannot provide evidence as to how well rapid antigen tests differentiate between individuals requiring isolation and those who provide no risk. The effectiveness of mass screening using these tests will only be established though outcome studies, such as cluster-randomised community trials.

Given the low false positive rate of rapid tests, when used in a period of outbreak, those found testing positive will have a high chance of being true positives, and thus the test can be used to identify cases requiring isolation. Consideration should be made as to whether test positives should be confirmed with PCR to identify false positives. With a 1% prevalence, a test with 40% sensitivity and 99.6% specificity would yield as many false positives as true positives.

However, the low and variable sensitivity, and lack of evidence that those who test negative are not, or will not become, infectious



indicates that those who are rapid antigen test-negative cannot be considered free of risk of being, or of becoming, infectious. In any screening or mass testing programme people testing negative may still have a non-negligible risk of infection.

4. We did not find any evidence of test accuracy in at-risk asymptomatic groups, such as contacts of confirmed cases, hospital workers, or during local outbreaks at schools, workplaces, or care homes. The impact of low-sensitivity tests in these settings is greater than in mass screening, as there will be higher numbers of false negatives, which could either create new outbreaks or will increase the severity of existing outbreaks. Positive cases will be more likely to be true positives than in mass screening settings.

5. We did not find any evidence evaluating the repeated use of tests. Although serial testing (over a number of days), or combinations of different rapid tests (e.g. an antigen test followed by a rapid molecular test) on the same sample are proposed to overcome the limitations of low test sensitivity, they all require validation. Use of multiple tests may increase false positive results, and there are likely to be many individuals with repeated false negative results reducing the expected benefit of subsequent tests. It is unlikely that models will be able to predict how well repeated tests and test combinations would work.

6. Some rapid molecular tests showed promising accuracy levels approximating those of laboratory-based RT-PCR and thus may have a role in small-capacity settings where obtaining test results within two hours will enable appropriate decision making. Results for Xpert Xpress, COVID Nudge and SAMBA II all showed high sensitivity and specificity. However, we identified methodological concerns with many of the evaluations such that we cannot be certain as to how the tests will perform when used in a point-of-care setting. Any application in practice should be accompanied with a proper evaluation to ascertain performance in real-world settings. Rapid molecular tests do not have all the logistical advantage of rapid antigen tests and the resource implications of their use at scale are potentially high, but they may be well suited for some testing scenarios. There is no evidence for use of rapid molecular tests in asymptomatic populations.

Our conclusions are in line with those in the first version of this review despite the increase in the evidence base. Ultimately, decisions around rapid testing will be driven not only by diagnostic accuracy but by acceptable levels of test complexity, time to result, access and acceptability to those being tested, and how test results influence individual behaviour, all of which might vary according to the setting in which the tests are to be used.

Implications for research

There is now a considerable volume of research for point-of-care tests for SARS-CoV-2 infection. However further well designed prospective and comparative evaluations of individual tests and test strategies in clinically relevant settings are urgently needed. Studies should recruit consecutive series of eligible participants and should clearly describe the clinical status, document time from symptom onset or time since exposure. Point-of-care tests must be conducted in accordance with manufacturer instructions for use, and across the spectrum of point-of care settings and test operators.

There needs to be evaluations of both individual tests and strategies of use of repeated tests. For molecular assays field trials are needed, not only to demonstrate test accuracy in these groups but acceptability and ease of use outside of centralised laboratories.

We observed a number of studies of molecular assays employing discrepant analysis to confirm the disease status of samples with false positive results in particular. There is a considerable risk of this type of selective re-testing leading to distorted results. If there is sufficient concern about the reliability of a single RT-PCR test then all samples should be tested with two RT-PCR assays. Finally, any future research study needs to be clear about eligibility and exclusion decisions throughout the whole diagnostic pathway, and should conform to the updated Standards for Reporting of Diagnostic Accuracy (STARD) guideline (Bossuyt 2015).

Consideration needs to be made of the best method for evaluating mass screening programmes. Whilst test accuracy studies help indicate which tests are likely to detect the greatest numbers of cases with the fewest false positives, assessing whether detecting asymptomatic cases leads to worthwhile reductions in disease spread will only be properly answered by studies of impact not accuracy.

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Albert 2020

Study characteristics	
Patient Sampling	Single group study estimating sensitivity and specificity: Patients with clinical suspicion of COVID-19 (compatible signs or symptoms appearing within the prior week) attending one of 8 primary care centres (n=412)
	Recruitment: Not stated; likely consecutive
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Primary care
	Location: 8 primary care centres of the Health Department Clínico-Malvarrosa in Valencia.
	Country: Spain
	Dates: Sep 2nd to Oct 7 2020
	Symptoms and severity: All symptomatic (<7 days p.s.o)
	Demographics: median age, 31 y (range, 1-91); 42% male 327 adults; median, 36 y (17-91y) 85 children; median, 11 y (1-16y)
	Exposure history: Not stated
Index tests	Test name: Panbio™ COVID-19 AG Rapid Test Device (no product code reported)
	Manufacturer: Abbott Diagnostic GmbH, Jena, Germany
	Antibody: Nucleoprotein
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; collected by trained nurses using flocked swabs
	Transport media: None for Ag testing
	Sample storage: None
	Test operator: Not stated; immediate testing

^{*} Indicates the major publication for the study



lbert 2020 (Continued)			
	Definition of test positivity: Visible line within 15 mins; As per manufacturer		
	Blinding reported: Yes		
	Timing of samples: Day <7 pso		
Target condition and reference standard(s)	Reference standard: RT-PCR; TaqPath COVID-19 Combo Kit (Thermo Fisher Scientific, Massachusetts, USA)		
	Definition of non-COVID cases: As for cases; single negative		
	Genetic target(s): ORF1ab, N and S genes		
	Samples used: NP in UTM		
	Timing of reference standard: As for index; tested within 24h		
	Blinded to index test: Not stated; presume Yes		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Simultaneous; paired		
	All patients received same reference standard: Yes		
	Missing data: None reported; no participant flow diagram reported		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Patients		
Comparative			
Notes	Funding: This work received no public or private funds. Abbott Diagnostics provided Panbio™ COVID-19 AG Rapid Test Device kits.		
	Publication status: Pre-print		
	Source: medRxiv		
	Author COI: The authors declare no conflicts of interest		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
	Yes		
Was a case-control design avoided?			
Was a case-control design avoided? Did the study avoid inappropriate exclusions?	Yes		



Albert 2020 (Continued)			
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	



Alemany 2020

Study characteristics		
Patient Sampling	Single group study including particpants from three settings: [1] symptomatic individuals with suspected COVID-19 seen in routine practice (n=446) [2] contacts exposed to positive PCR confirmed COVID-19 cases (n=473) [3] preventive screening of unexposed asymptomatic individuals in the general population (n=487)	
	Recruitment: Retrospective (frozen swabs)	
	Prospective or retrospective: Not stated	
Patient characteristics and setting	Setting: Mixed/Unclear (laboratory-based)	
	Location: Not reported; multiple author institutions reported	
	Country: Spain	
	Dates: Not stated	
	Symptoms and severity: Not stated; 15/1406 (1.1%) reportedly hospitalised (all PCR+) Viral load of cases: Ct <20: 258 (18.3%); Ct 20-24 305 (21.7%); Ct 25-29 285 (30.3%); Ct >30 (7.3%)	
	Demographics: All samples: mean age 40.4y (SD 24.5), 453 (32.2% male)	
	Exposure history: 473/1406 (33.6%) identified through contact tracing;	
Index tests	Test name: Panbio TM COVID-19 Ag Test (no product codes) [Selected following validation exercise using 40 NP samples to compare PanBio with Coris Bioconcept COVID-19 Ag RespiStrip, SD Biosensor Standard F COVID-19 Ag FIA and Standard Q COVID-19 Ag Test]	
	Manufacturer: Abbott Laboratories, Illinois, USA	
	Antibody: Not stated	
	Antigen target: SARS-CoV-2	
	Test method: CGIA	
	Samples used: [1] and [2] NP, [3] nasal mid-turbinate; collection not reported	
	Transport media: VTM (DeltaSwab Virus)	
	Sample storage: stored at 2-8C prior to PCR then frozen (-80C) prior to Ag testing; "Internal validation showed no significant change in the test performance using Abbot test Kit buffer or a mix of the Kit buffer and transport media at 1:3 dilution; likewise, the use of frozen specimens showed no significant differences compared with fresh ones"	
	Test operator: two laboratory technicians	
	Definition of test positivity: Visible line; as per manufacturer	
	Blinding reported: Yes	
	Timing of samples: Not stated	
Target condition and reference stan-	Reference standard: RT-PCR; in-house following CDC protocol	
dard(s)	Definition of non-COVID cases: As per cases; single negative PCR for absence of infection	
	Genetic target(s): Not stated; as per CDC protocol	



Alemany 2020 (Continued)			
	Samples used: NP or nasa	mid-turbinate; as per inde	ex test
	Timing of reference stands RT-PCR	ard: fresh samples stored a	at 2 – 8 °C for up to 72 hours prior to
	Blinded to index test: Yes;	conducted first	
	Incorporated index test: N	0	
Flow and timing	Time interval between ind	ex and reference tests: Sin	nultaneous (same swab)
	All patients received same	reference standard: Yes	
	Missing data: None report	ed; no participant flow dia	gram reported
	Uninterpretable results: N	one reported	
	Indeterminate results (ind	ex test): None reported	
	Indeterminate results (ref	erence standard)։ None rep	ported
	Unit of analysis: Patients		
Comparative			
Notes	Funding: The test kits were purchased to Abbott Rapid Diagnostics Healthcare SL (Spain). The funders of the study had no role in the study conception, design, conduct, data analysis, or writing of the report.		
	Publication status: Pre-pri	nt	
	Source: medRxiv		
	Author COI: Authors decla	e no conflicts of interest	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			High
	,		



Alemany 2020 (Continued)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular to	ests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	



Assennato 2020

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity and specificity: - samples from symptomatic individuals with suspected COVID-19 sent for routine laboratory diagnosis; supplied via PHE (n = 172)
	Recruitment: not stated
	Prospective or retrospective: retrospective
	Number of samples (samples with confirmed SARS-CoV-2): 172 (88)
Patient characteristics and setting	Setting: not stated; supplied by PHE
	Location: PHE, Cambridge Laboratory (samples from East of England)
	Country: UK
	Dates: not stated
	Symptoms and severity: symptomatic; no further details
	Demographics: not stated
	Exposure history: not stated
Index tests	Test name: SAMBA II SARS-CoV-2 Test
	Manufacturer: Diagnostics for the Real World
	Antigen target: ORF1ab, N2
	Antibody: N/A
	Test method: rapid PCR
	Samples used: combined nose and throat swab samples, provided as VTM
	Transport media: samples diluted 1:2 with SAMBA SCoV buffer
	Sample storage: not stated
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: as per manufacturer; either target present
	Blinding reported: yes; states that samples were rendered anonymous and provided blinded for the purpose of test validation
	Timing of samples: not stated
Target condition and reference standard(s)	Reference standard: RT-PCR; (1) Cambridge RdRp gene (Wuhan) assay on the Rotor gene Q real-time PCR assay routinely used by PHE; Ct ≤ 36 considered positive. (2) Samples al so tested with the PHE Colindale (Reference Laboratory) assay
	Definition of non-COVID cases: Single RT-PCR negative
	Genetic target(s): (1) RdRp, E gene, (2) RdRp 'different region'
	Samples used: combined nose and throat swab in VTM; same as for index test
	Timing of reference standard: not stated; Cambridge assay seems to have been part of routine testing near to time of sample collection; not clear if Colindale assay was at a later date after a period of storage



Assennato 2020 (Continued)	Bl. I I I I I I I I I I I I I I I I I I I			
	Blinded to index test: not		ambridge assay	
	Incorporated index test: n)		
Flow and timing		Time interval between index and reference tests: not stated; seems likely reference was carried out for routine diagnostic testing		
	All participants received so PCR tests)	nme reference standard: yo	es (all samples underwent both RT-	
	Missing data: none reporte	d, no participant flow dia	gram reported	
	Uninterpretable results: n	one reported		
	Indeterminate results (ind sults obtained on repeat	ex test): 3 FP and 1 FN resu	ılt retested using SAMBA-II; same re	
		lerline positive for ≥ 1 targ ssified as TP)	1 FN result were re-tested et gene on either Colindale or Cam ays	
	Unit of analysis: refers to p	articipants rather than sai	mples	
Comparative				
Notes	Funding: RKG is funded by WT108082AIA	Wellcome Senior Fellowsl	nip In Clinical Science award no	
	Publication status: preprir	t		
	Source: medRxiv			
	Author COI: no COI statem	ent reported; 3 co-authors	are affiliated to test manufacturer	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes		,	
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Antigen tests)				



Assennato 2020 (Continued) **DOMAIN 2: Index Test (Rapid molecular tests)** Were the index test results interpreted Yes without knowledge of the results of the reference standard? If a threshold was used, was it pre-speci-Yes fied? Could the conduct or interpretation of Low risk the index test have introduced bias? Are there concerns that the index test, Unclear its conduct, or interpretation differ from the review question? **DOMAIN 3: Reference Standard** Is the reference standards likely to cor-No rectly classify the target condition? Were the reference standard results inter-Yes preted without knowledge of the results of the index tests? Reference standard does not incorporate result of index test? Could the reference standard, its con-High risk duct, or its interpretation have introduced bias? Are there concerns that the target con-High dition as defined by the reference standard does not match the question? **DOMAIN 4: Flow and Timing** Was there an appropriate interval be-Yes tween index test and reference standard? Did all patients receive the same refer-Yes ence standard? Were all patients included in the analysis? Unclear Did all participants receive a reference Yes standard? Were results presented per patient? Yes

Could the patient flow have introduced

bias?

Unclear risk



Billaud 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity:
	- teachers (n=90) and students (n=419) screened for COVID-19 as part of a cluster investigation (n=509) $$
	Recruitment: Not stated; appears to be open to all
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Screening
	Location: College, Lyon
	Country: France
	Dates: September 16 and 17
	Symptoms and severity: 166/509, 32.6% symptomatic including 152/419 (36%) students
	Demographics: Mean, median age Students 21.6y, 21y (18 to 37y) Teachers 47.2y, 49y (26 to 64y)
	Exposure history: Outbreak investigation
Index tests	Test name: Described as "ABBOTT SARS-COV2 Antigenic Test"; presumed to be Panbio COVID-19 Ag Test
	Manufacturer: Abbott
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; collected by firefighters
	Transport media: None used
	Sample storage: n/a; tested immediately on site
	Test operator: Not stated
	Definition of test positivity: Visual line; as per manufacturer
	Blinding reported: Yes, performed first
	Timing of samples: Not stated but includes people >7 days pso
Target condition and reference standard(s)	Reference standard: RT-PCR; SARS-COV-2 (Thermofisher)
	Definition of non-COVID cases: As for cases; single negative
	Genetic target(s): Not stated
	Samples used: NP (paired)
	Timing of reference standard: As for index
	Blinded to index test: Not stated



Billaud 2020 (Continued)	Incorporated index test:	: No		
Flow and timing	Time interval between index and reference tests: Simultaneous			
	All patients received sar	me reference standard	: Yes	
	Missing data: 47 missing, including 11 uninterpretable			
	Uninterpretable results:	11 uninterpretable or	n Ag test	
	Indeterminate results (i	ndex test): None repor	ted	
	Indeterminate results (r	eference standard): N	one reported	
	Unit of analysis: Patient	S		
Comparative				
Notes	Funding: Not stated, pu	blic funding		
	Publication status: Publ	ished		
	Source: Report accessed	d via SFM Microbiologi	e website	
	Author COI: None			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability con- cerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Yes			
Did the study avoid inappropriate inclusions?	Yes			
Could the selection of patients have introduced bias?		Low risk		
Are there concerns that the included patients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Low risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear	



Billaud 2020 (Continued)

DOMAIN 2: Index Test (Rapid molecular tests)

DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	

Blairon 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: sampled from cohort of suspected COVID-19 patient samples sent for laboratory diagnosis (n=56) [Excluded data for full cohort, as only those with negative antigen test underwent confirmatory RT-PCR; of 912 submitted samples during time period, 776 remained after removing repeat tests and were reported in main study]
	Recruitment: Selection of 56 for verification analysis was not reported.
	Prospective or retrospective: prospectively
Patient characteristics and setting	Setting: Unclear; swabs obtained at hospital site (no further detail)
	Location: Not stated; author institution Iris Hospitals South, Brussels
	Country: Belgium
	Dates: April 5 - May 4 2020



Blairon 2020 (Continued)	
	Symptoms and severity: Not stated
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: COVID-19 Ag Respi-Strip (no product code reported)
	Manufacturer: Coris Bioconcept (Gembloux, Belgium)
	Antibody: Not stated
	Antigen target: Not stated
	Test method: LFA
	Samples used: NP swabs; collection not reported
	Transport media: Samples for antigen testing taken from UTM-RT swabs (Copan spa, Brescia, IT)
	Sample storage: No storage described; infer that antigen test was conducted immediately on receipt of sample at on-site laboratory 'after antigenic testing was performed, the molecular assessment of SARS-CoV-2 was outsourced to a university centre'
	Test operator: Not stated; infer laboratory staff
	Definition of test positivity: As per manufacturer
	Blinding reported: Not stated; infer yes as conducted prior to PCR confirmation
	Timing of samples: Not stated; appears to be on presentation (repeat tests ordered at clinician's discretion were excluded)
Target condition and reference standard(s)	Reference standard: qRT-PCR
	Definition of non-COVID cases: As above, single PCR negative to confirm absence of disease
	Genetic target(s): E gene
	Samples used: NP swabs (same as for Ag test)
	Timing of reference standard: Not stated
	Blinded to index test: Not stated
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Not stated but infer short interval; samples sent to university centre laboratory for PCR confirmation
	All patients received same reference standard: Yes (only if author confirms Ag+ also got PCR)
	Missing data: None reported; review team excluded main cohort data as no reference standard for antigen test positive samples
	Uninterpretable results: None reported; 1 'invalid' sample excluded from main cohort
	Indeterminate results (index test): None reported; 1 'non-conform' sample excluded from main cohort
	Indeterminate results (reference standard): None reported



Blairon 2020 (Continued)	Unit of analysis: Unclear; ed for separate group of 5		que patient samples but not report
Comparative			
Notes	Funding: None to declare		
	Publication status: Publis	hed	
	Source: Journal of Clinica	l Virology	
	Author COI: None to decla	ire	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have in- troduced bias?		Unclear risk	
Are there concerns that the included pa- tients and setting do not match the re- view question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tes	sts)		,
DOMAIN 3: Reference Standard			



Blairon 2020 (Continued)			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		
Could the patient flow have introduced bias?		High risk	

Broder 2020

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity: - samples positive on Roche cobas 6800 assay in lower range of viral load (E target Ct \geq 30) (n = 35)
	Recruitment: not stated; deliberate sampling according to viral load
	Prospective or retrospective: unclear
	Number of samples (samples with confirmed SARS-CoV-2): 35 (35)
Patient characteristics and setting	Setting: not stated
	Location: not stated; author institution Emory University School of Medicine, At lanta
	Country: USA



Broder 2020 (Continued)	Dates: not stated	
	Symptoms and severity: not stated; lower viral load	
	Demographics: not stated	
	Exposure history: not stated	
Index tests	Test name: GeneXpert Xpress SARS-CoV-2 assay (no product code reported)	
	Manufacturer: Cepheid	
	Antigen target: not stated E gene	
	Antibody: N/A	
	Test method: rapid PCR	
	Samples used: NP swabs in VTM	
	Transport media: not stated	
	Sample storage: within 3 days of initial testing (with RT-PCR)	
	Test operator: not stated; presume laboratory staff	
	Definition of test positivity: "all specimens were tested using the manufacturer protocol", no mention of presumptive positives	
	Blinding reported: not stated	
	Timing of samples: not stated	
Target condition and reference standard(s)	Reference standard: Roche cobas 6800 SARS-CoV-2 assay	
	Definition of non-COVID cases: N/A	
	Genetic target(s): E gene (unclear if other genetic targets as well)	
	Samples used: NP swabs (as for index test)	
	Timing of reference standard: not stated; presume on presentation	
	Blinded to index test: not stated; presume yes	
	Incorporated index test: no	
Flow and timing	Time interval between index and reference tests: same samples; index within 3 days of reference	
	All participants received same reference standard: yes	
	Missing data: none reported	
	Uninterpretable results: none reported, no participant flow diagram reported	
	Indeterminate results (index test): none reported	
	Indeterminate results (reference standard): discrepancies resolved using modified CDC RT-PCR; 1 FN confirmed as disease negative (i.e. a TN)	
	Unit of analysis: not stated; refers only to samples	
Comparative		
Notes	Funding: no funding described	



Broder 2020 (Continued)

Publication status: accepted manuscript

Source: Journal of Clinical Microbiology

Author COI: Dr. Kraft participated on a Roche advisory board regarding COVID

serology. All other study authors have no conflicts

Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		



Broo	ler 2020	(Continued)
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Diodei 2020 (continueu)			
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		

Unclear risk

Cerutti 2020

Could the patient flow have introduced bias?

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity in two cohorts: (1) symptomatic patients attending one of two Emergency departments (n=185) (2) asymptomatic travellers returning home from European high risk countries (Croatia, Spain, Malta) (n=145)
	Recruitment: (1) Random; (2) Not stated, presume consecutive
	Prospective or retrospective: Not stated
Patient characteristics and setting	Setting: Mixed; (1) Emergency department; (2) Possible contacts
	Location: (1) two Infectious Disease reference centres in North-Italy (ASL Citt`a di Torino, Turin and San Martino University Hospital, Genoa); (2) Not stated; samples sent to Microbiology and Virology Laboratory, Amedeo di Savoia Hospital, Torino
	Country: Italy
	Dates: (1) Mar 3 to May 1; (2) August 2020
	Symptoms and severity: Not stated; cohort (2) were asymptomatic
	Demographics: (1) mean age 44.6, 95 %CI: 40.7–48.6; (2) mean age 35.9, 95 % CI: 32.7–39.1
	Exposure history: (1) Not stated; (2) High risk country visit
Index tests	Test name: STANDARD Q COVID-19 Ag
	Manufacturer: SD-Biosensor, RELAB, I
	Antibody: NP



Cerutti 2020 (Continued)	Antigen target: Not stated	
	Test method: Not stated	
	Samples used: NP; collection not stated	
	Transport media: UTM (Copan, I)	
	Sample storage: Primarily run in parallel with standard of care RT-PCR; 13 were frozen residual samples	
	Test operator: Not stated; laboratory staff presumed	
	Definition of test positivity: Visual line after 15-30 mins; as per manufacturer.	
	Blinding reported: Not stated	
	Timing of samples: Not stated	
Target condition and reference standard(s)	Reference standard: RT-PCR; Seegene Allplex® 2019 n-CoV Assay (N = 159), DiaSorin Simplexa® (n = 28), and Cobas 6800 Roche® (N = 118).	
	Definition of non-COVID cases: Single negative	
	Genetic target(s): Not stated	
	Samples used: Not stated	
	Timing of reference standard: Not stated	
	Blinded to index test: Unclear	
	Incorporated index test: No	
Flow and timing	Time interval between index and reference tests: Simultaneous; not clear if same sample used or paired swabs obtained	
	All patients received same reference standard: Yes; different assays	
	Missing data: None reported; no participant flow diagram reported	
	Uninterpretable results: None reported	
	Indeterminate results (index test): None reported	
	Indeterminate results (reference standard): None reported	
	Unit of analysis: Patients	
Comparative		
Notes	Funding: Authors thank RELAb for the donation of the STANDARD Q COVID-19 SD-Biosensor kits to pursue the study. No other specific grant from public funding agencies was received.	
	Publication status: Published	
	Source: J Clin Virol	
	Author COI: The authors report no declarations of interest.	
Methodological quality		
Item	Authors' judgement Risk of bias Applicability concerns	



Cerutti 2020 (Continued)

Cerutti 2020 (Continued) DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have intro- duced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests))		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			



Could the patient flow have introduced bias?	Unclear risk
Were results presented per patient?	Yes
Did all participants receive a reference standard?	Yes
Were all patients included in the analysis?	Unclear
Did all patients receive the same reference standard?	Yes
Was there an appropriate interval between index test and reference standard?	Yes
Cerutti 2020 (Continued)	

Chen 2020a

Study characteristics		
Patient Sampling	Single group study using: - archived paired samples from COVID-19 inpatients (n=58). Aim is to compare diagnos tic yield between saliva and NP swabs but can also extract sensitivity for each using rapid test.	
	Recruitment: Not stated	
	Prospective or retrospective: Retrospective	
Patient characteristics and setting	Setting: In-patients	
	Location: Queen Mary Hospital, Pokfulam, Hong Kong	
	Country: People's Republic of China	
	Dates: Not stated	
	Symptoms and severity: Not stated	
	Demographics: Median age 38 y; 28, 48% male	
	Exposure history: Not stated	
Index tests	Test name: Xpert Xpress SARS-CoV-2 assay (no product codes reported)	
	Manufacturer: Cepheid, Sunnyvale, CA, USA	
	Target gene(s): E and N2 gene	
	Antigen target: n/a	
	Test method: Automated RT-PCR	
	Samples used: NP, saliva (posterior oropharyngeal, self-collected by clearing the throa and spitting c1 mL saliva directly into a sterile bottle in the early morning before mout rinsing and breakfast)	
	Transport media: Both sample types immersed in 2ml of viral transport solution	
	Sample storage: Not stated; archived	



Item	Authors' judgement	Risk of bias	Applicability concerns	
Methodological quality				
	Author COI: No potential conflict of interest was reported by the author(s); Xpert Xpress cartridges provided by the test manufacturer via an Investigator-Initiated Study agreement (Cepheid-IIS-2020-0009).			
	Source: Emerging microbes and infections			
	Publication status: Publish	ed		
	oratory Surveillance of Em timicrobial Resistance, and search Grants Council, the Hong Kong Michael Seak-K Foundation Limited, Hui M	erging Infectious Diseases a I the Theme-Based Researd donations of Richard Yu an an Tong, May Tam Mak Mei ing, Hui Hoy, and Chow Sin able Foundation, Marina M	and Research Capability on An- ch Scheme (T11/707/15) of the Re- d Carol Yu, the Shaw Foundation Yin Respiratory Viral Research Lan Charity Fund Limited, Chan an-Wai Lee, the Jessie & George Ho	
Comparative Notes	Funding: This study was pa	rtly supported by Consulta	ncy Services for Enhancing Lab-	
C	Unit of analysis: Patients			
	Indeterminate results (refe	rence standard): None repo	orted	
	Indeterminate results (inde			
	Uninterpretable results: No	ot stated		
	Missing data: None reporte tive only on saliva excluded		ram reported. THree samples posi-	
	All patients received same	reference standard: Yes		
Flow and timing	Time interval between index and reference tests: Simultaneous; same samples			
	Incorporated index test: No)		
	Blinded to index test: Not s	tated; infer yes		
	Timing of reference standa	rd: Not stated; prior to inde	ex test	
	Samples used: same as ind	ex test		
	Genetic target(s): RdRp			
	RT–PCR assay Definition of non-COVID ca	ses: n/a only cases include	d	
Target condition and reference standard(s)	Reference standard: in-hou (RdRp/Hel) real-time	Reference standard: in-house SARS-CoV-2 RNA dependent RNA polymerase/ Helicase		
	Timing of samples: Not sta	ted		
	Blinding reported: Not stat	ed;		
	Definition of test positivity no mention of presumptive		ing to manufacturer's instruction'	
	Test operator: Not stated; i	nfer laboratory staff		



Chen 2020a (Continued)

Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions?	
Patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Unclear	
Did the study avoid inappropriate exclusions?	
sions?	
Did the study avoid inappropriate inclu- Unclear sions?	
Could the selection of patients have introduced bias? High risk	
Are there concerns that the included High patients and setting do not match the review question?	
DOMAIN 2: Index Test (Antigen tests)	
DOMAIN 2: Index Test (Rapid molecular tests)	
Were the index test results interpreted unclear without knowledge of the results of the reference standard?	
If a threshold was used, was it pre-specified?	
Could the conduct or interpretation of Unclear risk the index test have introduced bias?	
Are there concerns that the index test, High its conduct, or interpretation differ from the review question?	
DOMAIN 3: Reference Standard	
Is the reference standards likely to cor- rectly classify the target condition?	
Were the reference standard results interpreted without knowledge of the results of the index tests?	
Reference standard does not incorporate Yes result of index test?	
Could the reference standard, its conduct, or its interpretation have introduced bias?	
Are there concerns that the target condition as defined by the reference standard does not match the question?	



Chen 2020a (Continued)

Could the patient flow have introduced bias?	Unclear risk
Were results presented per patient?	Yes
Did all participants receive a reference standard?	Yes
Were all patients included in the analysis?	Unclear
Did all patients receive the same reference standard?	Yes
Was there an appropriate interval between index test and reference standard?	Yes

Collier 2020

Collier 2020			
Study characteristics			
Patient Sampling	Single group study to estimate sensitivity and specificity: suspected COVID-19 patients admitted with a possible diagnosis of COVID-19 (n=149)		
	Recruitment: Consecutive		
	Prospective or retrospective: prospectively		
Patient characteristics and setting	Setting: In-patients		
	Location: Cambridge University Hospitals NHS Foundation Trust		
	Country: UK		
	Dates: April 6 - May 2 2020		
	Symptoms and severity: Not stated		
	Demographics: Mean age 62.7 y, 70, 47% male		
	Exposure history: Not stated		
Index tests	Test name: SAMBA II SARS-CoV-2 test (no product code reported)		
	Manufacturer: Diagnostics for the Real World (DRW), University of Cambridge, Cambridge		
	Target gene(s): Orf1 and the E genes		
	Antigen target: n/a		
	Test method: RT-PCR		
	Samples used: combined nasal/throat swab (NOP) on dry sterile swab. Collection not reported		
	Transport media: None used; samples inactivated in SCov buffer prior to testing		
	Sample storage: Not stated. Test performed within 18 hours of reference test		
	Test operator: Not stated; infer laboratory staff		



Collier 2020 (Continued)				
	Definition of test positivity: As per manufacturer			
	Blinding reported: Unclear; yes if always conducted before reference test but not explicitly described, i.e. 'SAMBA swab must be taken within 18 hours of the standard laboratory swab'			
	Timing of samples: Not stated; appears to be on presentation/admission but no further details			
Target condition and reference	Reference standard: RT-PCR; in-house PHE assay			
standard(s)	Definition of non-COVID cases: As above, single PCR negative to confirm absence of disease			
	Genetic target(s): Not stated			
	Samples used: Not stated; separate swab used as participants were excluded if >18h interval between swab collections			
	Timing of reference standard: Not stated			
	Blinded to index test: Yes; 'The results of the SAMBA II SARS-CoV-2 was not known to the assessors of the standard lab RT-PCR prior.' Not stated. Possibly if done prior to index test.			
	Incorporated index test: No			
Flow and timing	Time interval between index and reference tests: <18 hours			
	All patients received same reference standard: Yes			
	Missing data: Yes; 5 discarded VTM, 1 timing of PHE swab not reported, 1 inadequate SAMBA swab, 2 interval between swabs >24h			
	Uninterpretable results: None reported			
	Indeterminate results (index test): Not stated 'Indeterminate SAMBA II SARS CoV-2 tests were repeated with a 1:2 dilution of sample to inactivation buffer according to manufacturer standard operating procedures until a valid result was obtained.' Discrepant results between index and reference were also re-tested using SAMBA-II on original samples			
	Indeterminate results (reference standard): 1 false negative Indeterminate standard lab RT PCR tests were repeated on a replicate nose/throat swab until a valid result was obtained. Discrepant results between index and reference were re-tested using RT-PCR on original sam- ples, with reference to clinical notes to determine clinical suspicion. Remaining discrepant re- sults were re-tested using alternative sample, i.e. sample in SCov buffer tested on RT-PCR and sample in VTM tested on SAMBA-II			
	Unit of analysis: Patients			
Comparative				
Notes	Funding: The Wellcome Trust (Senior Research Fellowship to RKG WT108082AIA and PhD Research Fellowship to DAC; Principal Research Fellowship 210688/Z/18/Z to PJL), Addenbrooke' Charitable Trust to PJL, National Institute of Health Research (NIHR) Cambridge BRC			
	Publication status: Pre-print and published version (25-8-20)			
	Source: Pre-print; Cell Reports Medicine			
	Author COI: Pre-print - Dr. Besser reports personal fees from STAGO, personal fees from Novartis, personal fees from Cosmopharma, personal fees from Werfen, personal fees from Agios, grants from Mitsubishi Pharma, outside the submitted work; RKG reports fees from ad hoc con sulting from ViiV, Gilead and UMOVIS.			



Collier 2020 (Continued)

Published version - The authors declare no competing interests (Three co-authors affiliated to test manufacturer)

Methodological quality

местойоюдісаї quaiity			
Item	Authors' judgement		Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen test	ts)		
DOMAIN 2: Index Test (Rapid mole	cular tests)		
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it prespecified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		



Collier 2020 (Continued)				
Reference standard does not in- corporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	No			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	No			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow have introduced bias?		High risk		
Courtellemont 2020				
Study characteristics				

Courtellemont 2020	
Study characteristics	
Patient Sampling	Unclear design estimating sensitivity and specificity (coded as two group because of deliberate sampling of PCR positive cases): (1) Symptomatic (headache, fatigue, fever, or respiratory signs) or asymptomatic people voluntarily accessing the COVID-19 Screening Department (n=231) (2) hospitalized SARS-CoV-2 positive patients (n=17)
	[review team excluded 20 cases with a previous positive RT-qPCR within 5 days but a negative RTqPCR at the time of study sampling]
	Recruitment: Unclear
	Prospective or retrospective: Unclear
Patient characteristics and setting	Setting: Mixed
	Location: COVID-19 Screening Department and SARS CoV-2 positive patients hospitalized in the Infectious Diseases Department of the Centre Hospitalier Régional (CHR) of Orléans, France, or the Department of Infectious and Tropical Diseases of the Centre Hospitalier Universitaire (CHU) Tenon, Paris



Courtellemont 2020 (Continued)			
	Country: France		
	Dates: Oct 12 to Oct 19		
	Symptoms and severity: 99/121, 82% cases were symptomatic; 22 asymptomatic		
	Demographics: median age 38y, mean age 43y (range: 18-96), 117, 47% male		
	Exposure history: Not stated		
Index tests	Test name: COVID-VIRO®		
	Manufacturer: AAZ, Boulogne Billancourt, France		
	Antibody: Nucleocapsid		
	Antigen target: monoclonal		
	Test method: CGIA		
	Samples used: NP; collected by trained personnel (nurse, doctors, or biologist); subgroup also had OP or saliva collected		
	Transport media: Direct testing for Ag test		
	Sample storage: None		
	Test operator: Not stated		
	Definition of test positivity: Visible line; As per manufacturer		
	Blinding reported: Yes		
	Timing of samples: median 5 days pso, mean 5.3 days, range 1 to 20d		
Target condition and reference standard(s)	Reference standard: RT-PCR; TaqPath Covid-19 Multiplex RT-PCR, Thermofisher		
	Definition of non-COVID cases: single negative PCR		
	Genetic target(s): ORF1ab, S and N genes		
	Samples used: NP in VTM; paired		
	Timing of reference standard: As for index		
	Blinded to index test: Not stated		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Simultaneous; paired		
	All patients received same reference standard: Yes		
	Missing data: None reported, no participant flow diagram reported; review team excluded 20 cases with a previous positive RT-qPCR within 5 days but a negative RTqPCR at the time of study sampling		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Patients		



Courte	llemont 2020	(Continued)
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Notes

Funding: No funding statement reported

Publication status: Preprint

Source: medRxiv

Author COI: No COI statement reported

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		



Courtellemont 2020 (Continued)			
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Cradic 2020(a)

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: - symptomatic patients suspected of COVID-19 that met criteria for testing, either presenting to ED or as inpatients at single hospital (n=184)
	Recruitment: Not stated
	Prospective or retrospective: Prospective
	[Second cohort of paired samples from patients presenting to ED with signs/symptoms of COVID-19 submitted for routine laboratory testing (n=182), extracted as Cradic 2020(b)]
Patient characteristics and setting	Setting: Mixed (ED/inpatients)
	Location: OhioHealth Riverside Methodist Hospital, Columbus
	Country: USA
	Dates: Not stated
	Symptoms and severity: All symptomatic, no further details.
	Demographics: Not stated
	Exposure history: Not stated



Cradic 2020(a) (Continued) Index tests Test name: [A] ID NOW COVID-19 EUA [Study also evaluates [B] Diasorin Simplexa and [C] Roche cobas 6800 SARS-CoV-2; not eligible for this review] Manufacturer: Abbott Laboratories Target gene(s): RdRp Antigen target: n/a Test method: Isothermal PCR Samples used: NP swabs in UTM; collected on flocked swab, no other details, Transport media: 3 mL of sterile UVT (Becton Dickinson) Sample storage: asap, or stored for up to 72 hours at 2°C to 8°C. Following routine testing, samples were stored frozen (≤–80°C) until comparator testing with the Roche cobas assay could be completed Test operator: Not stated; infer laboratory staff. Definition of test positivity: as per manufacturer Blinding reported: Not stated Timing of samples: Unclear, infer upon presentation Target condition and reference standard(s) Reference standard: Composite reference standard, defined as the result obtained from at least 2 of the 3 assays conducted (Abbot ID NOW, Diasorin Simplexa or Roche cobas 6800 SARS-CoV-2) Definition of non-COVID cases: Same as index test; single negative for absence disease Genetic target(s): RdRp, S or ORF1ab gene (either present), ORF1ab or E gene (both present for +ve, either present for presumptive +ve) Samples used: Same as index test Timing of reference standard: Not stated Blinded to index test: No (>=2 +ve) Incorporated index test: Yes Flow and timing Time interval between index and reference tests: Simultaneous - same swab All patients received same reference standard: Yes Missing data: None reported, no participant flow diagram reported Uninterpretable results: None reported Indeterminate results (index test): None reported Indeterminate results (reference standard): None reported Unit of analysis: Patients Comparative Notes Funding: No funding statement reported

Publication status: published



Cradic 2020(a) (Continued)

Source: American Journal of Clinical Pathology

Author COI: No COI statement reported

	Author COI: No COI statement reported		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Reference standard does not incorporate result of index test?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	



Crad	ic 2020(a)	(Continued)
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Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Yes
Did all patients receive the same reference standard?	No
Were all patients included in the analysis?	Unclear
Did all participants receive a reference standard?	Yes
Were results presented per patient?	Yes
Could the patient flow have introduced bias?	High risk

Cradic 2020(b)

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: paired samples from patients presenting to ED with signs/symptoms of COVID-19 submitted for routine laboratory testing (n=182)
	Recruitment: Not stated
	Prospective or retrospective: Prospective
	[Second cohort of symptomatic patients suspected of COVID-19 that met criteria for testing, either presenting to ED or as inpatients at single hospital (n=184), extracted as Cradic 2020(a)]
Patient characteristics and setting	Setting: Emergency department
	Location: OhioHealth Laboratory Services, Columbus (presume ED at Ohio-Health Riverside Methodist Hospital)
	Country: USA
	Dates: Not stated
	Symptoms and severity: All symptomatic, no further details.
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: [A] ID NOW COVID-19 EUA [Study also evaluates [B] Diasorin Simplexa and [C] Roche cobas 6800 SARS-CoV-2; not eligible for this review]
	Manufacturer: Abbott Laboratories
	Target gene(s): RdRp



Methodological quality	Authors' judgement Risk of bias Applicability concerns
Methodological quality	
	Author COI: No COI statement reported
	Source: American Journal of Clinical Pathology
	Publication status: published
Notes	Funding: No funding statement reported
Comparative	
	Unit of analysis: Patients
	Indeterminate results (reference standard): None reported
	Indeterminate results (index test): None reported
	Uninterpretable results: None reported
	Missing data: None reported, no participant flow diagram reported
	All patients received same reference standard: Yes
Flow and timing	Time interval between index and reference tests: Simultaneous; paired swabs
	Incorporated index test: No
	Blinded to index test: Not stated
	Timing of reference standard: Not stated
	Samples used: NP swab in UTM
	disease Genetic target(s): S or ORF1ab gene (either present)
	Definition of non-COVID cases: Same as index test; single negative for absence
Target condition and reference standard(s)	Reference standard: RT-PCR; Diasorin Simplexa
	Timing of samples: Unclear, infer upon presentation
	Blinding reported: Not stated
	Definition of test positivity: as per manufacturer
	Test operator: Not stated; infer laboratory staff.
	Sample storage: not stated
	tions) Transport media: presume as above for NP in UTM
	Samples used: NP swabs in UTM (collected as part of standard of care), plus direct testing of OP swabs and of nasal swabs (collected according to CDC instruc
	Test method: Isothermal PCR



Unclear		
Yes		
Unclear		
Yes		
	Unclear risk	
		Low concern
Unclear		
Yes		
	Unclear risk	
		Unclear
No		
Unclear		
Yes		
	High risk	
		High
Yes		
	Yes Unclear Yes Unclear Ves Unclear Yes No Vocation of the properties of the p	Yes Unclear Yes Unclear risk Unclear Unclear Ves Unclear High risk



Cradic 2020(b) (Continued)		
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Unclear	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		Unclear risk

Diao 2020

Study characteristics	
Patient Sampling	Single group estimating sensitivity and specificity for detecting active disease - samples from cases of suspected SARS-CoV-2 infection (n = 239)
	Recruitment: not stated if participants were consecutive
	Prospective or retrospective: retrospective
	Number of samples (samples with confirmed SARS-CoV-2): 239 (208)
Patient characteristics and setting	Setting: hospital (inpatients)
	Location: 7 centres, including General Hospital of Central Theatre Command, Wuhar No.7 People's Hospital, Wuhan Pulmonary Hospital, Hubei Maternal and Child Hospital, Taikang Hospital, Hanyang Hospital and Wuguo Hospital. Urine study done in Southwest Hospital in Chongqing
	Country: China
	Dates: not stated
	Symptoms and severity: not stated
	Demographics: not stated
	Exposure history: not stated
ndex tests	Test name: not stated
	Manufacturer: in house (but study authors affiliated to Bioeasy Technology)
	Antibody: monoclonal antibody
	Antigen target: nucleocapsid protein (N-antigen)
	Test method: FIA (fluorescence immunochromatographic); requires immunofluorescence analyser
	Samples used: NP (all), urine (subgroup)
	Transport media: samples diluted and mixed in 500 μL saline solution; 100 μL transferred to the sample well of the test card
	Sample storage: not reported
	Test operator: not stated; presume laboratory staff



Diao 2020 (Continued)	Definition of test positivity: cut-off value was determined by testing 100 nasal swab		
	samples of healthy people and calculated as the mean value of the fluorescence signal plus 5 SD.		
	Blinding reported: done in parallel; blinded		
	Timing of samples: not stated		
Target condition and reference standard(s)	Reference standard: RT-PCR (Daan Gene kit); performed on ABI Prism 7500 and Light Cycler 480 real-time PCR system. Threshold < 40 Ct; threshold < 30 Ct also investigated Definition of non-COVID cases: all participants underwent 3 nucleic acid tests, and the results of each nucleic acid test were verified by 2 COVID-19 nucleic acid test kits.		
	Genetic target(s): ORF1ab and N gene		
	Samples used: NP swab, same as for index test		
	Timing of reference standard: not stated		
	Blinded to index test: done in parallel; blinded		
	Incorporated index test: no		
Flow and timing	Time interval between index and reference tests: done in parallel		
	All participants received same reference standard: yes		
	Missing data: not reported, no participant flow diagram reported		
	Uninterpretable results: not reported		
	Indeterminate results (index test): none reported		
	Indeterminate results (reference standard): none described		
	Unit of analysis: participants		
Comparative			
Notes	Funding: this research was supported by grants from National Key R&D Program of China (2016YFA0502204); Chongqing Health Commission COVID-19 Project (2020ZX01).		
	Publication status: preprint (not peer-reviewed)		
	Source: medRxiv preprint		
	Author COI: study authors declare no COI present; 1 affiliated to Shenzhen Bioeasy Biotechnology Co. Ltd.		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		



Diao 2020 (Continued)			
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tes	sts)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		



Diao	202	(Continued)	
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Did all participants receive a reference standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

Unclear risk

Dust 2020

Study characteristics		
Patient Sampling	Design unclear; coded as two group study: [1] SARS-CoV-2 positive samples submitted for routine viral diagnostic testing (n=20 evaluated with Xpert Xpress) [2] samples positive for other respiratory infection from those submitted for routine viral diagnostic testing (n=18) (Sampled from total n of 177; 65 SARS-CoV-2 positive, 112 SARS-CoV-2 negative including 57 positive for other respiratory viruses) [Study also reports results for reference panel of simulated specimens; not extracted for this review)	
	Recruitment: Convenience	
	Prospective or retrospective: Retrospective	
Patient characteristics and setting	Setting: Unclear; submitted to laboratory	
	Location: Cadham Provincial Laboratory (CPL), Manitoba	
	Country: Canada	
	Dates: Not stated	
	Symptoms and severity: Not reported	
	Demographics: Not reported	
	Exposure history: Not reported	
Index tests	Test name: Xpert Xpress (no product code) [also evaluates cobas SARS-CoV-2 RT-PCR (Roche) and three in-house RT-PCR assays; not eligible for this review]	
	Manufacturer: Cepheid Inc	
	Antibody: E, N2	
	Antigen target: n/a	
	Test method: automated RT-PCR	
	Samples used: NP swabs in VTM; collection not reported	
	Transport media: VTM; no further detail	
	Sample storage: Not stated, could be archived samples	
	Test operator: Not stated	



Dust 2020 (Continued)	Definition of test positivitive positives not mentio		as per manufacturer (presump-	
	Blinding reported: Not st	•		
	Timing of samples: Not s			
Target condition and reference standard(s)	Reference standard: In-h	ouse RT-PCR (extraction (Thermo Scientific™) an	n with MagMAX™ reagents on a d RT-PCR performed on a Bio- nreshold NR	
	Definition of non-COVID cases: As for cases; single negative			
	Genetic target(s): E, N1			
	Samples used: NP (as for	index)		
	Timing of reference stan	dard: Not stated		
	Blinded to index test: No	t stated		
	Incorporated index test:	No		
Flow and timing	Time interval between in	dex and reference tests	: Simultaneous (same swab)	
	All patients received sam	ne reference standard: Y	es	
	Missing data: None repor	ted, no participant flow	diagram reported	
	Uninterpretable results: None reported			
	Indeterminate results (index test): None reported			
	Indeterminate results (reference standard): None reported			
	Unit of analysis: Not stated			
Comparative				
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection		,		
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	No			
Did the study avoid inappropriate exclusions?	No			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	



Dust 2020 (Continued)

DOMAIN 2: Index	(Test	(Antigen	tests)
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DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?		U	nclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?		н	igh
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		
Could the patient flow have introduced bias?		Unclear risk	

Fenollar 2020(a)

Study characteristics



Fenollar 2020(a) (Continued)			
Patient Sampling	Two cohorts of patients presenting for COVID-19 testing at the same institution. This extraction relates to: [1] Single group study to estimate sensitivity alone: symptomatic patients, all PCR positive (n=182) Fenollar 2020(b) reports data for [2] Single group study to estimate both sensitivity and specificity: asymptomatic contacts of confirmed cases (n=159)		
	Recruitment: Prospective		
	Prospective or retrospective: Unclear		
Patient characteristics and setting	Setting: Unclear; COVID-19 testing		
	Location: Institut Hospitalo-universitaire Méditerranée Infection, Marseille,		
	Country: France		
	Dates: Sep 21 to Oct 2 2020		
	Symptoms and severity: Not stated; all symptomatic Ct values for 154 pts: Ct <=20: 58, 38%; Ct 21-25: 49, 32%; Ct 26-30: 39, 25%; Ct 31-34: 8, 5%		
	Demographics: Not reported		
	Exposure history: [1] Not stated		
Index tests	Test name: Panbio COVID-19 Ag		
	Manufacturer: Abbott		
	Antibody: NP		
	Antigen target: Not stated		
	Test method: Not stated		
	Samples used: NP		
	Transport media: Not stated; appears to be direct testing		
	Sample storage: Tested within 1 hour		
	Test operator: Not stated		
	Definition of test positivity: Visual line; as per manufacturer		
	Blinding reported: Not stated, but presume yes as conducted within 1h of collection		
	Timing of samples: Not stated		
Target condition and reference standard(s)	Reference standard: Automated RT-PCR; VitaPCR (Credo diagnostics, Singapore)		
	Definition of non-COVID cases: n/a		
	Genetic target(s): Not stated		
	Samples used: NP (paired, from opposite nostril)		
	Timing of reference standard: Not stated		
	Blinded to index test: Unclear		
	Incorporated index test: No		



Fenollar 2020(a) (Continued)				
Flow and timing	Time interval between index and reference tests: Simultaneous; paired swabs			
	All patients received sam	ne reference standard: \	'es	
	Missing data: None repo	rted		
	Uninterpretable results:	None reported, no part	icipant flow diagram reported	
	Indeterminate results (ir	idex test): None reporte	d	
	Indeterminate results (re	eference standard): Nor	e reported	
	Unit of analysis: Patients	i .		
Comparative				
Notes	Funding: Supported by the Méditerranée-Infection Foundation and the French Agence Nationale de la Recherche under reference Investissements d'Avenir Méditerranée Infection 10-IAHU-03 and Région Provence-Alpes-Côte d'Azur and European funding FEDER IHUBIOTK.			
	Source: Accepted manus	script		
	Author COI: Pr Raoult and Pr Drancourt are co-founders of the Pocrame startu that develops diagnostic devices for infectious diseases			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	No			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen tests)	,			

Were the index test results interpreted without

knowledge of the results of the reference stan-

If a threshold was used, was it pre-specified?

dex test have introduced bias?

Could the conduct or interpretation of the in-

dard?

Low risk

Yes



enollar 2020(a) (Continued)	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?	Unclear
DOMAIN 2: Index Test (Rapid molecular tests)	
DOMAIN 3: Reference Standard	
Is the reference standards likely to correctly classify the target condition?	No
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear
Reference standard does not incorporate result of index test?	Yes
Could the reference standard, its conduct, or its interpretation have introduced bias?	High risk
Are there concerns that the target condition as defined by the reference standard does not match the question?	High
DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Yes
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Unclear
Did all participants receive a reference standard?	Yes
Were results presented per patient?	Yes
Could the patient flow have introduced bias?	Unclear risk
enollar 2020(b)	
Study characteristics	
Patient Sampling	Two cohorts of patients presenting for COVID-19 testing at the same institution. This extraction relates to: [2] Single group study to estimate both sensitivity and specificity: asymptomatic contacts of confirmed cases (n=159) See Fenollar 2020(a) for extraction of additional cohort: [1] Single group study to estimate sensitivity alone: symptomatic patients, all

PCR positive (n=182)

Recruitment: Prospective



Genollar 2020(b) (Continued)	Prospective or retrospective: Unclear
Patient characteristics and setting	Setting: Unclear
-	Location: Institut Hospitalo-universitaire Méditerranée Infection, Marseille,
	Country: France
	Dates: Sep 21 to Oct 2 2020
	Symptoms and severity: All asymptomatic; 21/22 cases had Ct >25
	Demographics: Not reported
	Exposure history: [2] All described as contacts
Index tests	Test name: PANBIO COVID-19 Ag
	Manufacturer: Abbott
	Antibody: NP
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP
	Transport media: Not stated; appears to be direct testing
	Sample storage: Tested within 1 hour
	Test operator: Not stated
	Definition of test positivity: Visual line; as per manufacturer
	Blinding reported: Not stated, conducted first
	Timing of samples: Not stated
Target condition and reference standard(s)	Reference standard: Automated RT-PCR; VitaPCR (Credo diagnostics, Singapore
	Definition of non-COVID cases: As for cases; single negative
	Genetic target(s): Not stated
	Samples used: NP (paired, from opposite nostril)
	Timing of reference standard: Not stated
	Blinded to index test: Unclear
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Simultaneous; paired swabs
	All patients received same reference standard: Yes
	Missing data: None reported, no participant flow diagram reported
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported



Fenollar 2020(b) (Continued)	Unit of analysis: Patients	:		
Comparative				
Notes	Funding: Supported by the Méditerranée-Infection Foundation and the French Agence Nationale de la Recherche under reference Investissements d'Avenir Méditerranée Infection 10-IAHU-03 and Région Provence-Alpes-Côte d'Azur and European funding FEDER IHUBIOTK.			
	Source: Accepted manus	cript		
	Author COI: Pr Raoult and that develops diagnostic		ounders of the Pocrame startup liseases	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Low risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear	
DOMAIN 2: Index Test (Rapid molecular tests)				
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	No			



Fenollar 2020(b) (Continued)			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

FIND 2020a

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: - patients with symptoms consistent with COVID-19 (meeting national definitio for testing) presenting at a community testing clinic
	Recruitment: Consecutive recruitment
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community (COVID-19 testing clinic)
	Location: Institution not described; Marica, Rio de Janeiro
	Country: Brazil
	Dates: 30 Jul to 21 Aug 2020
	Symptoms and severity: All symptomatic; no further details
	Demographics: mean age 40y (range 4 to 84); reported for 396 participants 181 (45%) male
	Exposure history: Not stated
Index tests	Test name: NowCheck COVID-19 Ag test (RG1901DG)



FIND 2020a (Continued)	
	Manufacturer: Bionote Inc
	Antibody: SARS-CoV-2 nucleocapsid antigen
	Antigen target: Mouse monoclonal SARS-CoV-2 antibodies
	Test method: Rapid chromatographic immunoassay in lateral flow format
	Samples used: Proprietary NP swab collected by HCW
	Transport media: No transport media. Sample is immediately transferred to proprietary tube containing extraction buffer.
	Sample storage: Test should be performed as soon as possible after collection. Specimens may be stored at RT for 1h or 2-8°C for 4h.
	Test operator: HCW
	Definition of test positivity: Presence of visible control and test lines
	Blinding reported: Yes
	Timing of samples: median 4 days p.s.o (IQR 3, 6 days); day <0 to 3 152, 39% day 4 to 7 180, 46% day >=8 58, 15%
Target condition and reference standard(s)	Reference standard: RT-PCR (in-house assay based on the US CDC protocol); Ct threshold of 37
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection
	Genetic target(s): N1, N2
	Samples used: NP swabs
	Timing of reference standard: Same timing as per NP swabs for index test
	Blinded to index test: Yes
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: 0 to several days based on PCR turnaround times at the lab
	All patients received same reference standard: Yes
	Missing data: Reports 0 invalid results
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported
	Unit of analysis: Patients
Comparative	
Notes	Funding: FIND
	Publication status: published
	Source: FIND website/IFU index test



FIND 2020a (Continued)

Author COI: None stated (these are independent evaluations)

Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			,
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		,
Did the study avoid inappropriate inclusions?	Yes		,
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	



FIND 2020a (Continued)

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing		
Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Yes	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		Low risk

FIND 2020b

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity at single site: - patients seeking COVID-19 testing at main testing centre; described as presenting either with symptoms compatible with a SARS-CoV-2 infection, or with a known positive contact or asymptomatic HCWs (n=535)
	Recruitment: Consecutive recruitment
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community (main testing centre)
	Location: Hopitaux Universitaires de Geneve (HUG), Geneva
	Country: Switzerland
	Dates: 9-16 Oct 2020
	Symptoms and severity: 534/535 symptomatic (99%)
	Demographics: Mean age 38.5y (16 to 85y) 247, 46% male
	Exposure history: Not stated
Index tests	Test name: PanbioTM Covid-19 Ag Rapid Test (41FK10)
	Manufacturer: Abbott
	Antibody: Not reported
	Antigen target: Not reported
	Test method: CGIA (from product insert)
	Samples used: NP



FIND 2020b (Continued)				
	Transport media: No transport media; assay buffer used			
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU			
	Test operator: HCW			
	Definition of test positivity: Presence of visible control and test lines			
	Blinding reported: Yes			
	Timing of samples: time pso recorded for 115/124, 92%. Day 0-3 89, 78%; Day 4-7 23, 20%; Day 8+ 3, 3%			
Target condition and reference standard(s)	Reference standard: RT-PCR Roche Cobas; Ct threshold <40 (from Figure)			
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection			
	Genetic target(s): Not stated			
	Samples used: NP swab (paired, from contralateral nostril)			
	Timing of reference standard: Not stated; author contact advises only paired swabs used.			
	Blinded to index test: Yes			
	Incorporated index test: No			
Flow and timing	Time interval between index and reference tests: Paired swabs; 0 to several days based on PCR turnaround times at the lab			
	All patients received same reference standard: Yes			
	Missing data: Reports 0 invalid.			
	Uninterpretable results: None reported			
	Indeterminate results (index test): None reported			
	Indeterminate results (reference standard): None reported			
	Unit of analysis: Patients			
Comparative				
Notes	Funding: FIND			
	Publication status: published			
	Source: FIND/HUG website/IFU index test			
	Author COI: None stated (these are independent evaluations)			
Methodological quality				
Item	Authors' judgement Risk of bias Applicability concerns			
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			



FIND 2020b (Continued)			
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		



FIND 2020b (Continued)

Were results presented per patient? Yes

Could the patient flow have introduced bias?	Low risk

FIND 2020c (BR)

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity at three sites; this extraction is for data from Brazil (see FIND 2020c (CH) and Kruger 2020(c) for extraction of data from other sites): - ambulatory patients meeting national suspect definition for COVID-19 testing presenting at a community testing clinic in Brazil
	Recruitment: Consecutive recruitment
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community testing clinic
	Location: Macae, state of Rio de Janeiro
	Country: Brazil
	Dates: 13-30 Jul 2020
	Symptoms and severity: 392/397 (99%) symptomatic; no further details
	Demographics: mean age 37y (2-94) (397 participants); 229/398 male (57%)
	Exposure history: Not stated
Index tests	Test name: STANDARD Q COVID-19 Ag (09COV30D)
	Manufacturer: SD Biosensor Inc
	Antibody: Not reported
	Antigen target: Not reported
	Test method: Rapid chromatographic immunoassay in lateral flow format
	Samples used: NP; collected by HCW
	Transport media: Proprietary swab/media provided by SD Biosensor
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU
	Test operator: HCW
	Definition of test positivity: Presence of visible control and test lines
	Blinding reported: Yes
	Timing of samples: median 5 days p.s.o (IQR 4, 6 days) (for 397 patients); day <0 to 3 85, 21%; day 4 to 7 273, 69%; day >=8 39, 10%
Target condition and reference standard(s)	Reference standard: RT-PCR (In-house; Lab-developed assay based on the US CDC protocol; Ct threshold not stated; author contact advises Ct thresholds as per assay IFUs



FIND 2020c (BR) (Continued)					
	Definition of non-COVID absence of infection	cases: Same as for cases	. Single negative PCR required for		
	Genetic target(s): N1 and N2				
	Samples used: NP swabs	5			
	Timing of reference stan swabs used.	Timing of reference standard: Not stated; author contact advises only paired swabs used.			
	Blinded to index test: Yes	5			
	Incorporated index test:	No			
Flow and timing	Time interval between ir based on PCR turnaroun		Paired swabs; 0 to several days		
	All patients received sam	ne reference standard: Ye	28		
	Missing data: Reports 0 r	missing data			
	Uninterpretable results:	None reported			
	Indeterminate results (ir	ndex test): None reported	I		
	Indeterminate results (re	eference standard): None	e reported		
	Unit of analysis: Patients	3			
Comparative					
Notes	Funding: FIND				
	Publication status: published				
	Source: FIND website/IFU index test				
	Author COI: None stated	(these are independent	evaluations)		
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	Yes				
Was a case-control design avoided?	Yes				
Did the study avoid inappropriate exclusions?	Yes				
Did the study avoid inappropriate inclusions?	Yes				
Could the selection of patients have introduced bias?		Low risk			
Are there concerns that the included patients and setting do not match the review question?			Low concern		



FIND 2020c (BR) (Continued)			
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Low risk	



FIND 2020c (CH)

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity at single site; this extraction is for data from Switzerland (see FIND 2020c (BR) and Kruger 2020(c) for extraction of data from other sites): - patients seeking COVID-19 testing at main testing centre; described as presenting either with symptoms compatible with a SARS-CoV2 infection, or with a known positive contact or asymptomatic HCWs (n=529; from total cohort of 1064 volunteers)
	Recruitment: Consecutive recruitment
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community (main testing centre)
	Location: Hopitaux Universitaires de Geneve (HUG), Geneva
	Country: Switzerland
	Dates: 9-23 Oct 2020
	Symptoms and severity: Not stated; time pso recorded for $183/191$, 96% $141/183$ COVID positive cases had symptoms for 0-4days (77%)
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: STANDARD Q COVID-19 Ag (09COV30D)
	Manufacturer: SD Biosensor Inc
	Antibody: Not reported
	Antigen target: Not reported
	Test method: Rapid chromatographic immunoassay in lateral flow format
	Samples used: NP
	Transport media: Proprietary swab/media provided by SD Biosensor
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU
	Test operator: HCW
	Definition of test positivity: Presence of visible control and test lines
	Blinding reported: Yes
	Timing of samples: median not reported (range 0 to 15); day <0 to 3 - 122, 67%; day 4-7 - 54, 29%; Day 8+ - 7, 34%
Target condition and reference standard(s)	Reference standard: RT-PCR Roche Cobas; Ct threshold <40 (from Figure)
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection
	Genetic target(s): Not stated
	Samples used: NP swab (paired, from contralateral nostril)



FIND 2020c (CH) (Continued)	Timing of reference stand	dard: Not stated; author	contact advises only paired		
	swabs used.				
	Blinded to index test: Yes	Blinded to index test: Yes			
	Incorporated index test:	No			
Flow and timing	Time interval between index and reference tests: Paired swabs; 0 to several day based on PCR turnaround times at the lab				
	All patients received sam	e reference standard: Ye	es		
	Missing data: Reports 0 n	nissing data			
	Uninterpretable results:	None reported			
	Indeterminate results (in	dex test): None reported	I		
	Indeterminate results (re	ference standard): None	ereported		
	Unit of analysis: Patients				
Comparative					
Notes	Funding: FIND				
	Publication status: publis	Publication status: published			
	Source: FIND & HUG websites/IFU index test				
	Author COI: None stated	(these are independent	evaluations)		
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	Yes				
Was a case-control design avoided?	Yes				
Did the study avoid inappropriate exclusions?	Yes				
Did the study avoid inappropriate inclusions?	Yes				
Could the selection of patients have introduced bias?		Low risk			
Are there concerns that the included patients and setting do not match the review question?			Low concern		
DOMAIN 2: Index Test (Antigen tests)					
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes				
If a threshold was used, was it pre-specified?	Yes				



IND 2020c (CH) (Continued)			
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
		Low risk	
Could the patient flow have introduced bias?		LOW HISK	

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity at two sites; this extraction is for data from Brazil (see FIND 2020d (DE) for extraction of data from other site): - adults in community meeting national suspect definition for COVID-19 testing presenting at [1] a community testing clinic or [2] a tertiary level hospital
	Recruitment: Consecutive recruitment



FIND 2020d (BR) (Continued)	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Mixed; community testing clinic and tertiary hospital
	Location: [1] Macae, state of Rio de Janeiro, [2] Universidade Federal do Rio de Janeiro (UFRJ)
	Country: Brazil
	Dates: [1] 17 Aug to 9 Sept, [2] 11 Jul to 8 Aug
	Symptoms and severity: 421/450 (94%) symptomatic; no further details
	Demographics: mean age 39 y (0-95y) (451 participants); 185 male (41%)
	Exposure history: Not stated
Index tests	Test name: STANDARD F COVID-19 Ag FIA (F-NCOV-01G, 10COV30D)
	Manufacturer: SD Biosensor Inc
	Antibody: Not reported
	Antigen target: Not reported
	Test method: FIA
	Samples used: NP; collected by HCW
	Transport media: Proprietary swab/media provided by SD Biosensor
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU
	Test operator: HCW
	Definition of test positivity: As per STANDARD F Analyzer; cut-off index (COI) \geq 1.0 (a per IFU)
	Blinding reported: Yes
	Timing of samples: median 4 days p.s.o (IQR 3, 6 days) (for 421 patients). Day <0 to 3 - 131, 31%; day 4 to 7 - 248, 59%; day >=8 - 42, 10%
Target condition and reference standard(s)	Reference standard: RT-PCR; one of two in-house assays: 1. Lab-developed assay based on the US CDC protocol; 2. Lab-developed assay based on the Charité Universitätsmedizin Berlin protocol. Ct thresholds not stated; author contact advises Ct thresholds as per assay IFUs
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection
	Genetic target(s): 1. N1 and N2; 2. E and RdRp
	Samples used: NP swabs
	Timing of reference standard: Not stated; author contact advises only paired swabs used.
	Blinded to index test: Yes
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired swabs; 0 to several days based on PCR turnaround times at the lab
anid, point-of-care antigen and molecular-based	tests for diagnosis of SARS-CoV-2 infection (Review)



FIND 2020d (BR) (Continued)			
	All patients received sam	e reference standard: Yes	
	Missing data: Reports 0 m	nissing data	
	Uninterpretable results: I	None reported	
	Indeterminate results (in	dex test): None reported	
	Indeterminate results (re	ference standard): None r	eported
	Unit of analysis: Patients		
Comparative			
Notes	Funding: FIND		
	Publication status: publis	shed	
	Source: FIND website/IFU	for index test	
	Author COI: None stated	these are independent ev	valuations)
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)		



FIND	2020d	(BR)	(Continued)
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FIND 2020d (BR) (Continued)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes	 	
Could the patient flow have introduced		Low risk	

FIND 2020d (DE)

bias?

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity at two sites; this extraction is for data from Germany (see FIND 2020d (BR) for extraction of data from other site): - adults in community meeting national suspect definition for COVID-19 testing presenting at [1] a drive-in testing centre or [2] ambulatory testing clinic
	Recruitment: Consecutive recruitment
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community
	Location: [1] Heidelberg drive in testing, [2] Berlin: Ambulatory testing clinic of Charité – University Hospital



FIND 2020d (DE) (Continued)	
	Country: Germany
	Dates: [1] Heidelberg: 15 June-18July 2020, [2] Berlin: 6 July – 23 Sept 2020
	Symptoms and severity: 517/669 (77%) symptomatic; no further details
	Demographics: mean age 38 y (18-85y) (676 participants); 307 male (46%)
	Exposure history: Not stated
Index tests	Test name: STANDARD F COVID-19 Ag FIA (F-NCOV-01G, 10COV30D)
	Manufacturer: SD Biosensor Inc
	Antibody: Not reported
	Antigen target: Not reported
	Test method: FIA
	Samples used: [1] NP; [2] Combined NOP swabs; collected by HCW
	Transport media: Proprietary swab/media provided by SD Biosensor
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU
	Test operator: HCW
	Definition of test positivity: As per STANDARD F Analyzer; cut-off index (COI) \geq 1.0 (as per IFU)
	Blinding reported: Yes
	Timing of samples: median 3 days p.s.o (IQR 2,5 days) (for 505 patients). Day <0 to 3 - 257, 51%; day 4 to 7 - 202, 47%; day >=8 - 46, 9%
Target condition and reference standard(s)	Reference standard: RT-PCR; one of 5 assays: 1. Cobas SARS-CoV-2 (Roche Diagnostics Inc); N = 342 2. Abbott RealTime SARS-CoV-2 (Abbott Molecular, Inc) N = 1 3. Allplex 2019-nCov Assay (Seegene Inc); N = 20 4. LightMix® Modular SARS-CoV (COVID19) E-gene (Tib Molbiol); N = 233 5. Cobas (Roche) or Thermofisher (Multiplex TaqPath COVID-19 CE-IVD RT-PCR Kit); N = 80 Ct thresholds not stated; author contact advises Ct thresholds as per assay IFUs
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection
	Genetic target(s): Not stated apart from 3. E gene
	Samples used: NP (n=305), NOP (n=342) and/or OP swabs (n=32)
	Timing of reference standard: Not stated; author contact advises only paired swabs used.
	Blinded to index test: Yes
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired swabs; 0 to several days based on PCR turnaround times at the lab
	All patients received same reference standard: Yes



FIND 2020d (DE) (Continued)			
	Missing data: Reports 0 m	issing data	
	Uninterpretable results: N	lone reported	
	Indeterminate results (ind	dex test): None reported	
	Indeterminate results (re	erence standard): None i	reported
	Unit of analysis: Patients		
Comparative			
Notes	Funding: FIND		
	Publication status: publis	hed	
	Source: FIND website/IFU	for index test	
	Author COI: None stated (these are independent e	valuations)
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern



FIND 2020d (DE) (Continued)

DOMAIN 2: Index Test	(Rapid molecular tests)
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Donnate 2. maex rese (rapid motecular tests)				
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	No			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	Yes			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow have introduced bias?		Low risk		

FIND 2020e (BR)

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Patient Sampling	Single group study to estimate sensitivity and specificity; this extraction is for data from Brazil (see FIND 2020e (DE) for extraction of data from other site): - adults in community meeting national suspect definition for COVID-19 testing presenting at a community testing clinic (n=476)
	Recruitment: Consecutive recruitment
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community testing clinic
	Location: Marica, state of Rio de Janeiro



IND 2020e (BR) (Continued)	Country Provil
	Country: Brazil
	Dates: 27 Jul to 16 Sep
	Symptoms and severity: 470/476 (99%) symptomatic; no further details
	Demographics: mean age 45 y (0-106 y) (473 participants); 252 male (53%)
	Exposure history: Not stated
Index tests	Test name: BIOCREDIT COVID-19 Ag (G61RHA20)
	Manufacturer: RapiGEN Inc
	Antibody: Not reported
	Antigen target: Not reported
	Test method: LFA (CGIA, from IFU)
	Samples used: NP; collected by HCW
	Transport media: Assay diluent provided by manufacturer
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU
	Test operator: HCW
	Definition of test positivity: Visual appearance of test and control lines
	Blinding reported: Yes
	Timing of samples: median 5 days p.s.o (IQR 4, 7 days) (for 470 patients). Day <0 to 3 - 95, 20%; day 4 to 7 - 296, 63%; day >=8 - 79, 17%
Target condition and reference standard(s)	Reference standard: RT-PCR; Lab-developed assay based on the US CDC proto-
	col. Ct threshold not stated; author contact advises Ct thresholds as per assay IFUs
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection
	Genetic target(s): N1 and N2
	Samples used: NP swabs
	Timing of reference standard: Not stated; author contact advises only paired swabs used.
	Blinded to index test: Yes
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired swabs; 0 to several day based on PCR turnaround times at the lab
	All patients received same reference standard: Yes
	Missing data: Reports 0 missing data
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported



FIND 2020e (BR) (Continued)	Unit of analysis: Patients	:	
Comparative			
Notes	Funding: FIND		
	Publication status: publi	shed	
	Source: FIND website/IFU	J for index test	
	Author COI: None stated	(these are independent	evaluations)
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		



FIND 2020e	(BR)	(Continued)
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Reference standard does not incorporate result of index test?

Could the reference standard, its conduct, or
its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index
test and reference standard?

Yes

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis?

Yes

Did all participants receive a reference standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

Low risk

FIND 2020e (DE)

Study	characte	ristics
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Patient Samp	ling
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Single group study to estimate sensitivity and specificity at two sites; this extraction is for data from Germany (see FIND 2020e (BR) for extraction of data from other site):

- adults in community meeting national suspect definition for COVID-19 testing pre-

senting at

[1] a drive-in testing centre or

[2] ambulatory testing clinic

Recruitment: Consecutive recruitment

Prospective or retrospective: Prospective

Patient characteristics and setting

Setting: Community

Location: [1] Heidelberg drive in testing; [2] Berlin: Ambulatory testing clinic of Charité

- University Hospital

Country: Germany

Dates: [1] Heidelberg: 4 May - 3 Sept; [2] Berlin: 4 May - 18 Aug

Symptoms and severity: 733/1223 symptomatic; no further details

Demographics: mean age 39.5 y (17,59.2 y) (1239 participants); 607 male (50%)

Exposure history: Not stated

Index tests

Test name: BIOCREDIT COVID-19 Ag (G61RHA20)



FIND 2020e (DE) (Continued)	Manufactures DesiCEN Inc
	Manufacturer: RapiGEN Inc
	Antibody: Not reported
	Antigen target: Not reported
	Test method: LFA (CGIA, from IFU)
	Samples used: [1] NP; [2] NOP; collected by HCW
	Transport media: Assay diluent provided by manufacturer
	Sample storage: Author contact advises tested as soon as possible and within the time limit specified in the IFU
	Test operator: HCW
	Definition of test positivity: Visual appearance of test and control lines
	Blinding reported: Yes
	Timing of samples: median 3 days p.s.o (IQR 2,4days) (for 701 patients). Day <0 to 3 - 472, 67%; day 4 to 7 - 161, 23%; day >=8 - 68, 10%
Target condition and reference standard(s)	Reference standard: RT-PCR; one of 5 assays: 1. Cobas SARS-CoV-2 (Roche Diagnostics Inc); N = 344 2. Abbott RealTime SARS-CoV-2 (Abbott Molecular, Inc) N = 114 3. Allplex 2019-nCov Assay (Seegene Inc); N = 571 4. LightMix® Modular SARS-CoV (COVID19) E-gene (Tib Molbiol); N = 132 5. RealStar® SARS-CoV-2 RT-PCR Kit (Altona Diagnostics); N = 80 Ct thresholds not stated; author contact advises Ct thresholds as per assay IFUs
	Definition of non-COVID cases: Same as for cases. Single negative PCR required for absence of infection
	Genetic target(s): Not stated
	Samples used: NP swabs
	Timing of reference standard: Not stated; author contact advises only paired swabs used.
	Blinded to index test: Yes
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired swabs; 0 to several days based on PCR turnaround times at the lab
	All patients received same reference standard: Yes
	Missing data: Reports 0 missing data
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported
	Unit of analysis: Patients
Comparative	
Notes	Funding: FIND



FIND 2020e (DE) (Continued)

Publication status: published

Source: FIND website/IFU for index test

Author COI: None stated (these are independent evaluations)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included pa- tients and setting do not match the re- view question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tes	sts)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		



FIND 2020e (DE) (Continued)		
Reference standard does not incorporate result of index test?	Yes	
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk
Are there concerns that the target condition as defined by the reference standard does not match the question?		High
DOMAIN 4: Flow and Timing		
Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Yes	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		Low risk

Fourati 2020 [A]

Study characteristics	
Patient Sampling	Two group study to estimate sensitivity and specificity: (1) residual samples from subjects with positive SARS-CoV-2 PCR tested when they presented symptoms at the time of the first epidemic wave (n=297) (2) pre-pandemic samples (n=337)
	Recruitment: Random (stratified by Ct and time pso)
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Mixed; likely outpatient and in-patient "consulted or were admitted"
	Location: Henri Hospital Mondor de Créteil
	Country: France
	Dates: March 9 to April 9, 2020.
	Symptoms and severity: Not stated; all apparently symptomatic Data by viral load reported for 293/297 cases: <=20 Ct - 39, 13%; 20 to 25 Ct - 88, 30%; 25 to 30 Ct - 72, 25%; >30 Ct - 88, 30%
	Demographics: Not stated
	Exposure history: Not stated



Fourati 2020 [A] (Continued)

Index tests

Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] data relate to test [A], see additional entries for tests [B] to [E]

[A] SARS-CoV-2 COVID-19 Respi-Strip

- [B] Standard Q COVID-19 Ag
- [C] PanBio COVID-19 Antigen Rapid Test
- [D] Biosynex COVID-19 Ag BSS
- [E] COVID-VIRO Antigen Rapid Test
- [F] NG Test SARS-CoV-2 Ag (assay excluded from review due to Vortex requirement as stated in IFU)

(no product codes reported)

Manufacturer:

[A] Coris BioConcept, Gembloux, Belgium [B] SD BIOSENSOR, Inc., Korea

- [C] Abbott, Chicago, Illinois, USA
- [D] Biosynex, Strasbourg, France
- [E] AAZ, Boulogne-Billancourt, France
- [F] NG Biotech, Guipry, France

Antibody: Not stated

Antigen target: Not stated

Test method: Not stated

Samples used: NP; collection not reported

Transport media: VTM (Cepheid® or Deltalab®); 100 μL used for testing

Sample storage: frozen at -80 °C until use

Test operator: Laboratory staff

Definition of test positivity: Visual, as per manufacturer.

Blinding reported: Yes; each test was interpreted independently by two different laboratory technicians. A third reading was carried out in the event of discrepancy

Timing of samples: post-symptom onset (reported for 289 samples): 0-3 days 97, 34%; 4-7

days 103, 36%; 8=11 days 63, 22%; >=12 days 26, 9%

No. samples reported at >7 days varied per test, maximum was 289

Target condition and reference standard(s)

Reference standard: RT-PCR; in-house assay developed by CNR (Institut Paster) or RealStar

SARS-CoV-2 (Altona Diagnostics, Germany)

Definition of non-COVID cases: Pre-pandemic

Genetic target(s): Not stated

Samples used: NP; same as for index

Timing of reference standard: As for index

Blinded to index test: Yes, seems to be at time of sampling

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Same swab; simultaneous

All patients received same reference standard: Yes

Missing data: Number of cases missing per assay varied; reasons for missing data not reported (presumably invalid assay results)



Fourati 2020 [A] (Continued)	[A] 5, 1.7% [B] 6, 2.0% [C] 2, 0.7% [D] 0 [E] 2, 0.7% [F] 0 Uninterpretable results: No Indeterminate results (indeterminate results (reference))	x test): Not stated rence standard): Not stated	d
Comparative			
Notes	Funding: Evaluation of [A] a tières and Epicenter Publication status: Publishe Source: Laboratory report of Author COI: No COI present	ed obtained via SFM Microbiol	ooration with Médecins sans Fron- logie website
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it prespecified?	Yes		



Fourati 2020 [A] (Continued)				
Could the conduct or interpretation of the index test have introduced bias?		Low risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High	
DOMAIN 2: Index Test (Rapid molecula	r tests)			
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Yes			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	No			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow have introduced bias?		High risk		
Fourati 2020 [B]				
Study characteristics				



Fourati 2020 [B] (Continued)

Patient Sampling

Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS

Patient characteristics and setting

Index tests

Comparative study of six Ag tests (no product codes reported); Fourati 2020 [B] relates to test [B] in the list below; see Fourati 2020 [A] for full study characteristics and QUADAS entries

[A] SARS-CoV-2 COVID-19 Respi-Strip

[B] Standard Q COVID-19 Ag

- [C] PanBio COVID-19 Antigen Rapid Test
- [D] Biosynex COVID-19 Ag BSS
- [E] COVID-VIRO Antigen Rapid Test
- [F] NG Test SARS-CoV-2 Ag (assay excluded from review due to Vortex requirement as stated in IFU) (no product codes reported)

Manufacturer:

[A] Coris BioConcept, Gembloux, Belgium

[B] SD BIOSENSOR, Inc., Korea

- [C] Abbott, Chicago, Illinois, USA
- [D] Biosynex, Strasbourg, France
- [E] AAZ, Boulogne-Billancourt, France
- [F] NG Biotech, Guipry, France

Antibody: Not stated

Antigen target: Not stated

Test method: Not stated

Samples used: NP; collection not reported

Transport media: VTM (Cepheid® or Deltalab®); 100 μL used for testing

Sample storage: frozen at -80 °C until use

Test operator: Laboratory staff

Definition of test positivity: Visual, as per manufacturer.

Blinding reported: Yes; each test was interpreted independently by two different laboratory technicians. A third reading was carried out in the event of discrepancy

Timing of samples: post-symptom onset (reported for 289 samples): 0-3 days 97, 34%; 4-7 days 103, 36%; 8=11 days 63, 22%; >=12 days 26, 9%

No. samples reported at >7 days varied per test, maximum was 289

Target condition and reference standard(s)

Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS

Flow and timing

Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS

Comparative

Notes



Fourati 2020 [C]

Study characteristics	
Patient Sampling	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Patient characteris- tics and setting	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Index tests	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [C] relates to test [C] in the list below; see Fourati 2020 [A] for full study characteristics and QUADAS entries
	 [A] SARS-CoV-2 COVID-19 Respi-Strip [B] Standard Q COVID-19 Ag [C] PanBio COVID-19 Antigen Rapid Test [D] Biosynex COVID-19 Ag BSS [E] COVID-VIRO Antigen Rapid Test [F] NG Test SARS-CoV-2 Ag (assay excluded from review due to Vortex requirement as stated in IFU)
	Manufacturer:
	 [A] Coris BioConcept, Gembloux, Belgium [B] SD BIOSENSOR, Inc., Korea [C] Abbott, Chicago, Illinois, USA [D] Biosynex, Strasbourg, France [E] AAZ, Boulogne-Billancourt, France [F] NG Biotech, Guipry, France
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; collection not reported
	Transport media: VTM (Cepheid® or Deltalab®); 100 μL used for testing
	Sample storage: frozen at -80 °C until use
	Test operator: Laboratory staff
	Definition of test positivity: Visual, as per manufacturer.
	Blinding reported: Yes; each test was interpreted independently by two different laboratory technicians. A third reading was carried out in the event of discrepancy
	Timing of samples: post-symptom onset (reported for 289 samples): 0-3 days 97, 34%; 4-7 days 103, 36%; 8=11 days 63, 22%; >=12 days 26, 9% No. samples reported at >7 days varied per test, maximum was 289
Target condition and reference stan- dard(s)	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Flow and timing	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Comparative	
Notes	



Fourati 2020 [D]

Study characteristics	5
Patient Sampling	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Patient characteris- tics and setting	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Index tests	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [D] relates to test [D] in the list below; see Fourati 2020 [A] for full study characteristics and QUADAS entries
	 [A] SARS-CoV-2 COVID-19 Respi-Strip [B] Standard Q COVID-19 Ag [C] PanBio COVID-19 Antigen Rapid Test [D] Biosynex COVID-19 Ag BSS [E] COVID-VIRO Antigen Rapid Test [F] NG Test SARS-CoV-2 Ag (assay excluded from review due to Vortex requirement as stated in IFU)
	Manufacturer:
	 [A] Coris BioConcept, Gembloux, Belgium [B] SD BIOSENSOR, Inc., Korea [C] Abbott, Chicago, Illinois, USA [D] Biosynex, Strasbourg, France [E] AAZ, Boulogne-Billancourt, France [F] NG Biotech, Guipry, France
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; collection not reported
	Transport media: VTM (Cepheid® or Deltalab®); 100 μL used for testing
	Sample storage: frozen at -80 °C until use
	Test operator: Laboratory staff
	Definition of test positivity: Visual, as per manufacturer.
	Blinding reported: Yes; each test was interpreted independently by two different laboratory technicians. A third reading was carried out in the event of discrepancy
	Timing of samples: post-symptom onset (reported for 289 samples): 0-3 days 97, 34%; 4-7 days 103, 36%; 8=11 days 63, 22%; >=12 days 26, 9% No. samples reported at >7 days varied per test, maximum was 289
Target condition and reference stan- dard(s)	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Flow and timing	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Comparative	



Fourati 2020 [D] (Continued)

Notes

Fourati 2020 [E]

Study characteristics	s
Patient Sampling	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Patient characteris- tics and setting	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS
Index tests	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [E] relates to test [E] in the list be low; see Fourati 2020 [A] for full study characteristics and QUADAS entries
	 [A] SARS-CoV-2 COVID-19 Respi-Strip [B] Standard Q COVID-19 Ag [C] PanBio COVID-19 Antigen Rapid Test [D] Biosynex COVID-19 Ag BSS [E] COVID-VIRO Antigen Rapid Test [F] NG Test SARS-CoV-2 Ag (assay excluded from review due to Vortex requirement as stated in IFU)
	Manufacturer: [A] Coris BioConcept, Gembloux, Belgium [B] SD BIOSENSOR, Inc., Korea [C] Abbott, Chicago, Illinois, USA [D] Biosynex, Strasbourg, France [E] AAZ, Boulogne-Billancourt, France [F] NG Biotech, Guipry, France
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; collection not reported
	Transport media: VTM (Cepheid® or Deltalab®); 100 μL used for testing
	Sample storage: frozen at -80 °C until use
	Test operator: Laboratory staff
	Definition of test positivity: Visual, as per manufacturer.
	Blinding reported: Yes; each test was interpreted independently by two different laboratory technicians. A third reading was carried out in the event of discrepancy
	Timing of samples: post-symptom onset (reported for 289 samples): 0-3 days 97, 34%; 4-7 days 103, 36%; 8=1 days 63, 22%; >=12 days 26, 9% No. samples reported at >7 days varied per test, maximum was 289
Target condition and reference stan- dard(s)	Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteristics and QUADAS



Fourati 2020 [E] (Continued)

Flow and timing Comparative study of six Ag tests (no product codes reported); Fourati 2020 [A] reports full study characteris-

tics and QUADAS

Comparative

Notes

Ghofrani 2020

Study characteristics		
Patient Sampling	Single group study to estimate sensitivity and specificity in patients with both RT-PCR and POCT results available (n=113), including: [1] symptomatic patients with a PCR swab test close to presentation and a re-swal for POC testing, [2] patients with positive RT-PCR results and remnant NP swabs available for POC test, [3] asymptomatic patients with positive POC result on admission who were re-swabbed for RT-PCR confirmation. N per group was not reported	
	Recruitment: Convenience	
	Prospective or retrospective: Retrospective	
Patient characteristics and setting	Setting: Unclear; primarily in-patients?	
	Location: PeaceHealth Medical Group (10 hospitals and numerous clinics serving suburban and rural communities in three states)	
	Country: USA	
	Dates: April 6- April 21 2020	
	Symptoms and severity: Majority' symptomatic, no further details.	
	Demographics: Not stated	
	Exposure history: Not stated	
Index tests	Test name: ID NOW COVID-19 assay (no product code reported)	
	Manufacturer: Abbott Laboratories	
	Target gene(s): RdRp region	
	Antigen target: n/a	
	Test method: Isothermal PCR	
	Samples used: Nasal 58 (51.3%), NP 33 (29.2%), not stated 22 (19.5%). Direct testing 58 (51.3%), UTM 26 (23.0%); not stated 29 (25.7%).	
	Transport media: None or UTM; no further details	
	Sample storage: Not stated	
	Test operator: Not stated; infer laboratory staff.	
	Definition of test positivity: Not stated; presume as per manufacturer	



Ghofrani 2020 (Continued)	Blinding reported: Not stated		
	Timing of samples: Not stated; implies mostly close to presentation		
Target condition and reference standard(s)	Reference standard: RT-PCR; not described (conducted at one of two commercial laboratories, one of two State Public Health laboratories, an academic medical center, or tested in-house)		
	Definition of non-COVID cases: Same as index test; infer single negative		
	Genetic target(s): not stated		
	Samples used: Mixed; either paired swabs (within 3 days of each other) or same samples used		
	Timing of reference standard: Not stated		
	Blinded to index test: unclear; probably mixed depending on where RT-PCR was conducted		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Some same sample; paired samples could be up to 3 days apart		
	All patients received same reference standard: Yes		
	Missing data: None reported		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Patients		
Comparative			
Notes	Funding: No funding received		
	Publication status: Published		
	Source: Unclear		
	Author COI: none reported		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Unclear		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	No		



Ghofrani 2020 (Continued)			
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	No		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Were results presented per patient?	Yes		



Ghofrani 2020 (Continued)

Could the patient flow have introduced bias?

High risk

Gibani 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity with three sources of participants: [1] self-referred, health-care workers or their family members with suspected COVID-19 who were not admitted to hospital (n=280) [2] emergency department patients with suspected COVID-19 (n=15) [3] hospital inpatient admissions with or without suspected COVID-19 (n=91) Total N was 418 paired samples; 32 excluded as invalid (patient group not reported), 24 invalid on DnaNudge and 8 on RT-PCR) Recruitment: [1] and [2] Not reported; [3] consecutive Prospective or retrospective: Prospective
Patient characteristics and	Setting: Mixed ([1] community, [2] A&E, [3] Inpatient)
setting	Location: [1] St Mary's Hospital and the John Radcliffe Hospital, [2] St Mary's Hospital, [3] Chelsea & Westminster Hospital
	Country: London or Oxford, UK
	Dates: [1] April 10 to May 12, [2] April 2 to 24, [3] May 12 to 18
	Symptoms and severity: Only group [3] were inpatient
	Demographics: median age 46 y (IQR 31–66); 124, 32% male
	Exposure history: Not reported
Index tests	Test name: CovidNudge (no product code)
	Manufacturer: DnaNudge, UK
	Antibody: rdrp1, rdrp2, e-gene, n-gene, n1, n2, and n3
	Antigen target: n/a
	Test method: Automated RT-PCR; Described as "integrated lab-on-chip device that enables sample-to-result (RT-)PCR"
	Samples used: NP; HCW obtained swabs using pediatric swab
	Transport media: None
	Sample storage: No delay reported
	Test operator: Unclear; possibly HCW
	Definition of test positivity: at least two replicates of at least one viral gene target amplified
	Blinding reported: Yes; results from CovidNudge testing reported before laboratory results were available
	Timing of samples: On presentation; timing not reported



Gibani 2020 (Continued)

Target condition and reference standard(s)

Reference standard: SARS-CoV-2 RT-PCR; assay varied by site.

A. AusDiagnostics MT-PCR (Orf1ab, Orf8); n=74

b. Roche RT-PCR (Orf1ab, E); N=81 c. Abbott RT-PCR (RdRp, N); n=66

d. Thermo Fisher (orf1ab, the spike (S) gene and the nucleocapsid (N) gene); $n\!=\!21$

e. PHE in-house RT-PCR (RdRp); n=120

f. Imperial Molecular Diagnostics Unit (E); n=24

Definition of non-COVID cases: As above (single negative)

Genetic target(s): See above Samples used: NOP (paired)

Timing of reference standard: Not stated

Blinded to index test: Yes; centralised laboratory testing and point-of-care testing were done by separate staff members. Staff doing the centralised laboratory testing were masked to the point of-

care test results and vice-versa
Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Simultaneous (paired)

All patients received same reference standard: Yes (different assays)

Missing data: Additional 47 samples not 'paired'; not collected on same date

Uninterpretable results: 32 samples excluded; 24 invalid on DNANudge (failed to amplify RNaseP; 22/24 with associated RT-PCR result were negative) and 8 on RT-PCR (all 8 from one site)

Indeterminate results (index test): None reported

Indeterminate results (reference standard): None reported

Unit of analysis: Patients

Comparative

Notes

Funding: Supported by the National Institute for Health Research (NIHR) Imperial NHS Trust Biomedical Research Centre (London, UK). Part of this work was supported by the NIHR Health Protection Research Unit in Healthcare Associated Infections and Antimicrobial Resistance at Oxford University (Oxford, UK) in partnership with Public Health England (grant HPRU-2012-10041). DnaNudge supplied the test cartridges and NudgeBox processing units.

Publication status: Published

Source: Lancet Microbe

Author COI: CT, RS, MS, MK, T-KH, SDM, K-YFL, JB, and AO are employees of DnaNudge. CT is the co-inventor of the DnaNudge CovidNudge system and is named on the patent for the method and apparatus for analysing biological specimens on the DnaNudge platform (US Patent No: US 10 093 965.B2).16 LSPM has consulted for bioMerieux (2013–20), DNAelectronics (2015), Dairy Crest (2017–18), Pfizer (2018–20), and Umovis Lab (2020), received speaker fees from Profile Pharma (2018), received research grants from the UK National Institute for Health Research (NIHR; 2013–2019), Leo Pharma (2016), and CW+ Charity (2018–19), and received educational support from Eumedica (2016–17). NM has received speaker fees from Beyer (2016) and Pfizer (2019), and received educational support from Eumedica (2016) and Baxter (2017). MMG and GC are partly supported by the NIHR Imperial Biomedical Research Centre. GC is an NIHR research professor and investigator within the NIHR London in-vitro diagnostic co-operative. All other authors declare no competing interests.



Gibani 2020 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of pa- tients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen	tests)		
DOMAIN 2: Index Test (Rapid m	olecular tests)		
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or inter- pretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard	I		
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		



Gibani 2020 (Cor	tinued)
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Reference standard does not incorporate result of index test?

Yes

Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

Yes

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis?

No

Did all participants receive a reference standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

High risk

Goldenberger 2020

Study characteristics

Patient Sampling

Design unclear but appears to be a two group study to estimate sensitivity and specificity:

[1] SARS-CoV-2 positive samples selected to reflect a broad range of Ct values (n=10)

[2] SARS-CoV-2 negative samples (n=9)

Groups [1] and [2] from patients suspected of COVID-19 undergoing routine

diagnostics within a one week period

[third cohort of pre-pandemic samples positive for other coronaviruses re-

ported but not included in review (n=8)]

Recruitment: Convenience

Prospective or retrospective: Unclear

Patient characteristics and setting

Setting: Unclear

Location: University Hospital Basel



Goldenberger 2020 (Continued)		
	Country: Switzerland	
	Dates: One week during 2020 pandemic	
	Symptoms and severity: Not reported	
	Demographics: Not reported	
	Exposure history: Not reported	
Index tests	Test name: Xpert Xpress (no product code)	
	Manufacturer: Cepheid Inc	
	Antibody: E, N2	
	Antigen target: n/a	
	Test method: Automated RT-PCR	
	Samples used: NP	
	Transport media: UTM or eSwab media (Copan)	
	Sample storage: frozen at –80 $^{\circ}\text{C}$ until batch-wise sample processing with the Xpert	
	Test operator: laboratory technician	
	Definition of test positivity: Not stated; both targets reported in all samples	
	Blinding reported: Unclear	
	Timing of samples: Not stated	
Target condition and reference standard(s)	Reference standard: Roche cobas RT-PCR; threshold not reported but all positive samples <33 Ct	
	Definition of non-COVID cases: [2] COVID-19 suspects; as for cases (single negative PCR)	
	Genetic target(s): E, ORF1	
	Samples used: NP (same as index)	
	Timing of reference standard: Not stated	
	Blinded to index test: Yes, conducted first	
	Incorporated index test: Not stated	
Flow and timing	Time interval between index and reference tests: Simultaneous (same swab)	
	All patients received same reference standard: Yes	
	Missing data: None reported, no participant flow diagram reported	
	Uninterpretable results: None reported	
	Indeterminate results (index test): None reported	
	Indeterminate results (reference standard): None reported	



Goldenberger 2020 (Continued)

Notes
Funding: None reported
Publication status: Published
Source: Journal of Virological Methods
Author COI: None reported

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		



Go	lde	enl	berg	zer	2020	(Continued)
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Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an	appropriate interval between index
test and refe	ence standard?

Yes

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis?

Unclear

Did all participants receive a reference standard?

Yes

Were results presented per patient?

Unclear

Could the patient flow have introduced bias?

Unclear risk

Gremmels 2020(a)

Studv	charac	teristics

Patient Sampling

Report of two cohorts of patients presenting for COVID-19 testing. Gremmels 2020(a) en-

try relates to:

[1] community-dwelling mildly symptomatic subjects in a medium endemic area

(n=1369)

Gremmels 2020(b) entry reports data for second cohort in a high endemic area

Recruitment: Yes; all individuals invited to participate

Prospective or retrospective: Prospective

Patient characteristics and setting

Setting: Community testing centre

Location: [1] University Medical Center Utrecht (UMCU)

Country: Netherlands

Dates: [1] Sep 22 to Oct 6

Symptoms and severity: Cohort [1] only. Data on symptoms were missing from nine sub-

iects

Asymptomatic 37, 2.7%, Sore throat 907, 66.3%; Coryza 943, 69%; Cough 780, 57.1%; Headache 601, 44.0%; Tiredness 565, 41.3%; General malaise 365, 26.7% (further 19 doc-

umented)

Demographics: median age 36.4y (IQR 27.0, 49.6y); 523, 38.3% male

Exposure history: 233, 17% contact with confirmed case

Index tests

Test name: Panbio™ COVID-19 Ag Rapid Test (lot 41ADF011A)



Gremmels 2020(a) (Continued)	Manufacturer: Abbott (Lake Country, IL, U.S.A)
	Antibody: NP
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; obtained after NOP swab for RT-PCR; implies collected by HCW
	Transport media: Unclear; states transferred to 3 ml UTM after collection until further processing but also describes collected swabs transferred into dedicated sample collection tubes containing a sampling buffer for Ag test
	Sample storage: Not stated; within 2 hours of collection
	Test operator: Two independent observers
	Definition of test positivity: Visual line within 15 mins; as per manufacturer
	Blinding reported: Yes; observers (blinded to each other and to the PCR results)
	Timing of samples: Cohort [1] (data on duration of symptoms reportedly missing for 201 subjects; total reported here is 1138 but denominator for %s is 1166) day 1-3 pso 387, 33.2%; day 4-7 560, 48.0%; day >7 191, 16.4%
Target condition and reference standard(s)	Reference standard: RT-PCR; Seegene Allplex positive result on amplification of any of the three SARS-CoV-2 genes
	Definition of non-COVID cases: As for cases; single negative result
	Genetic target(s): E-, N-, and RdRP-gene
	Samples used: NOP (paired)
	Timing of reference standard: NOP swab obtained first for RT-PCR
	Blinded to index test: Not stated
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired
	All patients received same reference standard: Yes
	Missing data: 2 patients excluded ('inappropriate application of NP swab and lab mislabelling'), disease status not reported. [Considered overall low risk of bias due to small numbers]
	Uninterpretable results: None reported
	Indeterminate results (index test): None; no bands were classified as unclear by the independent observers
	Indeterminate results (reference standard): Patients
	Unit of analysis:
Comparative	
Notes	Funding: This study was investigator initiated. No external funding was received
	Publication status: Pre-print



Gremmels 2020(a) (Continued)

Author COI: No COI statement reported

Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular t	ests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		



Gremmels 2020(a) (Continued)			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Low risk	

Gremmels 2020(b)

Study characteristics	
Patient Sampling	Report of two cohorts of patients presenting for COVID-19 testing. Gremmels 2020(b) entry relates to: [2] community-dwelling mildly symptomatic subjects in a high endemic area (n=208)
	Gremmels 2020(a) entry reports data for second cohort in a medium endemic area
	Recruitment: Yes; all individuals invited to participate
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Community testing centre
	Location: [2] Horacio Oduber Hospital on Aruba
	Country: Netherlands
	Dates: [2] Sep 23 to Oct 9
	Symptoms and severity: Not stated; 'mildly symptomatic', presume mixed as per Gremmels 2020a
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: Panbio™ COVID-19 Ag Rapid Test (lot 41ADF011A)



iremmels 2020(b) (Continued)	Manufacturer: Abbott (Lake Country, IL, U.S.A)
	Antibody: NP
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; obtained after NOP swab for RT-PCR; implies collected by HCW
	Transport media: No UTM used for Ag samples; collected swabs transferred into dedicated sample collection tubes containing a sampling buffer
	Sample storage: Not stated; within 2 hours of collection
	Test operator: Two independent observers
	Definition of test positivity: Visual line within 15 mins; as per manufacturer
	Blinding reported: Yes; observers (blinded to each other and to the PCR results)
	Timing of samples: Not stated; on presentation
Target condition and reference standard(s)	Reference standard: RT-PCR; Seegene Allplex positive result = amplification of any of the three SARS-CoV-2 genes
	Definition of non-COVID cases: As for cases; single negative result
	Genetic target(s): E-, N-, and RdRP-gene
	Samples used: NOP (paired)
	Timing of reference standard: NOP swab obtained first for RT-PCR
	Blinded to index test: Not stated
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired
	All patients received same reference standard: Yes
	Missing data: None reported for Aruba site
	Uninterpretable results: None reported
	Indeterminate results (index test): None; no bands were classified as unclear by the independent observers
	Indeterminate results (reference standard): none
	Unit of analysis: patients
Comparative	
Notes	Funding: This study was investigator initiated. No external funding was received
	Publication status: Pre-print
	Source: medRxiv
	Author COI: No COI statement reported



Gremmels 2020(b) (Continued)

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			



Gremmels 2020(b) (Continued)		
Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Yes	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		Low risk

Gupta 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: - symptomatic patients with suspected COVID-19 and asymptomatic contacts of laboratory-confirmed cases between 5 and 10 days of exposure, meeting Indian Council of Medical Research (ICMR) strategy for COVID-19 testing
	Recruitment: Consecutive
	Prospective or retrospective: Not stated; appears prospective
Patient characteristics and setting	Setting: Outpatient (tertiary care hospital)
	Location: All India Institute of Medical Sciences (AIIMS), New Delhi
	Country: India
	Dates: May 31 to July 24, 2020.
	Symptoms and severity: 204 (62%) symptomatic; 126 (38%) asymptomatic. median symptom duration: 1 day (range: 1-10). Symptoms included: fever (31.5%), cough (25.4%), fatigue/malaise (11.8%), headache (3.3%), runny nose (3.3%)
	Demographics: median age 34.1±12.6 yr; 231 (70%) male
	Exposure history: 127 asymptomatic were in contact with confirmed case
Index tests	Test name: Standard Q rapid antigen detection test
	Manufacturer: SD Biosensor, Inc., Gurugram
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP; collection method detailed but personnel not described; presume HCW. Sequence for specimen collection was random for both the samples (Agand RT-PCR)
	Transport media: None
	Sample storage: None



Gupta 2020 (Continued)			
	Test operator: Same person who obtained swab; HCW		
	Definition of test positivity: Visual; test and control lines		
	Blinding reported: Yes; conducted first		
	Timing of samples: Symptomatic: 192 (95%) <=5 days pso (incl 57 cases)		
Target condition and reference standard(s)	Reference standard: RT-PCR; commercial assay (BGI Genomics Co. Ltd., China). Psoitive defined as per manufacturer IFU		
	Definition of non-COVID cases: As for cases; single negative		
	Genetic target(s): ORF1 ab		
	Samples used: nasal and throat swabs (NOP) in VTM		
	Timing of reference standard: As for index test; states the sequence for specimen collection was random for both the samples		
	Blinded to index test: Not stated		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Simultaneous; paired swabs		
	All patients received same reference standard: Yes		
	Missing data: None reported, no participant flow diagram reported		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Patients		
Comparative			
Notes	Funding: Study was financially supported by the Indian Council of Medical Research, New Delhi (for the Regional Virus Research and Diagnostic Laboratory at the All India Institute of Medical Sciences, New Delhi).		
	Publication status: Published		
	Source: Indian J Med Res		
	Author COI: Author report no COI present		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		



Gupta 2020 (Continued)			
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have intro- duced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		



Gupta 2020 (Continued)

Were results presented per patient? Yes

Could the patient flow have introduced	Unclear risk
bias?	

Harrington 2020

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity and specificity: - symptomatic patients meeting diagnostic criteria for COVID-19 (n = 524)
	Recruitment: consecutive
	Prospective or retrospective: unclear; presume prospective
	Number of samples (samples with confirmed SARS-CoV-2): 524 (186)
Patient characteristics and setting	Setting: ED (n = 3) or urgent (immediate) care centres (n = 2)
	Location: not stated; author institutions Loyola University Medical Centre, Cedars-Sinai Medical Centre
	Country: USA
	Dates: not reported
	Symptoms and severity: not stated
	Demographics: not stated
	Exposure history: not stated
Index tests	Test name: ID NOW COVID-19 assay (no product code provided)
	Manufacturer: Abbott
	Antigen target: not stated
	Antibody: N/A
	Test method: not stated; isothermal PCR
	Samples used: nasal swabs (provider collected)
	Transport media: none; direct testing after heat inactivation
	Sample storage: ED swabs transported in sterile transport containers (using cups or conical tubes)
	Test operator: on-site medical personnel (urgent care centres); laboratory personnel a each separate location (EDs) - 2 sites reportedly experienced users of ID NOW (one ED and one urgent care centre) and 3 sites received training)
	Definition of test positivity: as per manufacturer
	Blinding reported: yes (RT-PCR performed at separate central lab)
	Timing of samples: not stated; on presentation



Harrington 2020 (Continued)

Target condition and reference standard(s)	Reference standard: RT-PCR (Abbott RealTime SARS-CoV-2 (ACOV) assay performed on the Abbott m2000 system (Abbott Molecular Inc. Des Plaines, IL); threshold not stated		
	Definition of non-COVID c	ases: not specifically stated	d; presume yes as central lab used
	Genetic target(s): not stat	ed	
	Samples used: NP swabs		
	Timing of reference stand	ard: VTM (no detail)	
	Blinded to index test: not heat inactivated for 30 mi		ral clinical laboratory; samples
	Incorporated index test: n	o (paired collection with s	wabs for index test)
Flow and timing	Time interval between index and reference tests: simultaneous swab collection (different swabs for index and reference)		nultaneous swab collection (differ-
	All participants received s	ame reference standard: y	es
	Missing data: none report	ed, no participant flow dia	gram reported
	Uninterpretable results: none reported		
	Indeterminate results (inc	esults (index test): none reported	
	Indeterminate results (reference standard): 2 initial FPs had repeat sampling: - 1 retested on RT-PCR only and was positive (designated as TP) - 1 retested on RT-PCR and ID NOW and was negative on both (designated as FP based on original sampling)		
	Unit of analysis: participa	nts	
Comparative			
Notes	Funding: study authors received "received no specific grant from any funding agency in the public, commercial, or not-for-profit sectors"		
	Publication status: accepted manuscript		
	Source: Journal of Clinical Microbiology		
	Author COI: COI not mentioned		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			_
Was a consecutive or random sample of patients enrolled?	Yes		

Yes

Yes

Yes

Was a case-control design avoided?

sions?

sions?

Did the study avoid inappropriate exclu-

Did the study avoid inappropriate inclu-



Harrington 2020 (Continued)			
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tes	sts)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		



Harrington 2020 (Continued)

Were results presented per patient? Yes

Could the patient flow have introduced	Unclear risk
bias?	

Hogan 2020

Study characteristics	
Patient Sampling	Single-group design to estimate sensitivity and specificity - samples from adult patients from 1 hospital and paediatric and adult samples from surrounding hospitals
	Recruitment: unclear; equal numbers of positive and negative RT-PCR samples (suspect deliberate sampling by PCR result)
	Prospective or retrospective: not stated
	Number of samples (samples with confirmed SARS-CoV-2): 100 (50)
Patient characteristics and setting	Setting: hospital; not stated if inpatient or outpatient (samples selected from clinical virology laboratory)
	Location: Stanford Health Care (hospital), and surrounding hospitals (not named)
	Country: USA
	Dates: 7-13 April 2020
	Symptoms and severity: not stated
	Demographics: not stated
	Exposure history: not stated
Index tests	Test name: Accula SARS-CoV-2 POCT (no product code reported)
	Manufacturer: Mesa Biotech, Inc., San Diego, CA
	Antigen target: N gene
	Antibody: N/A
	Test method: rapid PCR
	Samples used: NP swabs in VTM (n = 37) or saline (n = 63, including 37 positive on RT-PCR)
	Transport media: not stated; 10 μL of VTM or saline was transferred to 60 μL of SARS-CoV-2 buffer within a biosafety cabinet (not covered by manufacturer IFU)
	Sample storage: not stated; testing appears to have been conducted soon after sample collection
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: as per manufacturer
	Blinding reported: not stated
	Timing of samples: not stated



Hogan 2020	(Continued)
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duced bias?

Target condition and reference standard(s)	Reference standard: RT-PCR; in-house SHC assay (cites Hogan 2020 10.1016/			
	j.jcv.2020.104383:104383			
	Definition of non-COVID cases: single RT-PCR negative			
	Genetic target(s): E gene			
	Samples used: NP swabs, same as for index test			
	Timing of reference standard: not stated			
	Blinded to index test: not	stated		
	Incorporated index test: ı	10 		
Flow and timing		dex and reference tests: no atory soon after sample co	t stated but implies that both llection	
	All participants received	same reference standard: y	es	
	Missing data: none repor	ted		
	Uninterpretable results: 3	3 invalid results were re-tes	ted; 1 positive and 2 negative	
	Indeterminate results (index test): 1 known RT-PCR-positive sample that show faint positive test line was re-tested and again showed the same faint test line sidered positive)			
	Indeterminate results (reference standard): none reported			
	Unit of analysis: refers to	participants		
Comparative				
Notes	Funding: study authors report no specific funding			
	Publication status: prepr	int		
	Source: medRxiv			
	Author COI: authors decla	are no COI present		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Unclear			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have intro-		Unclear risk		



Hogan 2020 (Continued)			
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	



Hou 2020

Study characteristics		
Patient Sampling	Single group study using remnant OP swabs submitted for SARS-CoV-2 testing at three medical centers (n = 285)	
	Recruitment: Not stated	
	Prospective or retrospective: Retrospective	
Patient characteristics and setting	Setting: Mixed inpatient and outpatient	
	Location: Three sites in Wuhan: Wuhan Tongji hospital (n=99), Wuhan Pulmonary hospital (n=96); Wuhan No. 1 hospital (n=90)	
	Country: China	
	Dates: Feb to Apr 2020	
	Symptoms and severity: 178 (62.5%) inpatient; 107 (37.5%) outpatients. Site 2 were all inpatients	
	Demographics: 220 (77.2%) aged ≤65 years; 159 (55.8%) male	
	Exposure history: No details; all Wuhan	
Index tests	Test name: Xpert Xpress (no product code reported)	
	Manufacturer: Cepheid Inc	
	Target gene(s): E, N2	
	Antigen target: N/A	
	Test method: Automated RT-PCR	
	Samples used: OP	
	Transport media: Not stated; 'aliquot made'	
	Sample storage: stored at -80°C within 24 h of collection	
	Test operator: Not stated	
	Definition of test positivity: Not stated; presume as per manufacturer (company funded study) - no mention of presumptive positive results	
	Blinding reported: Not stated	
	Timing of samples: Not stated	
Target condition and reference standard(s)	Reference standard: RT-PCR assays approved by Chinese National Medical Products Administration (NMPA) for the detection of SARS-CoV-2	
	Definition of non-COVID cases: As for cases; single negative RT-PCR	
	Genetic target(s): Not stated	
	Samples used: OP (same as for rapid test)	
	Timing of reference standard: Not stated; conducted at time of sample collection	
	Blinded to index test: Yes	



Hou 2020 (Continued)				
	Incorporated index test:	No		
Flow and timing	Time interval between index and reference tests: Simultaneous (same swab); time period of frozen storage was not reported All patients received same reference standard: Yes, although could be different RTP PCR assays at different sites			
	Missing data: None reported, no participant flow diagram reported Uninterpretable results: None reported			
	Indeterminate results (index test): None reported			
	Indeterminate results (re	ference standard): None	reported	
	Unit of analysis: Patients	; states 'samples from uni	que patients'	
Comparative				
Notes		3005-007) and by the Cepl	ect on Major Infectious Disease heid Investigator-Initiated Study	
	Publication status: Accepted manuscript			
	Source: J Clin Microbiol			
	Author COI: YWT is an employee of Cepheid, the comme Xpert Xpress SARS-CoV-2 test. The other authors declare			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?	Unclear risk			
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen tests)				
DOMAIN 2: Index Test (Rapid molecular tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			



Hou 2020 (Continued)			
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Jin 2020

Study characteristics	
Patient Sampling	Laboratory-based study presenting data on a total of 8043 specimens for different RT-PCR tests (n=7251) and ID NOW (n=792). States that a significant proportion of specimens tested by ID NOW were pre-admission screening specimens for surgical patients but does not report percentage. Eligible data refer to [1] single group study to estimate sensitivity and specificity in paired dry swabs and NP or OP swabs in UTM (n=52)



in 2020 (Continued)	
	[Additional cases only set: [2] 124 RT-PCR positive NP/OP samples in UTM samples in cluded 117 'retested with ID NOW' and 7 samples diluted in UTM from 4 positive specimens (the diluted samples cannot be distinguished from the set of 117 and data hav been excluded from review)
	Recruitment: Unclear
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Unclear; may be predominantly screening of surgical patients
	Location: Molecular & Genomic Pathology Laboratory, Thomas Jefferson University Hospital, Philadelphia
	Country: USA
	Dates: April 23 to 26, 2020
	Symptoms and severity: Not stated
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: ID NOW (product code not reported)
	Manufacturer: Abbott Laboratories
	Target gene(s): RdRp
	Antigen target: n/a
	Test method: Isothermal PCR
	Samples used: 'dry swabs' as per manufacturer EUA protocol
	Transport media: None
	Sample storage: No storage reported (appears to be immediate testing)
	Test operator: Not stated; laboratory staff presumed
	Definition of test positivity: As per manufacturer
	Blinding reported: Not stated 'tested in parallel'
	Timing of samples: Not stated
Target condition and reference standard(s)	Reference standard: RT-PCR; cobas SARS-CoV-2 Test (Roche Molecular Systems, Inc., Pleasanton, CA) using a cobas 6800 analyzer (Roche Molecular Systems, Inc). Either target present considered positive
	Definition of non-COVID cases: As above; single PCR negative required
	Genetic target(s): ORF1/a, E gene
	Samples used: Not specifically described for subset of paired samples, but for full co- hort NP and OP swabs in VTM used (400 uL)
	Timing of reference standard: Not stated
	Blinded to index test: Not stated; tested in parallel
	,



Jin 2020 (Continued)				
	All patients received same reference standard: Yes			
	Missing data: None reported			
	Uninterpretable results: None reported, no participant flow diagram reported			
	Indeterminate results (index test): None reported			
	Indeterminate results (re	ference standard): None r	eported	
	Unit of analysis: Not state	ed; described as 'paired pa	tient specimens'	
Comparative				
Notes	Funding: No funding state	ement reported		
	Publication status: Publis	hed		
	Source: Arch Path Lab Me	d		
	Author COI: No COI stater	nent reported		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Antigen tests)				
DOMAIN 2: Index Test (Rapid molecular test	s)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



Jin 2020	(Continued)
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Are there concerns that the index test, its conduct, or interpretation differ from the review question?

Unclear

Is the reference standards likely to correctly classify the target condition?

No

Were the reference standard results interpreted without knowledge of the results of the index tests?

Unclear

Reference standard does not incorporate result of index test?

Yes

Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

Yes

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis?

Unclear

Did all participants receive a reference standard?

Yes

Were results presented per patient?

Unclear

Could the patient flow have introduced bias?

Unclear risk

Jokela 2020

Study characteristics

Patient Sampling

Two group study to estimate sensitivity and specificity including NP or OP swab samples sent to university laboratory:

[1] for SARS-CoV-2 testing (n=97),

 $\label{eq:continuous} \ensuremath{\text{[2]}} \ pre-pandemic \ samples \ sent \ for \ testing \ due \ to \ suspicion \ of \ other \ respiratory$

virus infection (n=10) Recruitment: Not stated

Prospective or retrospective: Not stated; presume retrospective



okela 2020 (Continued)	[Also reports results for third cohort of samples from participants attending ter-		
	tiary care EDs (n=362), however index test is ineligible for this review (Novodiag))		
Patient characteristics and setting	Setting: Not reported		
	Location: Helsinki University Hospital Laboratory (HUSLAB), Helsinki		
	Country: Finland.		
	Dates: Mar to May 2020		
	Symptoms and severity: Not stated		
	Demographics: Not stated		
	Exposure history: Not stated		
Index tests	Test name: Xpert Xpress (no product code reported)		
	Manufacturer: Cepheid Inc		
	Target gene(s): E, N2		
	Antigen target: n/a		
	Test method: Automated RT-PCR		
	Samples used: NP or OP; no details on collection		
	Transport media: Not stated		
	Sample storage: Not stated		
	Test operator: Not stated		
	Definition of test positivity: Not stated; presume as per manufacturer - no mention of presumptive positive results		
	Blinding reported: Not stated		
	Timing of samples: Not stated		
Target condition and reference standard(s)	Reference standard: RT-PCR, one of three assays including 1) in-house LDT, 2) cobas SARS-CoV-2 test kit (Roche), or 3) Amplidiag COVID-19 test on the Amplidiag Easy platform (Mobidiag)		
	Definition of non-COVID cases: As above for COVID-19 suspects (single PCR negative); for pre-pandemic either Allplex Respiratory Panel 1/2/3 (Seegene, Seoul, Republic of Korea) and two by xTAG RVP Fast (Luminex Diagnostics, Toronto, Canada).		
	Genetic target(s): 1) N gene, 2) orf1ab and E, 3) orf1ab and N		
	Samples used: NP or OP, as for index		
	Timing of reference standard: Not stated		
	Blinded to index test: Not stated		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Simultaneous (same samples)		
	All patients received same reference standard: Yes (different assays)		



Jokela 2020 (Continued)			
	Uninterpretable results:	None reported	
	Indeterminate results (in	dex test): None reported	I
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Not repo	orted	
Comparative			
Notes	Funding: No funding stat	ement reported	
	Publication status: Preprint		
	Source: medRxiv		
	Author COI: No COI state	ment reported	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			



Jokela 2020 (Continued)			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		
Could the patient flow have introduced bias?		High risk	

Kruger 2020(a)	
Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity of three assays (each tested on a separate cohort of individuals, and extracted as three entries Kruger 2020(a), Kruger 2020(b), Kruger 2020(c). Participants at risk for SARS-CoV-2 infection based on exposure to a confirmed case, suggestive symptoms, or travel to a high risk area, presenting at one of three sites: (1) drive-in testing station (n=1213) (2) a clinical ambulatory testing facility (n=1308) (3) secondary care facility (n=53)
	This entry (Kruger 2020(a)) relates to the 727 participants tested with assay (a) from Shenzhen Bioeasy Biotechnology; it is unclear whether some particpants may have received more than one assay *This study was also reported as three independent FIND evaluations; author contact advised including data from the Kruger et al pre-print
	Recruitment: Not stated; recorded as consecutive, as per FIND evaluation protocol
	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: Mixed; (1), (2) Community (drive-in or clinical ambulatory testing); (3) secondary care



Kruger 2020(a) (Continued)

Location: Three sites: (1) Heidelberg, Germany; (2) Berlin, Germany and (3) Liverpool University Hospital Foundation Trust, Liverpool

Country: (1), (2) Germany, (3) UK

Dates: April 17th and August 25th, 2020; dates varied by assay and site

Whole sample:

Symptomatic on testing day (n=1901/2355, 80.7%)

N with prior negative test result (n=236/1928, 12.2%)

Mean age (SD) (n=2405: 40.4y (14.3))

Male (%) (n=1115/2361, 47.2%)

Participants undergoing assay (a) (denominator back-calculated from n and %)

Symptomatic on testing day: 564/694, 81.2%

N with prior negative test result: 73/624, 11.7%

Mean age (SD): 42.7y (14.9y)

Male (%): 47.2%

Index tests

Study reports data for three Ag assays, each tested on a separate cohort of individuals. This entry (Kruger 2020(a)) relates to assay [A]. See Kruger 2020(b) and Kruger 2020(c) for assays (b) and (c)

Test name: Bioeasy 2019-nCoV Ag Fluorescence Rapid Test Kit (Time-Resolved Fluorescence)

Manufacturer: Shenzhen Bioeasy Biotechnology Co. Ltd., Guangdong Province, China

Antibody: Not stated

Antigen target: Not stated

Test method: FIA

Samples used: Drive-in centre: NP or OP; Other centres: combined NOP (OP conducted first) RT-PCR swab obtained first, then same technique repeated for Ag test.

Transport media: None; used manufacturer supplied buffer solution as per IFU (for the Bioeasy assay, "the developer requested for pipettes to be used to transfer adequate quantities of liquid; in the IFU no pipette is needed and a nozzle is provided").

Sample storage: Drive-in centre and ambulatory testing: tested on site (presume short time frame) Secondary care: transported on ice to a category 3 facility for testing RT-PCR swab obtained first, then same technique repeated for Ag test.

Test operator: Drive-in and ambulatory clinic: POC evaluation

Secondary care: laboratory staff

Definition of test positivity: as per Analyzer

Invalid results were repeated once using the remaining buffer according to the respective IFUs. Readouts were done within the recommended time for each Ag-RDT (10 minutes for Bioeasy, 15 minutes for Coris and 15 to 30 minutes for SD Biosensor).

Blinding reported: Yes; "Staff performing the Ag-RDTs were blinded to results of RT-PCR tests and vice versa"

Timing of samples: Overall: mean 5 days pso (SD 9.6); for this assay 7.0 days (SD 12.2);

Target condition and reference standard(s)

Reference standard: RT-PCR; varied by site



Kruger 2020(a) (Continued)

Drive-in samples (Heidelberg): TibMolbiol (Berlin, Germany); the Allplex SARS-CoV-2 Assay from Seegene

(Seoul, South Korea); or the Abbott (Illinois, US) RealTime 2019-nCoV assay

Ambulatory testing (Berlin): Roche Cobas SARS CoV-2 assay (Pleasanton, CA United States) on the

Cobas® 6800 or 8800 system; SARS CoV-2 assay from TibMolbiol (Berlin, Germany)

Secondary care (UK): Genesig® Real-Time Coronavirus COVID-19 PCR assay (Genesig, UK)

 $Samples\ that\ showed\ a\ signal\ above\ the\ threshold\ in\ the\ relevant\ RT-PCR\ target\ regions\ for\ each\ assay$

were considered to be positive

Definition of non-COVID cases: As per cases; single negative result

Genetic target(s): Not stated

Samples used: Paired swabs; as per index test (RT-PCR swab obtained first,)

Drive-in centre: NP or OP

Other centres: combined NOP (OP conducted first)

Timing of reference standard: As per index test

Blinded to index test: Yes; "Staff performing the Ag-RDTs were blinded to results of RT-PCR tests and vice

versa'

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Paired; simultaneous

All patients received same reference standard: Yes (different assays)

Missing data: 154 excluded following enrolment [116 2nd swab refused, 3 nose bleed after 1st swab, 3 in-

sufficient time for both swabs, 31 other reasons, 1 no reason available]

Uninterpretable results: 2 invalid (PCR negative); PCR: 3 excluded as invalid (n=2) or not available (n=1)

Indeterminate results (index test): None reported;

Indeterminate results (reference standard): None reported

Unit of analysis: Patients

Comparative

Notes

Study reports an ease of use assessment; for this assay:

a high number of test execution steps (including precision pipetting) ... challenges when performing
multiple tests at the same time possibly hindering the test's wide-spread us

Funding: Study was supported by FIND, Heidelberg University Hospital and Charité – University Hospital internal funds. Pfizer funded the clinical team in Liverpool, UK.

Publication status: Pre-print

Source: medRxiv

Author COI: No COI statement reported; "external funders of the study had no role in study design, data collection, or data analysis"

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patie	nt Selection		



Kruger 2020(a) (Continued)					
Was a consecutive or random sample of patients enrolled?	Yes				
Was a case-control design avoided?	Yes				
Did the study avoid inappropriate exclusions?	Yes				
Did the study avoid inappropriate inclusions?	Yes				
Could the selection of patients have introduced bias?		Low risk			
Are there concerns that the included patients and setting do not match the review question?			Low concern		
DOMAIN 2: Index Test (An	tigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes				
If a threshold was used, was it pre-specified?	Yes				
Could the conduct or interpretation of the index test have introduced bias?		Low risk			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern		
DOMAIN 2: Index Test (Rapid molecular tests)					
DOMAIN 3: Reference Standard					
Is the reference stan- dards likely to correctly classify the target condi- tion?	No				
Were the reference stan- dard results interpreted without knowledge of	Yes				



Kruger 2020(a) (Continued) the results of the index tests?				
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timi	ng			
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference stan- dard?	Yes			
Were all patients included in the analysis?	No			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow have introduced bias?		High risk		

Kruger 2020(b)

Study characteristics

Patient Sampling

Single group study to estimate sensitivity and specificity of three assays (each tested on a separate cohort of individuals, and extracted as three entries Kruger 2020(a), Kruger 2020(b), Kruger 2020(c). Participants at risk for SARS-CoV-2 infection based on exposure to a confirmed case, suggestive symptoms, or travel to a high risk area, presenting at one of three sites:

- (1) drive-in testing station (n=1213)
- (2) a clinical ambulatory testing facility (n=1308)
- (3) secondary care facility (n=53)

This entry (Kruger 2020(c)) relates to the 425 participants tested with assay (b) from Coris Bioconcept; it is unclear whether some participants may have received more than one assay



Kruger 2020(b) (Continued)

*This study was also reported as three independent FIND evaluations; author contact advised including data from the Kruger et al pre-print

Recruitment: Not stated; recorded as consecutive, as per FIND evaluation protocol

Prospective or retrospective: Prospective

Patient characteristics and setting

Setting: Mixed; (1), (2) Community (drive-in or clinical ambulatory testing); (3) secondary care

Location: Three sites: (1) Heidelberg, Germany; (2) Berlin, Germany and (3) Liverpool University Hospital Foundation Trust, Liverpool

Country: (1), (2) Germany, (3) UK

Dates: April 17th and August 25th, 2020; dates varied by assay and site

Whole sample:

Symptomatic on testing day (n=1901/2355, 80.7%)

N with prior negative test result (n=236/1928, 12.2%)

Mean age (SD) (n=2405: 40.4y (14.3))

Male (%) (n=1115/2361, 47.2%)

Participants undergoing assay (b) (denominator back-calculated from n and %)

Symptomatic on testing day: 283/411, 68.9%

N with prior negative test result: 38/301, 12.6%

Mean age (SD): 44.9y (15.4y)

Male (%): 39.7%

Index tests

Study reports data for three Ag assays, each tested on a separate cohort of individuals. See Kruger 2020(a) and Kruger 2020(c) for details of the other assays

Test name: COVID-19 Ag Respi-Strip

Manufacturer: Coris Bioconcept, Gembloux, Belgium

Antibody: Not stated

Antigen target: Not stated

Test method: CGIA

Samples used: Drive-in centre: NP or OP

Other centres: combined NOP (OP conducted first)

RT-PCR swab obtained first, then same technique repeated for Ag test.

Transport media: None; used manufacturer supplied buffer solution as per IFU

Sample storage: Drive-in centre and ambulatory testing: tested on site (presume short time frame)

Secondary care: transported on ice to a category 3 facility for testing RT-PCR swab obtained first, then same technique repeated for Ag test.

Test operator: Drive-in and ambulatory clinic: POC evaluation

Secondary care: laboratory staff

Definition of test positivity: Visual appearance were interpreted by two operators, each blinded to the result of the other. In case of discrepant results, both operators re-read the result and agreed on a final result.

Invalid results were repeated once using the remaining buffer according to the respective IFUs. Readouts were done within the recommended time for each Ag-RDT (15 minutes for Coris).



Kruger 2020(b) (Continued)

Blinding reported: Yes; "Staff performing the Ag-RDTs were blinded to results of RT-PCR tests and vice

Timing of samples: Overall: mean 5 days pso (SD 9.6); this assay 6.2 days (SD 14.0)

Target condition and reference standard(s)

Reference standard: RT-PCR; varied by site

Drive-in samples (Heidelberg): TibMolbiol (Berlin, Germany); the Allplex SARS-CoV-2 Assay from See-

gene (Seoul, South Korea); or the Abbott (Illinois, US) RealTime 2019-nCoV assay

Ambulatory testing (Berlin): Roche Cobas SARS CoV-2 assay (Pleasanton, CA United States) on the

Cobas® 6800 or 8800 system; SARS CoV-2 assay from TibMolbiol (Berlin, Germany)

Secondary care (UK): Genesig® Real-Time Coronavirus COVID-19 PCR assay (Genesig, UK)

Samples that showed a signal above the threshold in the relevant RT-PCR target regions for each assay

were considered to be positive

Definition of non-COVID cases: As per cases; single negative result

Genetic target(s): Not stated

Samples used: Paired swabs; as per index test (RT-PCR swab obtained first,)

Drive-in centre: NP or OP

Other centres: combined NOP (OP conducted first)

Timing of reference standard: As per index test

Blinded to index test: Yes; "Staff performing the Ag-RDTs were blinded to results of RT-PCR tests and

vice versa"

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Paired; simultaneous

All patients received same reference standard: Yes (different assays)

Missing data: 154 excluded following enrolment [116 2nd swab refused, 3 nose bleed after 1st swab, 3

insufficient time for both swabs, 31 other reasons, 1 no reason available]

Uninterpretable results: 8 invalid (PCR negative)

PCR: 3 excluded as invalid (n=2) or not available (n=1)

Indeterminate results (index test): None reported;

Indeterminate results (reference standard): None reported

Unit of analysis: Patients

Comparative

Notes

Study reports an ease of use assessment; for this assay:

• challenges due to inconsistent test result interpretation (often only very faint lines visible) and defi-

ciencies in both the test kit quality and design

Funding: Study was supported by FIND, Heidelberg University Hospital and Charité – University Hospi-

tal internal funds. Pfizer funded the clinical team in Liverpool, UK.

Publication status: Pre-print

Source: medRxiv

Author COI: No COI statement reported; "external funders of the study had no role in study design, data

collection, or data analysis"



Kruger 2020(b) (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns				
DOMAIN 1: Patient Selection							
Was a consecutive or random sample of patients enrolled?	Yes						
Was a case-control design avoided?	Yes						
Did the study avoid inap- propriate exclusions?	Yes						
Did the study avoid inap- propriate inclusions?	Yes						
Could the selection of patients have introduced bias?		Low risk					
Are there concerns that the included patients and setting do not match the review question?			Low concern				
DOMAIN 2: Index Test (Anti	gen tests)						
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes						
If a threshold was used, was it pre-specified?	Yes						
Could the conduct or in- terpretation of the in- dex test have introduced bias?		Low risk					
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern				
DOMAIN 2: Index Test (Rap	id molecular tests)						
DOMAIN 3: Reference Stan	dard						
Is the reference standards likely to correctly classify the target condition?	No						



Kruger 2020(b) (Continued)				
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Yes			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timin	g			
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	No			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow have introduced bias?		High risk		

Kruger 2020(c)

Study characteristics

Patient Sampling

Single group study to estimate sensitivity and specificity of three assays (each tested on a separate cohort of individuals, and extracted as three entries Kruger 2020(a), Kruger 2020(b), Kruger 2020(c).

Participants at risk for SARS-CoV-2 infection based on exposure to a confirmed case, suggestive symptoms, or travel to a high risk area, presenting at one of three sites:

- (1) drive-in testing station (n=1213)
- (2) a clinical ambulatory testing facility (n=1308)
- (3) secondary care facility (n=53)

This entry (Kruger 2020(c)) relates to the 1263 participants tested with assay (c) from SD Biosensor; it is unclear whether some participants may have received more than one assay



Kruger 2020(c) (Continued)

*This study was also reported as three independent FIND evaluations; author contact advised including data from the Kruger et al pre-print

Recruitment: Not stated; recorded as consecutive, as per FIND evaluation protocol

Prospective or retrospective: Prospective

Patient characteristics and setting

Setting: Mixed; (1), (2) Community (drive-in or clinical ambulatory testing); (3) secondary care

Location: Three sites: (1) Heidelberg, Germany; (2) Berlin, Germany and (3) Liverpool University Hospital Foundation Trust, Liverpool

Country: (1), (2) Germany, (3) UK

Dates: April 17th and August 25th, 2020; dates varied by assay and site

Whole sample:

Symptomatic on testing day (n=1901/2355, 80.7%)

N with prior negative test result (n=236/1928, 12.2%)

Mean age (SD) (n=2405: 40.4y (14.3))

Male (%) (n=1115/2361, 47.2%)

Participants undergoing assay (b) (denominator back-calculated from n and %)

Symptomatic on testing day: 1054/1249, 84.4%

N with prior negative test result: 125/1000, 12.5%

Mean age (SD): 37.6 (12.7)

Male (%): 49.8%

Exposure history: Not stated

Index tests

Study reports data for three Ag assays, each tested on a separate cohort of individuals. See Kruger 2020(a) and Kruger 2020(b) for details of the other assays

Test name: STANDARD Q COVID-19 Ag Test

Manufacturer: SD Biosensor, Inc. Gyeonggi-do, Korea

Antibody: Not stated

Antigen target: Not stated

Test method: CGIA

Samples used: Drive-in centre: NP or OP

Other centres: combined NOP (OP conducted first)

RT-PCR swab obtained first, then same technique repeated for Ag test.

Transport media: None; used manufacturer supplied buffer solution as per IFU

Sample storage: Drive-in centre and ambulatory testing: tested on site (presume short time frame)

Secondary care: transported on ice to a category 3 facility for testing RT-PCR swab obtained first, then same technique repeated for Ag test.

Test operator: Drive-in and ambulatory clinic: POC evaluation

Secondary care: laboratory staff

Definition of test positivity: Visual appearance were interpreted by two operators, each blinded to the result of the other. In case of discrepant results, both operators re-read the result and agreed on a final result.



Kruger 2020(c) (Continued)

Invalid results were repeated once using the remaining buffer according to the respective IFUs. Readouts were done within the recommended time for each Ag-RDT (10 minutes for Bioeasy, 15 minutes for Coris and 15 to 30 minutes for SD Biosensor).

Blinding reported: Yes; "Staff performing the Ag-RDTs were blinded to results of RT-PCR tests and vice ver-

Timing of samples: Overall: mean 5 days pso (SD 9.6); this assay 3.7 days (SD 5.6)

Target condition and reference standard(s)

Reference standard: RT-PCR; varied by site

Drive-in samples (Heidelberg): TibMolbiol (Berlin, Germany); the Allplex SARS-CoV-2 Assay from Seegene (Seoul, South Korea); or the Abbott (Illinois, US) RealTime 2019-nCoV assay

Ambulatory testing (Berlin): Roche Cobas SARS CoV-2 assay (Pleasanton, CA United States) on the Cobas®

6800 or 8800 system; SARS CoV-2 assay from TibMolbiol (Berlin, Germany)

Secondary care (UK): Genesig® Real-Time Coronavirus COVID-19 PCR assay (Genesig, UK)

Samples that showed a signal above the threshold in the relevant RT-PCR target regions for each assay

were considered to be positive

Definition of non-COVID cases: As per cases; single negative result

Genetic target(s): Not stated

Samples used: Paired swabs; as per index test (RT-PCR swab obtained first,)

Drive-in centre: NP or OP

Other centres: combined NOP (OP conducted first)

Timing of reference standard: As per index test

Blinded to index test: Yes; "Staff performing the Ag-RDTs were blinded to results of RT-PCR tests and vice

versa"

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Paired; simultaneous

All patients received same reference standard: Yes (different assays)

Missing data: 154 excluded following enrolment [116 2nd swab refused, 3 nose bleed after 1st swab, 3 insufficient time for both swabs, 31 other reasons, 1 no reason available]

Uninterpretable results: 2 invalid (PCR negative); [B] 8 invalid (PCR negative); [C] 0 invalid reported PCR: 3 excluded as invalid (n=2) or not available (n=1)

Indeterminate results (index test): None reported;

Ease of use assessment reported:

[A] a high number of test execution steps (including precision pipetting) ... challenges when performing multiple tests at the same time possibly hindering the test's wide-spread use

[B] challenges due to inconsistent test result interpretation (often only very faint lines visible) and deficiencies in both the test kit quality and design

[C] no dissatisfactory scores identified

Indeterminate results (reference standard): None reported

Unit of analysis: Patients

Comparative

Notes

Study reports an ease of use assessment; for this assay:

· no dissatisfactory scores identified

Funding: Study was supported by FIND, Heidelberg University Hospital and Charité - University Hospital internal funds. Pfizer funded the clinical team in Liverpool, UK.



Kruger 2020(c) (Continued)

Publication status: Pre-print

Source: medRxiv

Author COI: No COI statement reported; "external funders of the study had no role in study design, data

collection, or data analysis"

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selec	ction		
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included pa- tients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (A	Intigen tests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpre- tation differ from the review question?			Low concern

DOMAIN 2: Index Test (Rapid molecular tests)



K	ruge	r 2020	C	(Continued)
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DOMAIN 3: Reference St	tandard			
Is the reference stan- dards likely to correctly classify the target con- dition?	No			
Were the reference standard results inter- preted without knowl- edge of the results of the index tests?	Yes			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Tin	ning			
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	Yes			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow		Low risk		

Lambert-Niclot 2020

Study characteristics



Lambert-N	iclot	2020	(Continued)
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Patient Sampling Single-group study to estimate sensitivity and specificity: samples submitted for RT-PCR testing (n = 138) Recruitment: not stated Prospective or retrospective: unclear; testing conducted prospectively Number of samples (samples with confirmed SARS-CoV-2): 138 (94) Patient characteristics and setting Setting: not stated Location: samples collected from virology laboratories of 3 university hospital groups from Assistance-Publique-Hôpitaux de Paris (APHP), (Saint-Antoine-Tenon-Trousseau, Saint-Louis-Lariboisière and Kremlin Bicêtre-Paul Brousse) Country: France Dates: 1-15 April 2020 Symptoms and severity: not stated Demographics: not stated Exposure history: not stated Test name: COVID-19 Ag Respi-Strip CORIS (no product code) Index tests Manufacturer: BioConcept, Gembloux, Belgium Antigen target: SARS-CoV-2 NP Antibody: monoclonal antibodies Test method: CGIA Samples used: NP swabs in VTM (collection process not described) Transport media: either of: COPAN UTM 3 mL, Virocult 1 mL, Eswab Amies 1 mL, 4MRT 3 mL, 0.9% NaCl buffer and cobas ROCHE Sample storage: no cooling or freezing step used Test operator: not stated; presume laboratory staff Definition of test positivity: not stated; as per manufacturer Blinding reported: not stated Timing of samples: not stated; presume on presentation Target condition and reference standard(s) Reference standard: RT-PCR (different kits used including RealStar Altona®, Anatolia®, cobas 6800 Roche®, Allplex™ 2019-nCoV Assay Seegene®) Definition of non-COVID cases: single negative PCR Genetic target(s): E gene Samples used: NP swabs (same as for index) Timing of reference standard: within a few hours after collection; time post onset of symptoms not reported Blinded to index test: unclear Incorporated index test: no



Lambert-Niclot 2020 (Continued)

Lamber Chietot 2020 (Continued)				
Flow and timing	Time interval between index and reference tests: same sample, both tests conducted within a few hours All participants received same reference standard: yes (different kits)			
	Missing data: none report	red		
	Uninterpretable results: 4 samples collected in cobas VTM gave invalid results and all samples in cobas medium were excluded			
	Indeterminate results (index test): control lines reported as "barely visible" for 9 positive and 8 negative tests			
	Indeterminate results (re	ference standard): none r	reported	
	Unit of analysis: not repo to be 1 per participant	rted, but samples tested	on day of collection so considered	
Comparative				
Notes	Funding: no funding sour	ces reported		
	Publication status: accep	ted manuscript		
	Source: Journal of Clinica	l Microbioloby		
	Author COI: no conflict of	interest statement repor	ted	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



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.ambert-Niclot 2020 (Continued)			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests	s)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	
Lephart 2020 [A]			
Study characteristics			
Patient Sampling		ncluding samples from: ing to emergency departm	ent (n=75), or

Recruitment: Not stated

Prospective or retrospective: Not reported



ephart 2020 [A] (Continued)			
	[Study also reports results for second group of recovering inpatients with previously laboratory-confirmed COVID-19 (n=13); for purposes of this review only those in group [1] were included]		
Patient characteristics and setting	Setting: [1] ED		
	Location: Not stated; pathology lab at University of Michigan Medical School		
	Country: USA		
	Dates: 22 Apr to 5 May 2020		
	Symptoms and severity: Not reported		
	Demographics: Not reported		
	Exposure history: Not reported		
Index tests	Test name: [A] ID NOW (second index test [B] Xpert Xpress, extracted as Lephart 2020 [B]; two additional RT-PCR tests evaluated in study but not included in this review). No product codes reported		
	Manufacturer: [A] Abbott Molecular		
	Target gene: Not reported in paper		
	Test method: [A] isothermal PCR		
	Samples used: [A] Nasal; Presume collected by HCP but not reported		
	Transport media: [A] None - transported dry swabs in sealed sterile collection bags		
	Sample storage: [A] within 24h		
	Test operator: Not stated; presume lab staff		
	Definition of test positivity: Each assay was performed according to manufacturer's EUA instructions.		
	Blinding reported: Not stated; unlikely		
	Timing of samples: On presentation; timing pso not reported		
Target condition and reference standard(s)	Reference standard: Composite: positive on >=2 of 4 NATs tested considered D+, including [A] ID NOW, [B] Xpert Xpress, [C] Simplexa COVID-19 Direct (Diasorin) (this wa the standard of care assay), [D] RealTime m2000 SARS-CoV-2 Assay (Abbott Molecular)		
	Definition of non-COVID cases: Three negatives (on different assays) required for D-		
	Genetic target(s): Not stated		
	Samples used: NP swabs (Same as for Xpert Xpress)		
	Timing of reference standard: Within 24h of sample collection (on presentation at ED); no further detail		
	Blinded to index test: Not stated; seems unlikely		
	Incorporated index test: Yes		
Flow and timing	Time interval between index and reference tests: Same swab [B], or paired collection [A]		
	All patients received same reference standard: Yes, all had all 4 assays		



Lephart 2020 [A] (Continued)				
	Missing data: None reported, no participant flow diagram reported			
	Uninterpretable results: N	lone reported		
			ults, [B] 1 'invalid' result; not re- only) on Xpert Xpress or no result	
	Indeterminate results (ref	erence standard): None re	eported	
	Unit of analysis: Unclear;	text refers to 'patients' so	presumed patient-based	
Comparative				
Notes	Funding: No funding state	ement reported		
	Publication status: Pre-pr	int		
	Source: bioRxiv			
	Author COI: No COI staten	nent provided		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Yes			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the included patients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Antigen tests)				
DOMAIN 2: Index Test (Rapid molecular test	s)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



ephart 2020 [A] (Continued)			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Reference standard does not incorporate result of index test?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	No		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	

Lephart 2020 [B]

Lepilart 2020 [B]	
Study characteristic	S
Patient Sampling	See Lephart 2020 [A] for full study details and QUADAS entries
Patient characteris- tics and setting	
Index tests	Test name: [B] Xpert Xpress (second index test [A] ID NOW, extracted as Lephart 2020 [A], also see see Lephart 2020 [A] for full study details and QUADAS entries; two additional RT-PCR tests evaluated in study but not included in this review). No product codes reported
	Manufacturer: [B] Cepheid



Lephart 2020 [B] (Continued)

Target gene: Not reported in paper

Test method: [B] Automated RT-PCR

Samples used: [B] NP; presume collected by HCP but not reported

Transport media: [B] M4-RT VTM (Thermo Fisher)

Sample storage: [B] stored at 4°C and tested within 24h

Test operator: Not stated; presume lab staff

Definition of test positivity: each assay was performed according to manufacturer's EUA instructions (pre-

sumptive positives not described)

Blinding reported: Not stated; unlikely

Timing of samples: On presentation; timing pso not reported

Target condition and reference standard(s)

See Lephart 2020 [A] for full study details and QUADAS entries

dard(s)

Flow and timing

See Lephart 2020 [A] for full study details and QUADAS entries

Comparative

Notes

Lieberman 2020

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity and specificity: - samples submitted for clinical diagnostic testing (n = 169; not all samples analysed for all tests)
	Recruitment: not stated
	Prospective or retrospective: retrospective (residual samples)
	Number of samples (samples with confirmed SARS-CoV-2): 169 (87)
Patient characteristics and setting	Setting: not stated; sampled from laboratory
	Location: Washington State Public Health Laboratory
	Country: USA
	Dates: not stated
	Symptoms and severity: not stated
	Demographics: not stated
	Exposure history: not stated
Index tests	Test name: Xpert Xpress
	Manufacturer: Cepheid



Lieberman 2020 (Continued)

Antigen target: E, N2

Antibody: N/A

Test method: rapid PCR

Samples used: NP swabs (collection not described)

Transport media: 300 µL of VTM sample

Sample storage: all same-sample comparisons were performed on specimens

stored at 4 °C for < 72 h with no freeze-thaws

Test operator: not stated; presume laboratory staff

Common panel of 26 specimens tested at UW by the UW CDC EUA-based LDT or at Lab-

Corp Seattle

Definition of test positivity: 1 of 2 targets detected was considered positive for all assays; Xpert Xpress data extracted as per IFU definition (positive = both targets or N gene posi-

tive; E-gene-positive requires retest)

Blinding reported: not stated

Timing of samples: not stated

Also evaluates:

[B] Hologic Panther Fusion RUO, [C] Hologic Panther Fusion EUA, [D] Diasorin Simplexa,

[E] Roche cobas 6800

in same 26 samples and in additional residual specimens (n = 115) at UW (different N per

test)

Target condition and reference standard(s)

Reference standard: RT-PCR; UW CDC EUA-based in-house test (positive if 1 of 2 targets

detected - presume at < 40 Ct)

Definition of non-COVID cases: single negative PCR

Genetic target(s): NI, N2

Samples used: NP swabs, as for index test

Timing of reference standard: not stated

Blinded to index test: not stated

Incorporated index test: no

Flow and timing

Time interval between index and reference tests: all testing conducted within 72 h

All participants received same reference standard: yes

Missing data: none reported, no participant flow diagram reported; review team excluded data for 28 specimens comparing Panther Fusion with DiaSorin Simplexa

Uninterpretable results: not stated

Indeterminate results (index test): 'Inconclusive' results (i.e. 1 genetic target detected) were considered positive due to the high specificity of all assays and limited cross-reactivity seen for SARS-CoV-2 primer sets. For Xpert Xpress only 12/13 were positive according to IFU specifications on first test (both targets present, or N gene positive); on retesting the presumptive positive became positive (detection of E-gene but not N-gene)

Indeterminate results (reference standard): as for index test

Unit of analysis: not stated, only refers to samples



ieberman 2020 (Continued) Comparative			
Notes	Funding: no funding state	ment reported	
	Publication status: accept	ed manuscript	
	Source: Journal of Clinical	Microbioloby	
	Author COI: no COI statem		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular t	ests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		



Lieberman 2020 (Continued)			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		
Could the patient flow have introduced bias?		Unclear risk	

Linares 2020

Study characteristics		
Patient Sampling	Single group study estimating sensitivity and specificity, recruiting at two locations: [1] symptomatic patients admitted to ED with clinical suspicion of COVID-19 (n=135) or asymptomatic patients with history of contact with another COVID-19 patient (n=17) [2] symptomatic patients (n=50) or asymptomatic (n=55) patients attending one of two primary healthcare centres	
	Recruitment: Not stated	
	Prospective or retrospective: Unclear; appears to be prospective	
Patient characteristics and setting	Setting: Mixed; A&E or primary care	
	Location: Hospital Universitario Príncipe de Asturias, Madrid	
	Country: Spain	
	Dates: Sep 10 to Sep 15	
	Symptoms and severity: 185, 72% symptomatic	



inares 2020 (Continued)		
	ED (n=135): fever 40, dyspnoea 42, cough 22, headache 14 Prim care (n=50): fever 14, dyspnoea 1, cough 18, headache 17	
	Demographics: Mean(?) age (range): ED 51.5y (37.0 to 71.8y); primary care 39.0y (25.0 to 56.0y) Male: ED 77 (51%), primary care 49 (47%)	
	Exposure history: Not stated	
Index tests	Test name: PanBio COVID-19 Ag Rapid Test Device (no product code)	
	Manufacturer: Abbott Rapid Diagnostic Jena GmbH, Jena, Germany	
	Antibody: Nucleocapsid	
	Antigen target: Not stated	
	Test method: Not stated; qualitative membrane-based immunoassay (immunochro matography)	
	Samples used: NP; HCW obtained	
	Transport media: None reported	
	Sample storage: Not stated	
	Test operator: Not stated	
	Definition of test positivity: Not stated; as per manufacturer	
	Blinding reported: Not stated	
	Timing of samples: ED: 2 days pso (IQR? 1-5) PC: 4 days pso (IQR? 2-8) Table 3 reports range of 0 to 27 days post symptom onset or post COVID-19 contact and range of 0 to 16 days for days post symptoms onset for symptomatic cases only	
Target condition and reference standard(s)	Reference standard: RT-PCR; Allplex SARS-CoV-2 assay (Seegene, Seoul, South Korea); appears to be <40 Ct threshold	
	Definition of non-COVID cases: As for cases (single -ve)	
	Genetic target(s): Not stated	
	Samples used: NP (paired)	
	Timing of reference standard: Not stated	
	Blinded to index test: Unclear	
	Incorporated index test: No	
Flow and timing	Time interval between index and reference tests: Paired	
	All patients received same reference standard: Yes	
	Missing data: None reported however 257 reported in Methods and 255 in Results, no participant flow diagram reported	
	Uninterpretable results: None reported	
	offiniter pretable results. None reported	
	Indeterminate results (index test): None reported	



Linares 2020 (Continued)	Unit of analysis: Patients		
Comparative			
Notes	Funding: No funding state	ement provided	
	Publication status: Pre-pi	rint	
	Source: medRxiv		
	Author COI: No COI stater	nent provided	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests))		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		



Linares 2020 (Continued)			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Liotti 2020

Study characteristics	
Patient Sampling	Unclear design estimating sensitivity and specificity; residual samples selected from one of two virology laboratories at two Covid-19 reference hospitals: [1] RT-PCR positive for SARS-CoV-2 (n=104) [2] RT-PCR negative for SARS-CoV-2 (n=255) Recruitment: Not stated
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Unclear; laboratory samples
	Location: From authors' institutions: Fondazione Policlinico Universitario A. Gemelli IRCCS, and Istituto Nazionale per le Malattie Infettive (INMI) Lazzaro Spallanzani IRCCS, Rome
	Country: Italy
	Dates: Not stated
	Symptoms and severity: Not stated;



iotti 2020 (Continued)		
	Of SARS-CoV-2 positive samples, 21, 20% high viral load (<25 Ct), 83, 80%low viral load (>=25) [28, 27% with Ct >=35]	
	Demographics: Not stated	
	Exposure history: Not stated	
Index tests	Test name: STANDARD F COVID-19 Ag FIA (no product codes reported)	
	Manufacturer: SD Biosensor (Suwon, South Korea)	
	Antibody: NP	
	Antigen target: monoclonal anti-SARS-CoV-2 antibody	
	Test method: FIA	
	Samples used: NP; collection not reported	
	Transport media: Not stated	
	Sample storage: performed within 24 hr after collection on samples kept at 4 C until testing	
	Test operator: Not stated; presume laboratory staff	
	Definition of test positivity: As per manufacturer	
	Blinding reported: Not stated	
	Timing of samples: Not reported	
Target condition and reference standard(s)	Reference standard: RT-PCR (one of 4 assays); Altona Diagnostics RealStar® SARS-CoV-2 RT-PCR, the Seegene Allplex™ 2019-nCoV, the DiaSorin Simplexa™COVID-19 Direct or the Roche Diagnostics Cobas® SARS-CoV-2 test	
	Definition of non-COVID cases: As for cases (single negative)	
	Genetic target(s): Not stated	
	Samples used: NP (same as index)	
	Timing of reference standard: Not stated	
	Blinded to index test: Yes (performed first)	
	Incorporated index test: No	
Flow and timing	Time interval between index and reference tests: Simultaneous (same swab)	
	All patients received same reference standard: Yes	
	Missing data: None reported, no participant flow diagram reported	
	Uninterpretable results: None reported	
	Indeterminate results (index test): None reported; FP results were re-tested with Ag assay, 3 of 4 remained positive (all blood contaminated)	
	Indeterminate results (reference standard): None reported	
	Unit of analysis: Not stated	



Liotti 2020 (Continued)

Notes

Funding: Study supported by funds to the Istituto Nazionale per le Malattie Infettive (INMI) Lazzaro Spallanzani IRCCS, Rome, Italy, from the Ministero della Salute (Ricerca Corrente, linea 1; COVID- 2020-12371817), the European Commission e Horizon 2020 (EU project 101003544 e CoNVat; EU project 101003551 e EXSCALATE4CoV; EU project 12371675 e EXCALATE4CoV; EU project 101005075 e KRONO) and the European Virus Archive e GLOBAL (grants no. 653316 and no. 871029).

Publication status: Published letter

Source: Clin Microbiol Infect

Author COI: All authors report no relevant conflicts of interest

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Unclear		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular t	ests)		
DOMAIN 3: Reference Standard			



Liotti 2020 (Continued)			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	

Loeffelholz 2020

Study characteristics	
Patient Sampling	Two-group study to estimate sensitivity and specificity for diagnosis of active disease - suspected patients referred for COVID-19 testing at 7 sites according to the local criteria (n = 486); sampled to enrich for RT-PCR-positive specimens (not further described)
	Recruitment: convenience (in addition, 1 site (LAC+USC) tested specimens from a 4-day point prevalence survey of patients presenting with COVID-19 symptoms)
	Prospective or retrospective: retrospective
	Number of samples (samples with confirmed SARS-CoV-2): 486 (220)
Patient characteris-	Setting: not stated
tics and setting	Location: 7 sites:
	Johns Hopkins University, Baltimore;
	LAC+USC Medical Centre, University of Southern California, Los Angeles;
	Manchester University NHS Foundation Trust Manchester;



Loeffelholz 2020 (Continued)

Mondor Hospital, Paris; New York City Dept. Health and Mental Hygiene, NYC; Niguarda Hospital, Milan;

University Hospital, Newark.

Country: USA, UK, France, Italy

Dates: 1 March-2 April 2020

Symptoms and severity: not stated

Demographics: adults at all sites except New York City Dept. Health and Mental Hygiene and Niguarda Hospital where all age groups were tested (ages not stated)

Exposure history: not stated

Index tests

Test name: Cepheid Xpert Xpress SARS-CoV-2 (RUO version, no product code reported)

Manufacturer: Cepheid Europe

Antigen target: nucleocapsid gene (N2) and the envelope gene (E) (RUO version also detects RdRp gene but this does not contribute to definition of positive)

Antibody: N/A

Test method: automated point-of-care PCR

Samples used: swabs (NP (n = 339), OP (n = 15), combined NP/OP in the same transport vial (n = 97)), and TA (n = 30):

- 1. Baltimore 61 NP
- 2. Los Angeles 88 NP
- 3. Manchester 54 NP/OP, 11 NP
- 4. Paris 68 NP
- 5. NYC NP 11, OP 15, TA 30, NP/OP 43
- 6. Milan 79 NP
- 7. Newark 21 NP

Transport media: VTM (swabs), diluted in saline (TA). 1 site (Manchester) pretreated specimens with an equal volume (≥ 30-< 50% (w/w)) of a guanidine hydrochloride buffer and heated at 80 °C

Sample storage: stored at -80 °C prior to index test, except at 1 site (University Hospital, Newark) where specimens were tested in real time, within 2 h by the Xpert test (n = 21).

Test operator: not stated; presume laboratory staff

Definition of test positivity: as per manufacturer: if both targets are detected, or if only N2 is detected, the test reports a positive result. If only the E target is detected the test reports a presumptive positive result "because this target is shared among some members of the sarbecovirus subgenus of coronaviruses". The RUO version of the test shows the amplification curves and PCR cycle threshold for all 3 genetic targets. The study reports that "The EUA test version cartridge contains the same reagents as the RUO cartridge. The only difference between the tests is the software which in the EUA version allows the user to see amplification curves and results for the N2 and E targets only".

Blinding reported: not stated

Timing of samples: not stated, presume on presentation

Target condition and reference standard(s)

Reference standard: RT-PCR (sites using each kit not reported, added by review team based on number of samples per site and per RT-PCR kit)

- 1. New York SARS-CoV-2 Real-time Reverse Transcriptase (RT)- PCR Diagnostic Panel; NYC
- 2. Quest SARS-CoV-2 rRT-PCR (Quest Diagnostics, San Juan Capistrano, US); Los Angeles



Loeffelholz 2020 (Continued)

- 3. RealStar® SARS-CoV-2 RT-PCR Kit 1.0 (Altona Diagnostics, Hamburg, Germany); Baltimore and Paris
- 4. GeneFinder COVID-19 Plus RealAmp Kit (ELITechGroup, Puteaux, France); Milan
- 5. Allplex 2019-nCoV Assay (Seegene, Seoul, SK); Milan
- 6. Charité Virology (Berlin, Germany) (in-house); Manchester
- 7. Abbott RealTime SARS-CoV-2 Assay (Abbott, Des Plaines, US); Newark
- 8. Simplexa COVID-19 Direct (DiaSorin, Cypress, US); Newark

Definition of non-COVID cases: yes (performed prior to index test)

Genetic target(s): different targets depending on RT-PCR test used:

- 1. New York Panel; N (N1, N2)
- 2. Quest; N (N1, N3)
- 3. RealStar; S, E
- 4. GeneFinderTM; RdRp, E, N
- 5. Allplex; RdRp, E, N
- 6. Charité Virology; RdRp
- 7. Abbott RealTime; RdRp, N
- 8. Simplexa; ORF1ab, S

Tie-breaker methods (for discrepant results), included: Hologic Panther Fusion (San Diego, USA), Tib-Molbiol LightMix Modular Wuhan Coronavirus E-gene RT-PCR (Roche, Basel, Switzerland); and the CDC assay (IDT primers and probes)

Samples used: as for index test

Timing of reference standard: as for index test

Blinded to index test: no storage; tested in real time

Incorporated index test: no

Flow and timing

Time interval between index and reference tests: same samples but index performed after frozen storage for undefined period of time except at University Hospital, Newark where specimens were tested in real time, within 2 h by the Xpert test

All participants received same reference standard: no

Missing data: 4 Xpert Xpress test results were lost permanently due to a single instrument computer malfunction

Uninterpretable results: 1 Xpert Xpress test was invalid due to a cartridge error (inadequate sample volume)

Indeterminate results (index test) presumptive positive results on Xpert Xpress were not reanalysed by Xpert Xpress, but all discrepant results were reanalysed by a third RT-PCR method

Indeterminate results (reference standard): specimens with inconclusive results by a test, and those with discrepant results between Xpert and the RT-PCR tests were analysed by a third RT-PCR method

1 FN result was inconclusive on Quest SARS-CoV-2, and negative on CDC RT-PCR; re-considered as TN

Of 11 FPs (including 1 presumptive positive on Xpert Xpress), 2 were negative on both New York SARS-CoV-2 and Panther Fusion (remained as FPs), and 9 were negative on in-house RT-PCR but positive on Roche RT-PCR (reclassified as TP)

In addition, 12 specimens (8 NP, 4 NP/OP) were inconclusive by the NY (RT)- PCR Diagnostic Panel and considered positive for data analysis purposes in the study. Of these, 11 were positive by the Xpert test and 1 was presumptive positive (EUA version of Xpert test). In 4 of these only the N1 target was detected and in 8 only the N2 target was detected by the New York EUA method, all with Ct values > 36

One NP specimen was inconclusive by the Quest SARS-CoV-2 rRT-PCR test and negative by the Xpert test. The Quest test reports inconclusive if only a single target (N1 or N3) is detected. They were unable to determine which target was detected by the Quest test. This specimen was negative by a tie-breaker NAAT.

Unit of analysis: not stated; only samples reported



Loeffelholz 2020 (Continued)

Comparative

Notes

Funding: not stated; presume funded by test manufacturer (see COI statement)

Publication status: accepted manuscript

Source: Journal of Clinical Microbiolobyogy

Author COI: the study was designed and supervised by the sponsor, Cepheid. Data were collected by investigators at each study site, and statistical analyses were performed by a Cepheid author. Cepheid authors wrote the first draft of the manuscript. All study authors vouch for the accuracy and completeness of the data reported.

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	No				
Was a case-control design avoided?	No				
Did the study avoid inappropriate exclusions?	Unclear				
Did the study avoid inappropriate inclusions?	Yes				
Could the selection of patients have introduced bias?		High risk			
Are there concerns that the included patients and setting do not match the re- view question?			High		
DOMAIN 2: Index Test	(Antigen tests)				
DOMAIN 2: Index Test (Rapid molecular tests)					
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear				
If a threshold was used, was it pre- specified?	Yes				



Loeffelholz 2020 (Continu	ued)		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or in- terpretation dif- fer from the review question?			Unclear
DOMAIN 3: Reference	Standard		
Is the reference stan- dards likely to cor- rectly classify the tar- get condition?	No		
Were the reference standard results in- terpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its con- duct, or its interpre- tation have intro- duced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and 1	iming		
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		



Loeffelholz 2020	(Continued)
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Did all participants receive a reference standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

High risk

Mak 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity alone: [1] RT-PCR positive samples selected from Hong Kong's COVID-19 reference laboratory (n=160 samples from 152 patients)
	Recruitment: Convenience; deliberate sampling of specific numbers of different respiratory sample types (selected from cohort of all available positive samples with sufficient quantity)
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Not stated
	Location: Public Health Laboratory Services Branch, Hong Kong
	Country: Hong Kong
	Dates: Feb 1 to Apr 21 2020
	Symptoms and severity: Not stated; High viral load (<18.57 Ct) - 64, 40% 'Normal' viral load >18.57 - 96, 60%
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: BIOCREDIT COVID-19 Ag (no product code reported)
	Manufacturer: RapiGEN Inc
	Antibody: Not stated
	Antigen target: Not stated
	Test method: CGIA
	Samples used: throat saliva (TS, n = 45), nasopharyngeal swab and throat swab (NPS & TS, n=103), nasopharyngeal aspirate and throat swab (NPA & TS, n=81), sputum (n=45); no details of collection methods
	Transport media: Samples were placed in viral transport media (VTM) or Phosphate-Buffered Saline (PBS). 100 μL sample volume was used; less viscous samples were added directly to sample well of the device, for more viscous samples the swaprovided with the kit was used to collect the samples and was immersed in the pro-



Mak 2020 (Continued)			
	vided assay diluent tube. The subsequent procedures were carried out according to the manufacturer's instructions.		
	Sample storage: stored at −70 °C until used for study purposes		
	Test operator: Not stated; laboratory staff presumed		
	Definition of test positivity: Not stated		
	Blinding reported: Not stated but all positive samples		
	Timing of samples: Not stated		
Target condition and reference standard(s)	Reference standard: In-house RT-PCR; <=40Ct		
	Definition of non-COVID cases: n/a		
	Genetic target(s): RdRp		
	Samples used: NPA & TS, NPS & TS, sputum and throat saliva, as for index test		
	Timing of reference standard: Not stated		
	Blinded to index test: Yes, prior to index test		
	Incorporated index test: Not stated		
Flow and timing	Time interval between index and reference tests: Simultaneous; same samples		
	All patients received same reference standard: Yes		
	Missing data: None reported, no participant flow diagram reported		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Samples (160 from 152 patients)		
Comparative			
Notes	Funding: No funding statement reported		
	Publication status: Published		
	Source: J Clin Virol		
	Author COI: Authors report no COI		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		



Mak 2020 (Continued)			
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have intro- duced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?		High	
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?		High	
DOMAIN 2: Index Test (Rapid molecular tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?		High	
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		



Mak 2020 (Continued)

Were results presented per patient? No

Could the patient flow	have introduced
bias?	

High risk

Mertens 2020

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity and specificity for diagnosis of active disease: - samples from patients suspected of SARS-COV-2 infections (n = 328)
	Recruitment: random sampling of samples submitted to 3 laboratories 322/328 NP samples (NP swabs) were randomly selected
	Prospective or retrospective: retrospectively
	Number of samples (samples with confirmed SARS-CoV-2): 328 (132)
Patient characteristics and set- ting	Setting: unclear; samples from university laboratories (discussion states that no outpatient population has been sampled, therefore assume inpatients and HCW samples)
	Location: laboratories at Université Libre de Bruxelles (LHUB-ULB), UZ Leuven and Centre Hospitalier Universitaire Sart-Tilman (CHU) Liège
	Country: Belgium
	Dates: 19-30 March 2020
	Symptoms and severity: not reported
	Demographics: not reported
	Exposure history: unclear; 53/328 samples were from HCW
Index tests	Test name: COVID-19 Ag Respi-Strip
	Manufacturer: Coris BioConcept (Belgium)
	Antigen target: SARS-CoV and SARS-CoV-2 highly conserved nucleoprotein
	Antibody: monoclonal antibodies directed against SARS-CoV and SARS-CoV-2 highly conserved nucleoprotein antigen
	Test method: immunochromatographic assay using colloidal gold (CGIA)
	Samples used: remnant respiratory specimens (322 NP swabs, 4 NP aspirate and 2 BAL)
	Transport media: NP: flocked swab + UTM 3 mL (or 1 mL of Amies) (Copan, Brescia, Italy); NPA: 3 mL VTM (veal infusion broth (Difco, Becton Dickinson, Sparks, MD, USA) supplemented with bovine albumin (Sigma Aldrich, St Louis, MO, USA)) BAL: N/A
	Sample storage: not described
	Test operator: laboratory technician
	Definition of test positivity: visible reddish-purple band appearing at the Test line position (T)
	Blinding reported: not stated



Mertens 2	2020	(Continued)
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Timing of samples: not clear

Target condition and reference standard(s)

Reference standard: qRT-PCR: RealStar SARS-CoV-2 RT-PCR Kit from Altona-diagnostics with a cut-off set at 40 Ct (LHUB-ULB); Roche LC480 thermocycler using Taqman Fast Virus 1-Step Master Mix (Thermo Fisher) (Liege); QuantStudio Dx (Thermo Fisher Scientific) or Panther Fusion (PF, Hologic, San Diego, USA) (UZ Leuven)

Definition of non-COVID cases:

- Genetic target(s): RealStar: not stated;
- · Taqman Fast Virus: RdRp and E genes
- QuantStudio Dx; "slightly adapted" E-gene
- · Panther Fusion: E gene and ORF1-ab

Samples used: as for index test (respiratory specimens (322 NP swabs, 4 NP aspirate and 2 BAL)

Timing of reference standard: not stated; same samples as for index test but analysed at time of collection

Blinded to index test: yes (undertaken for diagnostic purposes at time of collection)

Incorporated index test: no

Flow and timing

Time interval between index and reference tests: same samples used; discussion report 'some delay' between PCR and antigen testing

All participants received same reference standard: yes but different RT-PCR kits

Missing data: none reported, no participant flow diagram reported

Uninterpretable results: none reported; discussion reports some difficulties in visualising the strip through the closed tube requiring the lab technician to open the test tube in the laminar air flow cabinet and pull out the strip with forceps

Indeterminate results (index test): weak T lines considered positive

Indeterminate results (reference standard): none reported

Unit of analysis: refers to participants

Comparative

Notes

Funding: not stated

Publication status: preprint (not peer-reviewed)

Sourcepreprint server (medRxiv)

Author COI: the IVD medical device has been developed by the investigator Pascal Mertens, Henri Magein, and Justine Bouzet working for Coris BioConcept (potential conflict of interest declared even though they don't have any share in this company); Thierry Leclipteux was involved in the development of this test and is the CEO of Coris

Bioconcept (potential conflict of interest declared). All scientific investigators that are external to Coris BioConcept declare having no conflict of interest.

Methodological quality

Item Authors' judgement Risk of bias Applicability concerns

DOMAIN 1: Patient Selection



Mertens 2020 (Continued)			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tes	sts)		
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpre- tation of the index test have in- troduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid mole	ecular tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	



Mertens 2020 (Continued)		
Are there concerns that the target condition as defined by the reference standard does not match the question?		High
DOMAIN 4: Flow and Timing		
Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Unclear	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	

Unclear risk

Mitchell 2020

troduced bias?

Could the patient flow have in-

Study characteristics		
Patient Sampling	Single-group study to estimate sensitivity and specificity for diagnosis of active disease: - samples positive and negative on 1 of 2 SARS-CoV-2 RT-PCR assays	
	Recruitment: not stated; suggests possible deliberate sampling of positive case	
	Prospective or retrospective: retrospective (residual samples)	
	Number of samples (samples with confirmed SARS-CoV-2): 61 (46)	
Patient characteristics and setting	Setting: not stated; 2 independent laboratories (Class II biosafety cabinet (BSC)	
	Location: not stated; author institutions University of Pittsburgh School of Med icine, Pittsburgh and Laboratory of Viral Diseases, Wadsworth Centre, New York State Department of Health, Albany, NY	
	Country: USA	
	Dates: not stated	
	Symptoms and severity: not stated	
	Demographics: not stated	
	Exposure history: not stated	
Index tests	Test name: ID NOW COVID-19 (product code not reported)	
	Manufacturer: Abbott, Chicago, USA	



Mitchell 2020 (Continued)	Antigen target: not stated
	Antibody: N/A
	Test method: not stated (should be isothermal PCR)
	Samples used: NP samples (residual samples)
	Transport media: VTM; no further detail (no longer covered on IFU)
	Sample storage: stored at −80 ℃ prior to testing
	Test operator: certified laboratory personnel
	Definition of test positivity: not stated; as per manufacturer
	Blinding reported: not stated
	Timing of samples: not stated
Target condition and reference standard(s)	Reference standard: CDC EUA or the New York EUA RT-PCR assays
	Definition of non-COVID cases: single RT-PCR negative
	Genetic target(s): not stated
	Samples used: as for index test
	Timing of reference standard: as for index test
	Blinded to index test: not stated; samples analysed at or near time of collection
	Incorporated index test: no
Flow and timing	Time interval between index and reference tests: same samples but used at different times (samples used for index test stored at $-80~\%$)
	All participants received same reference standard: no, either the CDC EUA or the New York EUA assays
	Missing data: none reported, no participant flow diagram reported
	Uninterpretable results: none reported
	Indeterminate results (index test): none reported
	Indeterminate results (reference standard): none reported
	Unit of analysis: not stated; only samples reported
Comparative	
Notes	Funding: not stated
	Publication status: accepted manuscript
	Source: Journal of Clinical Virology
	Author COI: COI not mentioned by study authors
Methodological quality	
Item	Authors' judgement Risk of bias Applicability concerns



Mitchell 2020 (Continued)

DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			

DOMAIN 3. Reference Standard	
Is the reference standards likely to correctly classify the target condition?	No
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes
Reference standard does not incorporate result of index test?	Yes

its interpretation have introduced bias?	
Are there concerns that the target condition as defined by the reference standard does not match the question?	High
DOMAIN 4: Flow and Timing	

High risk

Could the reference standard, its conduct, or



Mitchell 2020 (Continued)		
Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Unclear	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Unclear	
Could the patient flow have introduced bias?		Unclear risk

Moore 2020

Study characteristics	
Patient Sampling	2-group study to estimate sensitivity and specificity: - samples from symptomatic (fever or cough or shortness of breath) adult and paediatric outpatients, ED patients, and inpatients
	Recruitment: consecutive (first 94 participants), then all PCR-positive samples plus the next PCR-negative sample after each positive sample, to a total of 200 samples
	Prospective or retrospective: retrospective (participant and sample details extracted from the electronic medical record)
	Number of samples (samples with confirmed SARS-CoV-2): 200 (125)
Patient characteristics and setting	Setting: mixed (outpatients, ED patients and inpatients)
	Location: Rush University Medical Centre (RUMC) or Rush Oak Park Hospital (ROPH), Chicago
	Country: USA
	Dates: 27 March-9 April 2020
	Symptoms and severity: 79 (39.5%) hospitalised including 29 in ICU, 76 (38%) ambulatory care including 55 seen in a designated COVID-19 screening clinic), and 45 (23%) seen at ED
	Demographics: mean age 50 years (SD 17 years), 92 (46%) men
	Exposure history: not stated
Index tests	Test name: ID NOW (no product code)
	Manufacturer: Abbott
	Antigen target: RdRp
	Antibody: N/A
	Test method: isothermal amplification test
	Samples used: NP swabs in 3 mL VTM (collection not reported)
	Transport media: M4-RT VTM (Remel, Lenexa, KS)



Moore 2020 (Continued)	
	Sample storage: stored at 4 $^{\circ}\text{C}$ if all testing could not be completed on the same day; all tests completed within 72 h of collection
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: as per manufacturer
	Blinding reported: not stated
	Timing of samples: not stated; presumably on presentation but no information on symptom status
Target condition and reference stan-	Reference standard: RT-PCR; 2 methods used in the study
dard(s)	 modified CDC RT-PCR (positive result required Ct < 40 for both targets; negative if neither target detected and positive amplification curve for control (RP) gene; inconclusive if only 1 target detected at Ct < 40, and test repeated)
	Abbott RealTime SARS-CoV-2 RT-PCR (amplification curves reported as detected or not detected)
	Record review used to verify status of 8 samples positive on RealTime assay and negative (6) or inconclusive (2) on CDC assay (all considered disease-positive)
	Definition of non-COVID cases: single RT-PCR negative
	Genetic target(s):
	 N1, N2 N, RdRp
	Samples used: NP swabs in VTM, as for index test
	Timing of reference standard: not stated
	Blinded to index test: not stated
	Incorporated index test: no
Flow and timing	Time interval between index and reference tests: all 3 tests conducted within 72 h of sample collection
	All participants received same reference standard: no? (all received both RT-PCR tests, only discordant results on RT-PCR had record review)
	Missing data: none reported, no participant flow diagram reported
	Uninterpretable results: 2 results were invalid on ID NOW and were not retested (excluded)
	Indeterminate results (index test): none reported
	Indeterminate results (reference standard): discordant results between 2 RT-PCR assays had record review to determine presence/absence COVID-19 infection
	Unit of analysis: participants (specimens from 200 unique participants)
Comparative	
Notes	Funding: none reported (some reagents supplied from NIH)
	Publication status: preprint
	Source: medRxiv
	Author COI: no COI statement was reported



Moore 2020 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecula	r tests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it prespecified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		



Moore 2020	(Continued)
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Could the reference standard, its
conduct, or its interpretation have
introduced bias?

Unclear risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

Low concern

DOMAIN 4: Flow and Timing

Was there an appropriate interval be-
tween index test and reference stan-
dard?

Yes

Did all patients receive the same reference standard?

Yes

Were all patients included in the analy-

Unclear

Did all participants receive a reference

standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

Unclear risk

Test name: Xpert Xpress SARS-CoV-2 assay (no product code)

Moran 2020

Index tests

Study characteristics

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity and specificity: - specimens collected from inpatients and ambulatory patients at the University of Chicago
	Recruitment: not stated
	Prospective or retrospective: not stated
	Number of samples (samples with confirmed SARS-CoV-2): 103 (42)
Patient characteristics and setting	Setting: inpatient and ambulatory; samples selected from central laboratory
	Location: Clinical Microbiology Laboratory, University of Chicago
	Country: USA
	Dates: not stated
	Symptoms and severity: not stated
	Demographics: not stated
	Exposure history: not stated



Moran 2020 (Continued)	
	Manufacturer: Cepheid, Sunnyvale, CA
	Antigen target: E, N (N2 region)
	Antibody: N/A
	Test method: rapid PCR
	Samples used: 8 nasal and 95 NP swabs
	Transport media: none described
	Sample storage: not stated
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: not stated; re-testing using Xpert Xpress was undertaken for an N-gene positive result due discrepancy with RT-PCR (not in line with IFU recommendation)
	Blinding reported: not stated
	Timing of samples: not stated
Target condition and reference standard(s)	Reference standard: Roche cobas SARS-CoV-2 assay on the cobas 6800 system (Roche Molecular Systems, Branchburg, NJ)
	Definition of non-COVID cases: single RT-PCR negative
	Genetic target(s): ORF1, E
	Samples used: nasal and NP swabs; same as for index test
	Timing of reference standard: not stated
	Blinded to index test: not stated
	Incorporated index test: no
Flow and timing	Time interval between index and reference tests: not stated; same sample and appear to have both been conducted soon after sample collection
	All participants received same reference standard: yes
	Missing data: none reported, no participant flow diagram reported
	Uninterpretable results: none reported
	Indeterminate results (index test): single FP (negative on E gene and low positive on N gene) was retested with Xpert Xpress and considered negative on both targets
	Indeterminate results (reference standard): single FP was retested on RT-PCR and found to be repeatedly negative
	Unit of analysis: refers to participants
Comparative	
Notes	Funding: none described
	Publication status: accepted manuscript
	Source: Journal of Clinical Microbioloby



Moran 2020 (Continued)

Author COI: no COI statement was reported

Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	No		
Could the conduct or interpretation of the index test have introduced bias?		High risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	



Moran 2020 (Continued)

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Yes
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Unclear
Did all participants receive a reference standard?	Yes

Yes

Could the patient flow have introduced bias?

Were results presented per patient?

Unclear risk

Nagura-Ikeda 2020

Study characteristics	
Patient Sampling	Single group study of patients with laboratory confirmed COVID-19 referred for isolation and treatment (n=103); participants had undergone qRT-PCR tests using NP or OP swabs collected at public health institutes or hospitals (presumably symptomatic), asymptomatic patients were tested as a result of mass-screening due to an outbreak or family cluster
	Recruitment: Not stated
	Prospective or retrospective: NR; samples appear to be collected prospectively but states that patient information was retrospectively collected from the hospital electronic medical records.
Patient characteristics and setting	Setting: Inpatient and asymptomatic (admitted or quarantined)
	Location: Self-Defense Forces Central Hospital, Tokyo
	Country: Japan
	Dates: Feb 11 to May 13, 2020
	Symptoms and severity: 88 (85%) symptomatic, including 16 (15%) severe (showing clinical symptoms of pneumonia - dyspnea, tachypnea, saturation of percutaneous oxygen [SpO2] < 93%, and the need for oxygen therapy); 15 (15%) asymptomatic (including 4 presymptomatic)
	Demographics: IPD provided - median age 46, range 18-87; 66 (64%) male
	Exposure history: Not reported
Index tests	Test name: ESPLINE® SARS-CoV-2 (no product code reported) [Five other tests performed including RT-PCR and RT-LAMP, but not eligible for this review]
	Manufacturer: Fuji Rebio Inc



Nagura-Ikeda 2020 (Continued)	
	Antibody: NP
	Antigen target: Not stated
	Test method: LFA (no reader device required)
	Samples used: Saliva (self-collected)
	Transport media: None; around 500 μL saliva collected
	Sample storage: Stored at -80C until sample preparation
	Test operator: Not stated; implies laboratory staff
	Definition of test positivity: Not stated; appearance of test line implied
	Blinding reported: Not stated
	Timing of samples: saliva collected on admission to hospital; IPD reports this was median 7 days p.s.o (1-14)
Target condition and reference standard(s)	Reference standard: RT-qPCR on initial presentation (RT-PCR was conducted on saliva samples as part of the study but this did not form part of the reference standard diagnosis)
	Definition of non-COVID cases: Single RT-PCR negative
	Genetic target(s): Not reported
	Samples used: NP or OP
	Timing of reference standard: On presentation or as part of mass screening; specific timing in regard to symptom onset was not reported for the original RT-PCR and unclear if same day as saliva collection
	Blinded to index test: Yes
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Unclear; saliva collected on day of admission to quarantine/hospital but NP/OP conducted at some point prior to that
	All patients received same reference standard: Yes
	Missing data: Not stated, no participant flow diagram reported
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported
	Unit of analysis: Patients
Comparative	
Notes	Funding: work was supported by the Health, Labour and Welfare Policy Research Grants, Research on Emerging and Re-emerging Infectious Diseases and Immunization [grant number 20HA2002].
	Publication status: Accepted manuscript
	Source: J Clin Microbiol
	Author COI: The authors declare that they have no conflicts of interests



Nagura-Ikeda 2020 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular	tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	



Nagura-Ik	ceda 2020	(Continued)
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Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN	4: Flow	and	Timing
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Was there an appropriate interval be-
tween index test and reference stan-
dard?

Unclear

Did all patients receive the same reference standard?

Yes

Were all patients included in the analy-

Unclear

Did all participants receive a reference standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

Unclear risk

Nash 2020

Study ch	naracteristics
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Unclear design to estimate sensitivity and specificity: - samples from suspected patients submitted to 'PATH' (ww.path.org) for routine COVID diagnosis [Second cohort of samples also tested using Spike-based assay; excluded as assay requires use of centrifuge) Recruitment: Not stated Prospective or retrospective: Retrospective
Setting: Unclear; samples provided to study authors by PATH (non-profit organisa-
tion), protocol number 00004244
Location: Not reported
Country: Not reported
Dates: Not reported
Symptoms and severity: Not reported
Demographics: Not reported
Exposure history: Not reported
Test name: Direct antigen rapid test (DART TM); NP-based
Manufacturer: E25Bio Inc (Cambridge MA); not yet available
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Nash 2020 (Continued)	
	Antibody: NP
	Antigen target: anti-N mouse monoclonal antibodies
	Test method: immunochromatographic paper-based (CGIA)
	Samples used: Nasal; collection not described
	Transport media: Not stated
	Sample storage: banked frozen prior to testing
	Test operator: Not stated; presume lab staff
	Definition of test positivity: Visual line
	Blinding reported: Not stated
	Timing of samples: Not stated
Target condition and reference standard(s)	Reference standard: qRT PCR; ThermoFisher/ AppliedBiosystems TaqPATH COV-ID-19 Combo Kit (ThermoFisher, Waltham, MA USA)
	Definition of non-COVID cases: As for cases; single negative PCR required
	Genetic target(s): N, S, and ORF1ab genes
	Samples used: Nasal (same swab)
	Timing of reference standard: Not stated
	Blinded to index test: Yes, conducted first
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Simultaneous (Same swab)
	All patients received same reference standard: Yes
	Missing data: None reported
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported
	Unit of analysis: Not stated
Comparative	
Notes	Funding: The study is funded, in part, by a Bill and Melinda Gates Foundation Award (INV-017872) to E25Bio, Inc. EN is funded by Tufts University DISC Seed Grant. MLN is supported by a FAPESP grant (#2020/04836-0) and is a CNPq Research Fellow. AFV is supported by a FAPESP Fellow grant (#18/17647-0). GRFC is supported by a FAPESP Fellow grant (#20/07419-0). BHGAM 798 is supported by a FAPESP Scholarship (#19/06572-2).
	Publication status: pre-print
	Source: medRxiv
	Author COI: BN, AB, AR, MB, NS, AG, IB, and BBH are employed by or affiliated with E25Bio Inc. (www.e25bio.com), a company that develops diagnostics for epidemic viruses.



Nash 2020 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Unclear		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High



Nash 2020 (Continued)

		Timing

Could the patient flow have introduced bias?		Unclear risk
Were results presented per patient?	Unclear	
Did all participants receive a reference standard?	Yes	
Were all patients included in the analysis?	Unclear	
Did all patients receive the same reference standard?	Yes	
Was there an appropriate interval between index test and reference standard?	Yes	

PHE 2020(a)

PHE 2020(a)	
Study characteristics	
Patient Sampling	Set of studies conducted by PHE and University of Oxford. This extraction relates to a two group study estimating sensitivity and specificity: [1] residual frozen swabs from PCR+ in-patients (n=200) [2] residual fresh swab samples from PCR- patients (n=1000) Swabs were sent to PHE Porton Down aafter routine testing See other PHE 2020 extractions for other sub-studies of Innova assay Recruitment: Unclear
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Unclear; appears to be in-patients (samples obtained from secondary health care setting; cases decsribed as from patients admitted to hsopital)
	Location: John Radcliffe Hospital, Oxford (Ag testing at PHE Porton Down)
	Country: UK
	Dates: March-June 2020 (PCR+); August 2020 (PCR-)
	Symptoms and severity: Not stated
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: Innova SARS-CoV-2 Antigen Rapid Qualitative Test
	Manufacturer: Innova Medical Group
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: Naso- and oropharyngeal swabs



DOMAIN 1: Patient Selection	
Item	Authors' judgement Risk of bias Applicability concerns
Methodological quality	
	Author COI: None reported
	Source: Online PHE report
	Publication status: Published
Notes	Funding: PHE evaluation
Comparative	
	Unit of analysis: Patients
	Indeterminate results (reference standard): Unclear
	Indeterminate results (index test): Unclear
	Uninterpretable results: Failure rates reported as: [1] 12/212, 6%; [2] 50/1040, 5.1% NB remaining samples per group (200 and 990) does not match with final numbers re ported (178 and 940), however no explanation given in report.
	Missing data: See below, plus 1 void PCR
	All patients received same reference standard: Yes
Flow and timing	Time interval between index and reference tests: Same swab
	Incorporated index test: No
	Blinded to index test: Not stated
	Timing of reference standard: As for index test
	Samples used: Appears to be same sample as for Ag test
	Genetic target(s): Not stated
	Definition of non-COVID cases: single negative PCR
Target condition and reference standard(s)	Reference standard: RT-PCR; not described. The pre-print supplementary materials describes using the 'Roche platform' under the Phase 3b heading, and also provides the following text under the Phase 2 evaluation heading "Unless otherwise stated, all RT-PCR testing was undertaken on the Roche Cobas® 6800 or 8800 system using their proprietary SARS-CoV-2 assay as per manufacturer's instructions (with off-board lysis using AVL buffer (Qiagen) and 5% Triton-X100 (Sigma Aldrich)). This assay detects OR F-1a/b as a SARS-CoV-2 specific target, and the E-gene as a pan-sarbecovirus target."
	Timing of samples: Not stated
	Blinding reported: Not stated
	Definition of test positivity: Visual line; as per manufacturer
	Test operator: Laboratory staff
	Sample storage: Frozen (PCR+); fresh (PCR-)



PHE 2020(a) (Continued)				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	No			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High	
DOMAIN 2: Index Test (Rapid molecular test	es)			
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	No			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				



PHE 2020(a) (Continued)		
Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	No	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		High risk

PHE 2020(b)

Study characteristics	
Patient Sampling	Set of studies conducted by PHE and University of Oxford. This extraction relates to a single group study estimating sensitivity and specificity: - samples obtained during a COVID-19 outbreak at a Navy barracks (n=157 samples reported in pre-print; 2x2 data provided by study investigators) See other PHE extractions for other sub-studies of Innova assay Recruitment: Unclear; presume consecutive
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Outbreak investigation
	Location: Not stated
	Country: UK
	Dates: Not stated
	Symptoms and severity: Not stated
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: Innova SARS-CoV-2 Antigen Rapid Qualitative Test
	Manufacturer: Innova Medical Group
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: OP swab used; self-collected
	Transport media: VTM
	Sample storage: Transported at 4C to Porton Down for testing



PHE 2020(b) (Continued)			
	Test operator: Laboratory	staff	
	Definition of test positivit	y: Visual line; as per manı	ufacturer
	Blinding reported: Not sta	ated	
	Timing of samples: One w	eek after outbreak; no fu	rther details
Target condition and reference standard(s)	als describes using the 'R vides the following text u stated, all RT-PCR testing using their proprietary SA off-board lysis using AVL I	oche platform' under the nder the Phase 2 evaluati was undertaken on the R kRS-CoV-2 assay as per ma puffer (Qiagen) and 5% Tr	e-print supplementary materi- Phase 3b heading, and also pro- on heading "Unless otherwise oche Cobas® 6800 or 8800 system anufacturer's instructions (with iton-X100 (Sigma Aldrich)). This arget, and the E-gene as a pan-sar
	Definition of non-COVID o	ases: single negative PCR	
	Genetic target(s): Not star	ted	
	Samples used: Appears to	be same sample as for A	g test
	Timing of reference stanc	lard: As for index test	
	Blinded to index test: Not	stated	
	Incorporated index test: N	No	
Flow and timing	Time interval between index and reference tests: Same swab		
	All patients received same	e reference standard: Yes	
	Missing data: None report	ted	
			157, 3.8% (Table 4 of pre-print)NI Juite match with final number re-
	Indeterminate results (ind	dex test): Unclear	
	Indeterminate results (re	ference standard): Unclea	nr
	Unit of analysis: Patients		
Comparative			
Notes	Funding: PHE evaluation		
	Publication status: Publis	shed and unpublished	
	Source: Online PHE repor	t, plus additional data pro	ovided by evaluation team
	Author COI: None reporte	d	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		



Yes		
Unclear		
Unclear		
	Unclear risk	
		Low concern
Unclear		
Yes		
	Unclear risk	
		High
No		
Unclear		
Yes		
	High risk	
		High
Yes		
Yes		
	Unclear Unclear Unclear Ves No Unclear Yes	Unclear Unclear Unclear Ves Unclear risk Vas High risk Yes



PHE 2020(b) (Continued)		
Were all patients included in the analysis?	No	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		High risk

PHE 2020(c) [non-HCW tested]

Study characteristics	
Patient Sampling	Set of studies conducted by PHE and University of Oxford. This extraction relates to a single group study estimating sensitivity and specificity: - individuals presenting at a regional COVID-19 testing centre as part of a Phase 4 community field service evaluation (n=1946; according to Table 3 of pre-print) See other PHE extractions for other sub-studies of Innova assay Recruitment: Not stated; presume consecutive Prospective or retrospective: Not stated
Patient characteristics and setting	Setting: regional COVID-19 testing centres as part of an NHS Test and Trace service evaluation involving the general public
	Location: Not stated
	Country: UK
	Dates: Not stated
	Symptoms and severity: Not stated, presumed 'mainly symptomatic' for purposes of review analyses
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: Innova SARS-CoV-2 Antigen Rapid Qualitative Test
	Manufacturer: Innova Medical Group
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: Anterior nasal and combined oropharyngeal samples
	Transport media: Dry swab
	Sample storage: None; immediate testing
	Test operator: self-trained non-HCW ('Boots' member of staff); described in pre-print a an "operator" or as 'self-trained members of the public'.
	Definition of test positivity: Visual line; as per manufacturer



PHE 2020(c) [non-HCW tested] (Continued)	Blinding reported: Yes; conducted on site		
	Timing of samples: Not stated		
Target condition and reference standard(s)	Reference standard: RT-PCR; no details. The pre-print supplementary materials describes using the 'Roche platform' under the Phase 3b heading, and also provides the following text under the Phase 2 evaluation heading "Unless otherwise stated, all RT-PCR testing was undertaken on the Roche Cobas® 6800 or 8800 system using their proprietary SARS-CoV-2 assay as per manufacturer's instructions (with off-board lysis using AVL buffer (Qiagen) and 5% Triton-X100 (Sigma Aldrich)). This assay detects OR-F-1a/b as a SARS-CoV-2 specific target, and the E-gene as a pan-sarbecovirus target."		
	Definition of non-COVID cases: Cases only study		
	Genetic target(s): Not stated		
	Samples used: Not stated; paired swabs obtained		
	Timing of reference standard: As for index test		
	Blinded to index test: Not stated		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Paired swabs; simultaneous		
	All patients received same reference standard: Yes		
	Missing data: Initial sample of 1946 reported, 27 failed, leaving 1919 for inclusion, however data for only 1686 samples are provided in the pre-print (1314 PCR- in Table 3 and 372 PCR+ in text pg 7), a difference of 233 samples.		
	Uninterpretable results: Failure rate reported as 27/1946 failed, 1.4%		
	Indeterminate results (index test): Unclear		
	Indeterminate results (reference standard): Unclear		
	Unit of analysis: Patients		
Comparative			
Notes	Funding: PHE evaluation		
	Publication status: Published		
	Source: Online PHE report		
	Author COI: none reported		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		



PHE 2020(c) [non-HCW tested] (Continued)			
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tes	sts)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		



PHE 2020(c) [non-HCW tested] (Continued)		
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	No	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		High risk

Study characteristics	
Patient Sampling	Set of studies conducted by PHE and University of Oxford. This extraction relates to a single group study estimating sensitivity alone: - individuals presenting at one of 14 regional drive-through COVID-19 NHS test and trace centres as part of the FALCON C-19 (Facilitating Accelerated Clinical validation Of Novel diagnostics for COVID-19, 20/WA/0169, IRAS 284229) phase 3b study; those with a positive PCR result were asked to return for a re-test within 5 days of the original test result. From the originally published report (Nov 2020) it appears that only participants with samples that were positive on PCR at the second sampling were included.
	PHE 2020(d) [HCW tested] is for health care worker tested samples, and PHE 2020(d) [Lab tested] is for laboratory scientist tested samples See other PHE extractions for other sub-studies of Innova assay
	Recruitment: Not stated; presume consecutive
	Prospective or retrospective: Prospective
	Number of samples (cases): 479 (479); 267 tested by HCWs, 212 tested by laboratory scientists
Patient characteristics and setting	Setting: NHS drive through test and trace centres; no further details
	Location: 14 regional centres
	Country: UK
	Dates: 17 Sept to 23 Oct 2020
	Symptoms and severity: Only described for all 421 included participants in PHE 2020(d) [HCW tested] and PHE 2020(d) [Lab tested] combined: Suppl Table 2 reports 40 (9.5%) asymptomatic, 59 (14%) with no data, leaving 322 with >=1 symptom recorded. It is not stated whether symptoms were present at the time of the original swab or at the time of the second sampling therefore data for the asymptomatic group have not been included in analyses.
	NB: text reports data for 41 asymptomatic and 344 symptomatic from the Phase 3b study (total n = 385)
	Demographics: For the 421 participants: median age 33 y, 168, 40% male
	Exposure history: Not stated
Index tests	Test name: Innova SARS-CoV-2 Antigen Rapid Qualitative Test



PHE 2020(d) [HCW tested] (Continued)

Manufacturer: Innova Medical Group

Antibody: Not stated

Antigen target: Not stated

Test method: Not stated

Samples used: combined anterior nasal and oropharyngeal swabs (1 stored as a dry swab and 1 $\,$

swab placed in VTM; swabs were self-collected

Transport media: Dry swab

Sample storage: None; immediate testing (delay to testing at PHE for [B] is unclear)

Test operator: PHE 2020(d) [HCW tested] HCW on-site, PHE 2020(d) [Lab tested] Laboratory scien-

tist at PHE

Definition of test positivity: Visual line; as per manufacturer

Blinding reported: Yes

Timing of samples: Not stated

Target condition and reference standard(s)

Reference standard: RT-PCR; no details. The pre-print supplementary materials describes using the 'Roche platform' under the Phase 3b heading, and also provides the following text under the Phase 2 evaluation heading "Unless otherwise stated, all RT-PCR testing was undertaken on the Roche Cobas® 6800 or 8800 system using their proprietary SARS-CoV-2 assay as per manufacturer's instructions (with off-board lysis using AVL buffer (Qiagen) and 5% Triton-X100 (Sigma Aldrich)). This assay detects ORF-1a/b as a SARS-CoV-2 specific target, and the E-gene as a pan-sarbecovirus target."

Definition of non-COVID cases:

Genetic target(s): Not stated

Samples used: Appears to be combined NOP swabs in VTM; obtained at same time as second sampling for Ag testing (5 days after 1st positive PCR)

Timing of reference standard: As for index test

Blinded to index test: Not stated

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: appears to be simultaneous (if 2nd PCR result was used).

All patients received same reference standard: Yes

Missing data: Initial sample of 267 reported, 27 failed, leaving 240 for inclusion however data for only 223 HCW tested samples are provided in the pre-print (text pg 7). The original report (Nov 2020) documented 16 samples in this cohort that were either PCR- (n=15) or void (n=1) presumably at the time of the second sampling (as only PCR+ were invited for Ag testing. Although the numbers don't quite add up, it seems likely that this could explain the difference between the 240 and 223 samples.

Uninterpretable results: Failure rates reported as: [A] 28/296, 10.4%; [B] 9/221, 4.2%

Indeterminate results (index test): Unclear

Indeterminate results (reference standard): Unclear

Unit of analysis: Patients



PHE 2020(d) [HCW tested] (Continued)

Comparative

Notes

Method	اماما		
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Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Yes	'	
Did the study avoid inappro- priate inclusions?	Yes		
Could the selection of pa- tients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen	tests)		
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or inter- pretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 2: Index Test (Rapid m	olecular tests)		
DOMAIN 3: Reference Standard	I		
Is the reference standards likely to correctly classify the target condition?	Yes		



PHE 2020(d) [HCW tested] (Contin	ued)		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	No		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	

PHE 2020(d) [Lab tested]

Patient Sampling

Set of studies conducted by PHE and University of Oxford. This extraction relates to a single group study estimating sensitivity alone:

- individuals presenting at one of 14 regional drive-through COVID-19 NHS test and trace centres as part of the FALCON C-19 (Facilitating Accelerated Clinical validation Of Novel diagnostics for COV-ID-19, 20/WA/0169, IRAS 284229) phase 3b study; those with a positive PCR result were asked to return for a re-test within 5 days of the original test result. From the originally published report (Nov 2020) it appears that only participants with samples that were positive on PCR at the second sampling were included.

PHE 2020(d) [HCW tested] is for health care worker tested samples, and PHE 2020(d) [Lab tested] is for laboratory scientist tested samples



PHE 2020(d)	Lab tested	(Continued)
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See other PHE extractions for other sub-studies of Innova assay

Recruitment: Not stated; presume consecutive

Prospective or retrospective: Prospective

Number of samples (cases): 479 (479); 267 tested by HCWs, 212 tested by laboratory scientists

Patient characteristics and setting

Setting: NHS drive trhough test and trace centres; no further details

Location: 14 regional centres

Country: UK

Dates: 17 Sept to 23 Oct 2020

Symptoms and severity:

Only described for all 421 included participants in PHE 2020(d) [HCW tested] and PHE 2020(d) [Lab tested] combined: Suppl Table 2 reports 40 (9.5%) asymptomatic, 59 (14%) with no data, leaving 322 with >=1 symptom recorded. It is not stated whether symptoms were present at the time of the original swab or at the time of the second sampling therefore data for the asymptomatic group have not been included in analyses .

NB: text reports data for 41 asymptomatic and 344 symptomatic from the Phase 3b study (total n = 385)

Demographics: For the 421 participants: median age 33 y, 168, 40% male

Exposure history: Not stated

Index tests

Test name: Innova SARS-CoV-2 Antigen Rapid Qualitative Test

Manufacturer: Innova Medical Group

Antibody: Not stated

Antigen target: Not stated

Test method: Not stated

Samples used: combined anterior nasal and oropharyngeal swabs (1 stored as a dry swab and 1 swab placed in VTM; swabs were self-collected

Transport media: Dry swab

Sample storage: None; immediate testing (delay to testing at PHE for [B] is unclear)

Test operator: PHE 2020(d) [HCW tested] HCW on-site, PHE 2020(d) [Lab tested] Laboratory scien-

tist at PHE

Definition of test positivity: Visual line; as per manufacturer

Blinding reported: Yes for [A] unclear for [B]

Timing of samples: Not stated

Target condition and reference standard(s)

Reference standard: RT-PCR; no detailsThe pre-print supplementary materials describes using the 'Roche platform' under the Phase 3b heading, and also provides the following text under the Phase 2 evaluation heading "Unless otherwise stated, all RT-PCR testing was undertaken on the Roche Cobas® 6800 or 8800 system using their proprietary SARS-CoV-2 assay as per manufacturer's instructions (with off-board lysis using AVL buffer (Qiagen) and 5% Triton-X100 (Sigma Aldrich)). This assay detects ORF-1a/b as a SARS-CoV-2 specific target, and the E-gene as a pan-sarbecovirus target."



PHE 2020	(d)	[Lab tested	(Continued)
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Definition of non-COVID cases:

Genetic target(s): Not stated

Samples used: Appears to be combined NOP swabs in VTM; obtained at same time as second sampling for Ag testing (5 days after 1st positive PCR)

phing for hig testing (5 days after 1st positive ren

Timing of reference standard: As for index test

Blinded to index test: Not stated

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: appears to be simultaneous (if 2nd PCR result was

used).

All patients received same reference standard: Yes

Missing data: Initial sample of 212 reported, 9 failed, leaving 203 for inclusion however data for only 198 lab scientist tested samples are provided in the pre-print (text pg 7). The original report (Nov 2020) documented 8 samples in this cohort that were PCR- presumably at the time of the second sampling (as only PCR+ were invited for Ag testing. Although the numbers don't quite add up, it seems likely that this could explain the difference between the 203 and 198 samples.

Uninterpretable results: Failure rate reported as: 9/212, 4.2%

Indeterminate results (index test): Unclear

Indeterminate results (reference standard): Unclear

Unit of analysis: Patients

Comparative

Notes Funding: PHE evaluation

Publication status: Published Source: Online PHE report

Author COI: None reported

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of pa- tients have introduced bias?		High risk	



PHE 2020(d) [Lab tested] (Continue	ed)			
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen	tests)			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High	
DOMAIN 2: Index Test (Rapid m	olecular tests)			
DOMAIN 3: Reference Standard	i			
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
Reference standard does not incorporate result of index test?	Yes			
Could the reference stan- dard, its conduct, or its inter- pretation have introduced bias?		Unclear risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	No			



PHE 2020(d) [Lab tested] (Continued)			
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?	High risk		

PHE 2020(e)

Study characteristics	
Patient Sampling	Set of studies conducted by PHE and University of Oxford. This extraction relates to a single group study estimating specificity alone: - PHE and hospital staff volunteering for testing (n=538) See other PHE extractions for other sub-studies of Innova assay
	Recruitment: Not stated; presume consecutive
	Prospective or retrospective: Not stated
Patient characteristics and setting	Setting: Screening
	Location: PHE and John Radcliffe Hospital, Oxford
	Country: UK
	Dates: Not stated
	Symptoms and severity: Not stated; hospital staff described as asymptomatic
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: Innova SARS-CoV-2 Antigen Rapid Qualitative Test
	Manufacturer: Innova Medical Group
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: N OP swab for PHE staff; NP swab for hospital staff. All self-collected
	Transport media: Dry swab
	Sample storage: None; immediate testing
	Test operator: Not stated; presumably laboratory scientist at PHE



PHE 2020(e) (Continued)			
	Definition of test positivit	y: Visual line; as per man	ufacturer
	Blinding reported: Unclea	nr	
	Timing of samples: Not st	ated	
Target condition and reference standard(s)	The pre-print supplement the Phase 3b heading, and uation heading "Unless of Roche Cobas® 6800 or 880 manufacturer's instructio	tary materials describes of also provides the follow therwise stated, all RT-PC 00 system using their proins (with off-board lysis uh)). This assay detects Ol	gative PCR ok for asymptomatic). using the 'Roche platform' under ving text under the Phase 2 eval-CR testing was undertaken on the prietary SARS-CoV-2 assay as per sing AVL buffer (Qiagen) and 5% RF-1a/b as a SARS-CoV-2 specific vt."
	DGenetic target(s): Not st	ated	
	Samples used: Not stated	; presume same or paired	d swab
	Timing of reference stand	lard: As for index test	
	Blinded to index test: Not	stated	
	Incorporated index test: N	lo	
Flow and timing	Time interval between index and reference tests: Unclear, may have been a few day		
	All patients received same	e reference standard: Yes	
			ospital staff and 212 PHE staff), 3 leaving 534 for inclusion. Data fo
	Uninterpretable results: F 8.9% (PHE)	ailure rate reported as 1	7/358, 4.7% (hospital) 19/212,
	Indeterminate results (ind	dex test): Unclear	
	Indeterminate results (ref	erence standard): Unclea	ar
	Unit of analysis: Patients		
Comparative			
Notes	Funding: PHE evaluation		
	Publication status: Publis	hed	
	Source: Online PHE repor	t	
	Author COI: none reported	d	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	No		



PHE 2020(e) (Continued)	w.		
Did the study avoid inappropriate exclusions?	Yes		
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		



PHE 2020(e) (Continued)		
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		High risk

Porte 2020a

Patient Sampling	Two-group study to estimate sensitivity and specificity for diagnosis of active disease: - samples from suspected COVID-19 cases (n = 1453) with deliberate sampling of PCR-positive and negative cases on a 2:1 basis (n = 127)
	Recruitment: convenience sampling
	Prospective or retrospective: retrospectively
	Number of samples (samples with confirmed SARS-CoV-2): 127 (82)
Patient characteristics and setting	Setting: outpatients attending ED at private medical centre (hospital)
	Location: Clínica Alemana, Santiago
	Country: Chile
	Dates: 16-21 March 2020
	Symptoms and severity: cough 94 (74.6%) Fever 77 (61.1%) Median duration of symptoms of 2 days (IQR 1–4; range 0-12) Duration of symptoms: day 0-3 91 (72.2%); day 4-7 27 (22.4%); day ≥ 8 8 (6.3%)
	Demographics: 68 male (53.5%), median age 38 years (IQR 29.5–44; range 1–91)
	Exposure history: not stated
Index tests	Test name: diagnostic Kit for 2019-Novel Coronavirus (2019-nCoV) Ag Test (Cat. N° YRLF04401025, lot N° 2002N408)
	Manufacturer: Bioeasy Biotechnology Co., Shenzhen, China
	Antigen target: SARS-CoV-2 nucleocapsid protein
	Antibody: not stated
	Test method: FIA
	Samples used: remnant OP and NP swabs in 3 mL UTM
	Transport media: UTM-RT System, Copan Diagnostics, Murrieta, CA, USA
	Sample storage: stored at 4 °C and tested within 48 h
	Test operator: laboratory technician
	Definition of test positivity: not stated; test "automatically delivers a positive or negative qualitative result"



Porte 2020a (Continued)			
	Positive or negative defin	ned qualitatively	
	Blinding reported: yes		
	Timing of samples: on pr Within 48 h of the PCR te duration of symptoms re	st but it doesn't say when PCR t	est was performed (median
Target condition and reference standard(s)		PCR (COVID-19 Genesig Real-Tim)); Ct ≤ 40 considered positive	ne PCR assay (Primer Design
	Definition of non-COVID	cases: single RT-PCR negative	
	Genetic target(s): not sta	ted	
	Samples used: as for inde	ex test; same OP and NP swabs	used
	Timing of reference stand	dard: median 2 d post symptom	onset (IQR 1-4; range 0-12)
	Blinded to index test: yes	(index test done within 48 h of	PCR test)
	Incorporated index test:	no	
Flow and timing	Time interval between in	dex and reference tests: same s	ample used; within 48 h
	All participants received same reference standard: yes		
	Missing data: None; partipant flow diagram reported		
	Uninterpretable results: not reported		
	Indeterminate results (index test): not reported		
	Indeterminate results (reference standard): not reported		
	Unit of analysis: participa	ants	
Comparative			
Notes	Funding: this work did no	ot receive funding	
	Publication status: preprint (not peer-reviewed)		
	Source: SSRN		
	Author COI: all study aut	nors declare no competing inter	ests
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of pa-	No		
tients enrolled?			
tients enrolled?	No		
·	No Unclear		



Porte 2020a (Continued)			
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Yes		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		



Porte 2020a (Continued)

Could the patient flow have introduced bias?

Low risk

Porte 2020b [A]

Study characteristics	
Patient Sampling	Multi group study to estimate sensitivity and specificity: (1) Covid-19 patients presenting within 5 days of symptom onset (n=32) (2) symptomatic patients with negative PCR (n=20) (3) asymptomatic patients screened prior to surgery (n=12) [27 PCR+ and 19 PCR- samples were used in Weitzel 2020 (different assays)]
	Recruitment: Not stated; appears to be convenience
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Private clinic (classed as Emergence Dept)
	Location: Clínica Alemana, Santiago
	Country: Chile
	Dates: Not stated
	Symptoms and severity: Not reported; 12 asymptomatic
	Demographics: Total sample median age 39 y (IQR 36.7-57); 33, 52% male
	Exposure history: Not reported
Index tests	Comparative study of two Ag tests (no product codes reported); Porte 2020b [A] data relate to test [A], see Porte 2020b [B] tests [B] data.
	[A] SOFIA SARS Antigen FIA [B] STANDARD® F COVID-19 Ag FIA
	Manufacturer:
	[A] Quidel Corporation, San Diego, CA, USA [B] SD Biosensor Inc, Gyeonggi-do, Republic of Korea
	Antibody: NP (both)
	Antigen target: Not stated
	Test method: Both FIA
	Samples used: naso-oropharyngeal flocked swabs; obtained by trained personnel
	Transport media: UTM-RT® System, Copan Diagnostics
	Sample storage: stored at -80 degrees C following RT-PCR
	Test operator: Laboratory staff
	Definition of test positivity: As per manufacturer; both using analyzer device
	Blinding reported: Yes; blinded to RT-PCR result
	Timing of samples: All <5 days p.s.o; median



Porte 2020b [A] (Continued)	PCR+: 2 days (IQR 1-3); PC	R-: 1 day (IQR 0.75-4)	
Target condition and reference standard(s)	Reference standard: RT-PCR; COVID-19 Genesig®, Primerdesign Ltd., Chandler´s Ford, UK; (Ct) values ≤40 were considered positive		
	Definition of non-COVID c	ases: As for cases	
	Genetic target(s): Not stat	red	
	Samples used: NOP; as fo	r index test	
	Timing of reference stand	ard: Not stated	
	Blinded to index test: Unc	lear	
	Incorporated index test: N	lo	
Flow and timing	Time interval between inc	dex and reference tests: S	Simultaneous; same sample
	All patients received same	e reference standard: Yes	5
	Missing data: None report	ed, no participant flow c	liagram reported
	Uninterpretable results: N	lone reported	
	Indeterminate results (ind	dex test): None reported	
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Patients		
Comparative			
Notes	Funding: This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.		
	Publication status: Publis	hed	
	Source: Int J Infect Dis		
	Author COI: All authors de	clare no competing inte	rests
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		High risk	



Porte 2020b [A] (Continued)			
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	



orte 2020b [B]	
Study characteristic	s
Patient Sampling	Comparative study of two Ag tests; Porte 2020b [A] reports full study characteristics and QUADAS
Patient characteris- tics and setting	
Index tests	Comparative study of two Ag tests (no product codes reported); Porte 2020b [B] data relate to test [B], see Porte 2020b [A] for data relate to test [A] and QUADAS entries
	[A] SOFIA SARS Antigen FIA [B] STANDARD® F COVID-19 Ag FIA
	Manufacturer:
	[A] Quidel Corporation, San Diego, CA, USA [B] SD Biosensor Inc, Gyeonggi-do, Republic of Korea
	Antibody: NP (both)
	Antigen target: Not stated
	Test method: Both FIA
	Samples used: naso-oropharyngeal flocked swabs; obtained by trained personnel
	Transport media: UTM-RT® System, Copan Diagnostics
	Sample storage: stored at -80 degrees C following RT-PCR
	Test operator: Laboratory staff
	Definition of test positivity: As per manufacturer; both using analyzer device
	Blinding reported: Yes; blinded to RT-PCR result
	Timing of samples: All <5 days p.s.o; median PCR+: 2 days (IQR 1-3); PCR-: 1 day (IQR 0.75-4)
Target condition and reference stan- dard(s)	Comparative study of two Ag tests; Porte 2020b [A] reports full study characteristics and QUADAS
Flow and timing	Comparative study of two Ag tests; Porte 2020b [A] reports full study characteristics and QUADAS
Comparative	
Notes	

Rhoads 2020	
Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity: - samples positive using standard of care testing (n = 96) (14 negative controls (UTM) included to control for carry-over contamination only)
	Recruitment: convenience



thoads 2020 (Continued)	Drochostivo or retrochostivo vaturama etiva (vanama et
	Prospective or retrospective: retrospective (remnant samples)
	Number of samples (samples with confirmed SARS-CoV-2): 96 (96)
Patient characteristics and setting	Setting: not stated; includes self-collected and provided-collected samples
	Location: not stated; author institutions University Hospitals Cleveland Medical Cen tre
	and Case Western Reserve University
	Country: USA
	Dates: not stated
	Symptoms and severity: not stated
	Demographics: not stated
	Exposure history: not stated
Index tests	Test name: ID NOW (product codes not reported)
	Manufacturer: Abbott; Chicago, USA Also reports evaluation of Diasorin Simplexa (not eligible for this review)
	Antigen target: not stated
	Antibody: N/A
	Test method: isothermal amplification test
	Samples used: nasal swabs (self-collected) and NP swabs (provider collected); all remnant samples
	Transport media: nasal swabs (2 mL normal saline) and NP swabs (3 mL UTM)
	Sample storage: not stated
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: not stated; as per manufacturer
	Blinding reported: not stated
	Timing of samples: not stated
Target condition and reference standard(s)	Reference standard: standard of care testing for original samples; remnant samples re-tested with modified CDC RT-PCR (using 7500 Fast instrument and using alternate RNA extraction method (Maxwell RSC 6 instrument with Viral TNA Kit (Cat# AS1330; Promega, Madison, USA)); samples with 1 positive target detected considered positive instead of "inconclusive"
	Definition of non-COVID cases: as for index test
	Genetic target(s): N1 and N2
	Samples used: as for index test
	Timing of reference standard: as for index test
	Blinded to index test: as for index test
	Blinded to index test: as for index test Incorporated index test: as for index test



Rhoads 2020 (Continued)				
	All participants received s	ame reference standard: ye	es	
	Missing data: none reported, no participant flow diagram reported Uninterpretable results: none reported Indeterminate results (index test): none reported			
	Indeterminate results (reference standard): RT-PCR detected only 1 of 2 targets for 2 samples (both considered positive (diagnosed as positive on original sample testing); both were negative on index test)			
	Unit of analysis: not state	d; only samples reported		
Comparative				
Notes	Funding: no outside fundi	ng used to support the inve	estigation	
	Publication status: accept	ted manuscript		
	Source: Journal of Clinica	l Microbioloby		
	Author COI: COI not menti	oned by study authors		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	No			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included pa- tients and setting do not match the re- view question?			High	
DOMAIN 2: Index Test (Antigen tests)				
DOMAIN 2: Index Test (Rapid molecular test	s)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



Rhoads 2020 (Continued)			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Unclear risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		

Schildgen 2020 [A]

bias?

Were results presented per patient?

Could the patient flow have introduced

childgen 2020 [A] Study characteristics	
Patient Sampling	Unclear design; appears to be single cohort with deliberate sampling of PCR+/
	[1] RT-PCR positive BAL or throat wash samples (n=42)
	[2] RT-PCR negative samples (n=31)
	Described as pilot sample panel
	Recruitment: Appears to be convenience
	Prospective or retrospective: Not stated; presume retrospective

Unclear risk

Unclear



Schildgen 2020 [A] (Continued)

Patient characteristics and setting Setting: Not stated Location: Authors institution: Kliniken der Stadt Köln gGmbH (Koln city clinics) Country: Germany Dates: Not stated Symptoms and severity: Not stated for BAL samples, throat wash from 23 symptomatic and 27 asymptomatic people. Demographics: Not stated Exposure history: Not stated Index tests Comparative study of three Ag tests (no product codes reported); Schildgen 2020 [A] data relate to test [A], see Schildgen 2020 [B] and Schildgen 2020 [C] for data relate to tests [B] and [C]. Test name: [A] **BIOCREDIT** [B] Panbio [C] SARS-CoV-2 Rapid Antigen test Manufacturer: [A] RapiGEN [B] Abbott [C] Roche Antibody: Not stated Antigen target: Not stated Test method: All LFA Samples used: BAL (n=13); throat wash (n=50, including 27 from asymptomatic) Transport media: Not stated Sample storage: Not stated Test operator: Not stated; presume lab staff Definition of test positivity: As per manufacturer Blinding reported: Not stated Timing of samples: Not stated Target condition and reference standard(s) Reference standard: RT-PCR; RealStar® SARS-CoV-2 RT-PCR Kit, Altona, Germany Definition of non-COVID cases: As for cases Genetic target(s): Not stated Samples used: BAL or throat wash; As per index test Timing of reference standard: Not stated Blinded to index test: Not stated Incorporated index test: No



Schildge	n 2020 [<i>A</i>	(Continued)
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Semila Sem 2020 [A] (continued)				
Flow and timing	Time interval between index and reference tests: Same swab			
	All patients received same reference standard: Yes			
	Missing data: 8 PCR invalid samples also tested; 2/8 invalid in one AG assay each, 3/8 negative in all 3 Ag assays			
	Uninterpretable results: I	None reported		
	Indeterminate results (in	dex test): None reported	I	
	Indeterminate results (re	ference standard): None	ereported	
	Unit of analysis: Unclear			
Comparative				
Notes	Funding: The study did no	ot receive any external f	unding	
	Publication status: prepr	int		
	Source: medRxiv			
	Author COI: The authors declare that they have no conflicts of interest			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	No			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the in- dex test have introduced bias?		Unclear risk		



Schildgen 2020 [A] (Continued)			
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		
Could the patient flow have introduced bias?		High risk	

Schildgen 2020 [B]

Semagen 2020 [D]	
Study characteristics	5
Patient Sampling	Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS
Patient characteris- tics and setting	Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS
Index tests	Comparative study of three Ag tests (no product codes reported); Schildgen 2020 [B] data relate to test [B], see Schildgen 2020 [A] and Schildgen 2020 [C] for data relate to tests [A] and [C], and for QUADAS entries.
	Test name:
	[A] BIOCREDIT



Schildgen 2020 [B] (Continued)

[B] Panbio

[C] SARS-CoV-2 Rapid Antigen test

Manufacturer:

[A] RapiGEN

[B] Abbott

[C] Roche

Antibody: Not stated

Antigen target: Not stated

Test method: All LFA

Samples used: BAL (n=13); throat wash (n=50, including 27 from asymptomatic)

Transport media: Not stated

Sample storage: Not stated

Test operator: Not stated; presume lab staff

Definition of test positivity: As per manufacturer

Blinding reported: Not stated
Timing of samples: Not stated

Target condition and reference standard(s)

Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS

Flow and timing

Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS

Comparative

Notes

Schildgen 2020 [C]

Study characteristics	S
Patient Sampling	Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS
Patient characteris- tics and setting	Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS
Index tests	Comparative study of three Ag tests (no product codes reported); Schildgen 2020 [C] data relate to test [C], see Schildgen 2020 [A] and Schildgen 2020 [B] for data relate to tests [A] and [B], and for QUADAS entries.
	Test name:
	[A] BIOCREDIT [B] Panbio [C] SARS-CoV-2 Rapid Antigen test
	Manufacturer:
	[A] RapiGEN [B] Abbott



Schildgen 2020 [C] (Continued)

[C] Roche

Antibody: Not stated

Antigen target: Not stated

Test method: All LFA

Samples used: BAL (n=13); throat wash (n=50, including 27 from asymptomatic)

Transport media: Not stated
Sample storage: Not stated

Test operator: Not stated; presume lab staff

Definition of test positivity: As per manufacturer

Blinding reported: Not stated
Timing of samples: Not stated

Target condition and reference standard(s)

 $Comparative \ study \ of \ three \ Ag \ tests; \ Schildgen \ 2020 \ [A] \ reports \ full \ study \ characteristics \ and \ QUADAS$

dard(s)

Comparative study of three Ag tests; Schildgen 2020 [A] reports full study characteristics and QUADAS

Comparative

Flow and timing

Notes

Scohy 2020

Study characteristics	
Patient Sampling	Single group study including NP swabs submitted to laboratory at a large tertiary hospital (n=148)
	Recruitment: Random sample
	Prospective or retrospective: Not stated
Patient characteristics and setting	Setting: Unclear; presume microbiology laboratory takes samples from number of sources
	Location: Cliniques universitaires Saint-Luc Hospital, Brussels
	Country: Belgium
	Dates: Apr 6 to Apr 21, 2020
	Symptoms and severity: 86 (58%) symptomatic, 45 (30%) asymptomatic, 17 (11%) symptom status not reported; Cases only: viral load <25 Ct 10 (9%), >=25 Ct 96 (91%)
	Demographics: median age 57.5 (0, 94y); 64 (43%) male
	Exposure history: Not reported
Index tests	Test name: COVID-19 Ag Respi-Strip (product code not reported)



scohy 2020 (Continued)			
	Manufacturer: Coris Bioconcept		
	Antibody: NP		
	Antigen target: monoclonal antibody		
	Test method: CGIA		
	Samples used: NP		
	Transport media: Not stated		
	Sample storage: "If the rapid antigen test was not performed immediately, samples were stored at 4 °C until the test"		
	Test operator: Not stated		
	Definition of test positivity: Visual appearance of T line; also states that "Two versions of the test were evaluated. On the second version, conjugate was coupled on a different way and the control line was optimized."		
	Blinding reported: Unclear		
	Timing of samples: Not reported		
Target condition and reference standard(s)	Reference standard: RT-PCR: genesig® Real-Time PCR assay (Primerdesign Ltd, Chandler's Ford, UK); <40 Ct		
	Definition of non-COVID cases: Single PCR negative		
	Genetic target(s): RdRp		
	Samples used: NP; same as for index		
	Timing of reference standard: Not stated		
	Blinded to index test: Yes		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Same sample		
	All patients received same reference standard: Yes		
	Missing data: None reported, no participant flow diagram reported		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (reference standard): None reported		
	Unit of analysis: Patients		
Comparative			
Notes	Funding: No funding statement reported; COVID-19 Ag Respi-Strip tests provided by Coris BioConcept.		
	Publication status: Published		
	Source: J Clin Virol		



Scohy 2020 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High



Scohy 2020 (Continued)

		Timing

Was there an appropriate interval between index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	Unclear	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?		Unclear risk

Shrestha 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: - subjects who were close contacts of confirmed cases identified through contact tracing, residing in quarantine centre (n=113)
	Recruitment: Convenience
	Prospective or retrospective: Not stated; appears prospective
Patient characteristics and setting	Setting: Contact tracing
	Location: Not applicable; author institutions include Shukraraaj Tropical and Infectious Disease Hospital, Kathmandu
	Country: Nepal
	Dates: Aug to Sep 2020
	Symptoms and severity: All asymptomatic
	Demographics: Range 13 to 74; 89, 79% male
	Exposure history: All exposed to confirmed case
Index tests	Test name: BIOCREDIT
	Manufacturer: RapiGen
	Antibody: Not stated
	Antigen target: Not stated
	Test method: Not stated
	Samples used: NP
	Transport media: None used
	Sample storage: None reported; other sample from the same individual was processed for the results as instructed by the manufacturing company of antigen kit



hrestha 2020 (Continued)	Test operator: Lab techni	cian (trained)		
	Definition of test positivity: Visual line; as per manufacturer.			
	Blinding reported: Uncle			
	Timing of samples: Day 5			
Target condition and reference standard(s)	Reference standard: RT-F	PCR; not detailed, "followe		
	Definition of non-COVID	cases: As for cases; single i	negative	
	Genetic target(s): Not sta	ted		
	Samples used: NP in 3mL	.VTM		
	Timing of reference stand	dard: As for index test		
	Blinded to index test: No	t stated		
	Incorporated index test:	No		
Flow and timing	Time interval between in ples	dex and reference tests: S	imultaneous, paired sam-	
	All patients received same reference standard: Yes			
	Missing data: None reported			
	Uninterpretable results: None reported			
	Indeterminate results (index test): Tests were repeated for samples with indistinct outcomes.			
	Indeterminate results (reference standard):			
	Unit of analysis: Patient			
Comparative				
Notes	Funding: No funding stat	ement provided		
	Publication status: Published			
	Source: KATHMANDU UNIVERSITY MEDICAL JOURNAL			
	Author COI: No COI statement provided			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability con- cerns	
DOMAIN 1: Patient Selection				
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled?	No			
Was a consecutive or random sample of patients en-	No Yes			



Shrestha 2020 (Continued)			
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (Antigen tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Unclear risk	



Smithgall 2020 [A]

Study characteristics	
Patient Sampling	Two-group study to estimate sensitivity and specificity: - patients undergoing routine clinical testing by RT-PCR (n = 113)
	Recruitment: unclear; describes deliberate sampling of samples with high, medium and low Ct values on the reference standard RT-PCR
	Prospective or retrospective: unclear; residual swabs used but testing undertaken within 48 h of sample collection
	Number of samples (samples with confirmed SARS-CoV-2): 113 (88)
Patient characteristics and setting	Setting: inpatient and ED (n from each not reported)
	Location: not stated; author institution is Columbia University Irving Medical Centre
	Country: USA
	Dates: 8-13 April 2020
	Symptoms and severity: not stated
	Demographics: 111 adult (range 23-101 years; average 65 years for RT-PCR-positive and 43 years for RT-PCR-negative); 2 paediatric (age 1 day and 5 days) 61, 54% male
	Exposure history: not stated
Index tests	Test name:
	[A] ID NOW (see Smithgall 2020 [B] for details of comparator test) (product codes not reported)
	Manufacturer: [A] Abbott
	Antigen target: [A] RdRp gene
	Antibody: N/A
	Test method: [A] isothermal PCR Samples used: residual NP swabs (collection not described)
	Transport media: 3 mL VTM (M4RT VTM; ThermoFisher Scientific, Waltham, MA) or UTM (UTM; Becton Dickinson and Co., Franklin Lakes, NJ)
	Sample storage: stored at 4 °C; testing completed within 48 h of sample collection
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: automated as per manufacturer
	Blinding reported: not stated
	Timing of samples: not stated; presume on admission or presentation at ED
Target condition and reference standard(s)	Reference standard: RT-PCR with cobas SARS-CoV-2 assay on the 6800 platform (Roche Diagnostics, Indianapolis, IN); threshold not stated, all Ct values < 37 on bot target genes
	Definition of non-COVID cases: not stated; presume single RT-PCR negative



Smithgall 2020 [A] (Continued)	C//) ODE1	/I =	
	Genetic target(s): ORF1 a	-	
	Samples used: as for inde		
	Timing of reference stand		
	Blinded to index test: as f		
	Incorporated index test:	10	
Flow and timing	Time interval between in	dex and reference tests: s	imultaneous; same samples used
	All participants received	same reference standard:	yes
	Missing data: none repor	ted	
	Uninterpretable results:		
	Indeterminate results (in based on detection of E-§		was a presumptive positive 2 target
	Indeterminate results (re	ference standard): none r	reported
	Unit of analysis: participa	ants	
Comparative			
Notes	Funding: none reported		
	Publication status: publis	shed	
	Source: Journal of Clinica	al Virology	
	Author COI: study author	s report no conflicts of int	terest present
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have intro- duced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular tests)		



Smithgall 2020 [A] (Continued)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	No			
Could the conduct or interpretation of the index test have introduced bias?		High risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	No			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	Unclear			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	Yes			
Could the patient flow have introduced bias?		Unclear risk		
Smithgall 2020 [B]				
Study characteristics				
Patient Sampling See Smithgall 2020 [A] for	or full study details	and QUADAS-2 entries		



Smithga	ll 2020 [E	(Continued)
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Patient characteristics and setting

See Smithgall 2020 [A] for full study details and QUADAS-2 entries

Index tests

Test name: [B] Xpert Xpress (product codes not reported) (see Smithgall 2020 [A] for details of comparator

test)

Manufacturer: [B] Cepheid

Antigen target: [B] N2, E genes

Antibody: N/A

Test method: [B] automated RT-PCR

Samples used: residual NP swabs (collection not described)

Transport media: 3 mL VTM (M4RT VTM; ThermoFisher Scientific, Waltham, MA) or UTM (UTM; Becton Dickin-

son and Co., Franklin Lakes, NJ)

Sample storage: stored at 4 °C; testing completed within 48 h of sample collection.

Test operator: not stated; presume laboratory staff

Definition of test positivity: presumptive positive (only E gene present) considered positive (re-testing recom-

mended on IFU)

Blinding reported: not stated

Timing of samples: not stated; presume on admission or presentation at ED

Target condition and reference standard(s)

See Smithgall 2020 [A] for full study details and QUADAS-2 entries

Flow and timing

See Smithgall 2020 [A] for full study details and QUADAS-2 entries

Comparative

Notes

See Smithgall 2020 [A] for full study details and QUADAS-2 entries

SoRelle 2020

Study characteristics	
Patient Sampling	Unclear design to estimate sensitivity and specificity: paired saliva and NP samples from participants symptomatic for COVID-19 (n=83) [Additional saliva samples included for comparison of ID NOW with Xpert Xpress; not extracted for this review] Recruitment: Not stated Prospective or retrospective: Not stated
Patient characteristics and setting	Setting: Unclear Location: From authors institutions: University of Texas Southwestern Medical Center, Dallas
	Country: USA



SoRelle 2020 (Continued)	
	Dates: Not reported
	Symptoms and severity: Not reported
	Demographics: Not reported
	Exposure history: Not reported
Index tests	Test name: ID NOW (no product codes)
	Manufacturer: Abbott Diagnostics
	Antibody: Not stated
	Antigen target: n/a
	Test method: Isothermal PCR
	Samples used: Saliva; collection not described
	Transport media: Not stated
	Sample storage: Not stated
	Test operator: Not stated; presume lab staff
	Definition of test positivity: As per manufacturer
	Blinding reported: Not stated
	Timing of samples: Not stated; chart review of patients with FN results against either RT-PCR (NP) Xpert Xpress (Saliva) (n=9) showed 6/9 tested >2 weeks after symptom onset
Target condition and reference standard(s)	Reference standard: RT-PCR; either Xpert® Xpress SARS-CoV-2 (Cepheid) or Abbott RealTime SARS-CoV-2 (Abbott Molecular) RT-PCR assays; n per assay is not reported
	Definition of non-COVID cases: As for cases (single negative)
	Genetic target(s): Not stated
	Samples used: NP in VTM (paired)
	Timing of reference standard: Not stated
	Blinded to index test: Not stated
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired
	All patients received same reference standard: Yes
	Missing data: None reported, no participant flow diagram reported
	Uninterpretable results: None reported
	Indeterminate results (index test): None reported
	Indeterminate results (reference standard): None reported; presumptive positives not mentioned
	Unit of analysis: Patients?



SoRelle 2020 (Continued)			
Comparative			
Notes	Funding: No funding sta	tement reported	
	Publication status: Publ	ished letter	
	Source: Clin Chim Acta		
	Author COI: No COI state	ement reported	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Unclear		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced		Unclear risk	

setting do not match the review question:		
DOMAIN 2: Index Test (Antigen tests)		

Are there concerns that the included patients and

DOMAIN 2	· Inday Too	+ /Danid m	olocular tosts	۱

bias?

Were the index test results interpreted without knowledge of the results of the reference standard?

Unclear

If a threshold was used, was it pre-specified?

Yes

Could the conduct or interpretation of the index test have introduced bias?

Unclear risk

Are there concerns that the index test, its conduct, or interpretation differ from the review question?

High

Unclear

DOMAIN 3: Reference Standard

Is the reference standards likely to correctly classify the target condition?

No

Were the reference standard results interpreted without knowledge of the results of the index tests?

Unclear

Reference standard does not incorporate result of index test?

Yes



SoRel	le 2020 ((Continued)
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Could the reference standard, its conduct, or its interpretation have introduced bias?	High risk
Are there concerns that the target condition as defined by the reference standard does not match the question?	High
DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Yes
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	Unclear
Did all participants receive a reference standard?	Yes
Were all patients included in the analysis?	

Unclear risk

Yes

Stevens 2020

Were results presented per patient?

Could the patient flow have introduced bias?

Study characteristics	
Patient Sampling	Unclear design to estimate sensitivity and specificity: - selected residual samples from symptomatic and asymptomatic individuals ur dergoing routine testing; selected to represent the full range of Ct values Recruitment: Convenience
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Unclear; laboratory-based, serving adult and pediatric tertiary care hospitals
	Location: Stanford Healthcare Virology Laboratory, Stanford
	Country: USA
	Dates: Mar 31 to Apr 7
	Symptoms and severity: Unclear; 'symptomatic and asymptomatic'; Of 54 cases, 10 (19%) were low viral low (Ct>35)
	Demographics: Not reported
	Exposure history: Not reported
Index tests	Test name: Xpert Xpress (no product code)
	Manufacturer: Cepheid Inc
	Antibody: E, N2
	Antigen target: n/a



	Samples used: NP in VTM Transport media: VTM (MicroTest M4RT, Remel Inc., San Diego, CA) Sample storage: All samples frozen at -80°C prior to testing on the Xpert system Test operator: Not stated; presume lab staff Definition of test positivity: Presence of N2 +/- E gene; E gene only considered presumptive positive Blinding reported: Not stated	
	Sample storage: All samples frozen at -80°C prior to testing on the Xpert system Test operator: Not stated; presume lab staff Definition of test positivity: Presence of N2 +/- E gene; E gene only considered presumptive positive	
	Test operator: Not stated; presume lab staff Definition of test positivity: Presence of N2 +/- E gene; E gene only considered presumptive positive	
	Definition of test positivity: Presence of N2 +/- E gene; E gene only considered presumptive positive	
	presumptive positive	
	Blinding reported: Not stated	
	Timing of samples: Not stated	
arget condition and reference standard(s)	Reference standard: RT-PCR; Panther Fusion SARS-CoV-2 Assay (Hologic, Inc., San Diego, CA); interpreted based on the manufacturer's cycle threshold cut-off value	
	Definition of non-COVID cases: As for cases; single negative	
	Genetic target(s): Two regions of ORF1ab	
	Samples used: NP in VTM; as for index test	
	Timing of reference standard: Not stated	
	Blinded to index test: Yes, conducted first	
	Incorporated index test: No	
low and timing	Time interval between index and reference tests: Same sample	
	All patients received same reference standard: Yes	
	Missing data: 6 samples excluded due to insufficient sample volume	
	Uninterpretable results: 1 RT-PCR positive sample re-tested on Xpert Xpress due to initial interpretation of no results (invalid); Xpert +ve on re-test	
	Indeterminate results (index test): No presumptive positives were observed	
	Indeterminate results (reference standard): 1 RT-PCR positive sample that was negative on both targets for Xpert Xpress (FN) was re-tested on Panther Fusion and found to be negative (TN)	
	Unit of analysis: Unclear	
omparative		
otes	Funding: No funding statement reported	
	Publication status: Accepted manuscript	
	Source: J Appl Lab Med	
	Author COI: No authors declared any potential conflicts of interest.	
lethodological quality		
em	Authors' judgement Risk of bias Applicability concerns	
OMAIN 1: Patient Selection		



No			
Unclear			
Unclear			
Unclear			
	High risk		
		High	
Unclear			
Yes			
	Unclear risk		
		High	
No			
Yes			
Yes			
	High risk		
		High	
Yes			
	Unclear Unclear Unclear Unclear Ves No Yes Yes	Unclear Unclear Unclear High risk Unclear Ves Unclear risk No Yes High risk High risk	Unclear Unclear Unclear High risk High Ves Unclear risk High No Yes High High High High



Stevens 2020 (Continued)		
Did all patients receive the same reference standard?	Yes	
Were all patients included in the analysis?	No	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Unclear	
Could the patient flow have introduced bias?		High risk

Szymczak 2020

Study characteristics		
Patient Sampling	Single group study to estimate sensitivity and specificity: - remnant samples from patients with symptomatic diarrhea submitted for rou tine diagnostic testing (n=79 from 77 patients)	
	Recruitment: Convenience	
	Prospective or retrospective: Retrospective	
Patient characteristics and setting	Setting: Unclear	
	Location: Clinical Microbiology Laboratory at Montefiore Medical Center, New York	
	Country: USA	
	Dates: Apr 21 to May 15 2020	
	Symptoms and severity: All symptomatic for diarrhoea	
	Demographics: Not stated	
	Exposure history: Not stated	
Index tests	Test name: Xpert Xpress (no product code reported)	
	Manufacturer: Cepheid Inc	
	Target gene(s): N2 and E	
	Antigen target: n/a	
	Test method: Automated RT-PCR	
	Samples used: Stool, collection not reported	
	Transport media: Not stated; coated swabs transferred to 1 ml 0.85% saline for testing	
	Sample storage: Stored at 2 to 8C for up to 7 days prior to testing	
	Test operator: Not stated	
	Definition of test positivity: Describes 'following the package insert instructions presumptive positives not reported	



Szymczak 2020 (Continued)	Blinding reported: Yes; conducted first		
	Timing of samples: PCR +ve stool samples collected 0 to 33 days from initial respiratory PCR; 8/27 collected at >=14 days and 6/27 collected at >=21 days		
Target condition and reference standard(s)	Reference standard: RT-PCR; Hologic Panther Fusion		
	Definition of non-COVID cases: As for cases (single PCR negative)		
	Genetic target(s): two ORF1a regions		
	Samples used: Stool, as for index		
	Timing of reference standard: Some samples frozen at -80oC prior to testing with Hologic Panther Fusion		
	Blinded to index test: Unclear		
	Incorporated index test: No		
Flow and timing	Time interval between index and reference tests: Simultaneous; same swabs		
	All patients received same reference standard: Yes		
	Missing data: None reported, no participant flow diagram reported		
	Uninterpretable results: None reported		
	Indeterminate results (index test): discrepant results re-tested with both index and reference test using both a new aliquot and a shared aliquot tested on both instruments on the same day		
	Indeterminate results (reference standard): discrepant results re-tested with both index and reference test using both a new aliquot and a shared aliquot tested on both instruments on the same day		
	Unit of analysis: Samples (79 from 77 patients)		
Comparative			
Notes	Funding: No funding statement reported		
	Publication status: Published		
	Source: J Clin Microbiol		
	Author COI: No COI statement reported		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Unclear		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Yes		
ora the study avoid mappropriate inclusions:			



Szymczak 2020 (Continued)				
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen tests)				
DOMAIN 2: Index Test (Rapid molecular tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Low risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	No			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
Reference standard does not incorporate result of index test?	Yes			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			High	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	Yes			
Were all patients included in the analysis?	Unclear			
Did all participants receive a reference standard?	Yes			
Were results presented per patient?	No			



Szymczak 2020 (Continued)

Could the patient flow have introduced bias?

Unclear risk

Takeda 2020

Study characteristics		
Patient Sampling	Two group study to estimate sensitivity and specificity, in: [1] RT-PCR confirmed COVID-19 samples selected from a total of 88 positive sample during time period (n=62); [2] Random sample of RT-PCR negative samples selected from 1363 negative specimens tested during same time frame (n=100)	
	Recruitment: Unclear for cases (may have been all 'initial' samples tested); random sample of non-cases	
	Prospective or retrospective: Unclear	
Patient characteristics and setting	Setting: Not stated; multiple clinical institutions	
	Location: SRL Inc, Tokyo	
	Country: Japan	
	Dates: early April' also later states 4 day period	
	Symptoms and severity: Not stated; High viral load (< 25 Ct) - 32/60, 53% Low viral load (>=25 Ct) - 28/60, 47%	
	Demographics: Not stated	
	Exposure history: Not stated	
Index tests	Test name: ESPLINE SARS-CoV-2 (no product code reported)	
	Manufacturer: Fujirebio Inc	
	Antibody: SARS-CoV-2 antigen (from IFU)	
	Antigen target: Anti-SARS-CoV-2 monoclonal antibodies (mouse) (from IFU)	
	Test method: LFA using alkaline phosphatase (ALP) labelled antibodies	
	Samples used: NP; collection not reported	
	Transport media: Not described	
	Sample storage: Swabs mixed with sample treatment solution; no storage reported	
	Test operator: Not stated; laboratory staff presumed	
	Definition of test positivity: Visual line, as per manufacturer	
	Blinding reported: Not stated	
	Timing of samples: Not stated but all cases are first samples presumed by authors to be from patient suspected of SARS-CoV-2 for the first time; negative samples were 'probably from COVID-19 patients for monitoring purposes and to check for negative conversion'	
Target condition and reference standard(s)	Reference standard: RT-PCR; QuantiTect Probe RT-PCR Kit (Qiagen).	



Takeda 2020 (Continued)	Definition of non-COVID o	ases: As for cases: single	negative required	
	Genetic target(s): N2			
	Samples used: NP, as for index test			
	Timing of reference standard: Not stated			
	Blinded to index test: Not			
	Incorporated index test: N			
Flow and timing			imultaneous, same samples	
	Time interval between index and reference tests: Simultaneous, same samples All patients received same reference standard: Yes			
			y because not initial samples but	
	Uninterpretable results: N	None reported		
	Indeterminate results (in	dex test): None reported		
	Indeterminate results (re	ference standard): None r	reported	
	Unit of analysis: Patients (for cases), not clear for non-cases			
Comparative				
Notes	Funding: None reported, however laboratory wholly owned by test manufacturer			
	Publication status: Pre-print			
	Source: medRxiv			
	Author COI: SRL Inc. is a s holds all stock of Fujirebi		ngs Inc. Miraca Holdings Inc.	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Unclear			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		Unclear risk		
Are there concerns that the included patients and setting do not match the review question?			Unclear	
DOMAIN 2: Index Test (Antigen tests)				



Takeda 2020 (Continued)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	

Thwe 2020

Study characteristics



hwe 2020 (Continued)	
Patient Sampling	Single group study to estimate sensitivity and specificity: symptomatic patients with paired samples tested with both ID NOW (dry NP swabs) and a real-time RT-PCR assay (NP swabs in VTM) (n=182) [samples with RT-PCR using Xpert Xpress (n=21) were excluded from this review
	Recruitment: Not stated
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Mixed (inpatient and ED); lab-based study
	Location: University of Texas Medical Branch, Galveston
	Country: USA
	Dates: April to May 2020 ('4 weeks data')
	Symptoms and severity: Not stated
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: ID NOW (no product code)
	Manufacturer: Abbott
	Antibody: Not stated
	Antigen target: n/a
	Test method: Isothermal PCR
	Samples used: dry NP swabs
	Transport media: None
	Sample storage: in plain untreated sterile urine collection tubes
	Test operator: Not stated
	Definition of test positivity: As per manufacturer
	Blinding reported: Yes; conducted first
	Timing of samples: Not stated
Target condition and reference standard(s)	Reference standard: One of 4 RT-PCR assays; 1. Abbott RealTime SARS-CoV-2 (Abbott Park, IL, USA) (n=22) 2. Panther Fusion® SARS-COV-2 (San Diego, CA, USA) (n=129) 3. Cepheid Xpert® Xpress SARS-CoV-2 (Sunnyvale, CA, USA)) (n=21; excluded from this review) 4. a laboratory developed test (LDT) (n=10)
	Definition of non-COVID cases: As for cases (single negative)
	Genetic target(s): Not stated
	Samples used: NP in VTM (paired)

Timing of reference standard: Not stated

Blinded to index test: Not stated



Thwe 2020 (Continued)	Incorporated index test:	No			
Flow and timing	Time interval between index and reference tests: Paired				
	All patients received same reference standard: Yes				
	Missing data: None reported (review team excluded 21 samples tested with RT-PCR)				
	Uninterpretable results:	Uninterpretable results: None reported			
	Indeterminate results (index test): None reported				
	Indeterminate results (resis	ference standard): Non	e reported; no discrepant analy-		
	Unit of analysis: Patient				
Comparative					
Notes	Funding: This project did the public, commercial, o		g support from any agencies in		
	Publication status: Published				
	Source: Diagnostic Microbiol Infect Dis				
	Author COI: All authors have no conflict of interest.				
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	Unclear				
Was a case-control design avoided?	Yes				
Did the study avoid inappropriate exclusions?	Unclear				
Did the study avoid inappropriate inclusions?	Unclear				
Could the selection of patients have introduced bias?		Unclear risk			
Are there concerns that the included patients and setting do not match the review question?			Unclear		
			Unclear		
and setting do not match the review question?			Unclear		
and setting do not match the review question? DOMAIN 2: Index Test (Antigen tests)	Yes		Unclear		



Thwe 2020 (Continued)	
Could the conduct or interpretation of the in-	

Low risk

Are there concerns that the index test, its conduct, or interpretation differ from the review question?

Unclear

DOMAIN 3: Reference Standard

dex test have introduced bias?

Is the reference standards likely to correctly classify the target condition?

No

Were the reference standard results interpreted without knowledge of the results of the index

Unclear

Reference standard does not incorporate result of index test?

Voc

Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

Yes

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis?

Unclear

Did all participants receive a reference standard?

Yes

Were results presented per patient?

Yes

Could the patient flow have introduced bias?

Unclear risk

Van der Moeren 2020(a)

Study characteristics

Patient Sampling

Study reports data for two cohorts. Van der Moeren 2020(a) relates to cohort [1] Single group study to estimate sensitivity and specificity: all adults presenting at a single community test centre for COVID-19 testing (n=354) see Van der Moeren 2020(b) for cohort [2] data

[2] Single group study to estimate sensitivity alone: patients with a positive PCR test result at one of 3 community testing facilities who were retested at home within 72h of initial positive result (n=132)

Recruitment: Consecutive; 'all' adults invited to participate



/an der Moeren 2020(a) (Continued)	Prospective or retrospective: Prospective
Patient characteristics and setting	Setting: COVID-19 test centre (community)
	Location: Municipal Health Service (GGD) regional test centre at Breda
	Country: Netherlands
	Dates: Sep 28 to Sep 30
	Symptoms and severity: Not stated; symptomatic
	Demographics: Not stated
	Exposure history: Not stated
Index tests	Test name: BD Veritor System for Rapid Detection of SARS-CoV-2
	Manufacturer: Becton Dickinson
	Antibody: NP
	Antigen target: Not stated
	Test method: LFA; no further detail
	Samples used: NOP; "specimen from the throat and the superficial nasal cavities (bilateral, 2.5 cm proximal from the nostril)"; collected by GGD employee
	Transport media: Direct testing
	Sample storage: stored dry in sterile test tubes and stored and transported on dry ice until processing at the laboratory; tested within 6 hours after collection
	Test operator: trained laboratory technicians
	Definition of test positivity: reported using Analyzer (included in main analysis for review), and by naked eye inspection alone
	Blinding reported: Not stated
	Timing of samples: Not reported; on presentation time pso only provided for PCR+ cases: 12 < 7d; 1 ≥ 7d; 4=no pso data
Target condition and reference stan-	Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott).
dard(s)	Definition of non-COVID cases: As for cases; single negative
	Genetic target(s): E- and RDRP-gene (Cobas) or E-gene and N-gene (Abbott)
	Samples used: NOP; specimen from the throat and nasal cavity up to the nasal bridge
	Timing of reference standard: As for index test
	Blinded to index test: Not stated
	Incorporated index test: No
Flow and timing	Time interval between index and reference tests: Paired
	All patients received same reference standard: Yes; different assays
	Missing data: 2 samples excluded due to RT-PCR coding error [Considered overall low risk of bias due to small numbers]



an der Moeren 2020(a) (Continued)				
	Uninterpretable results: 1 invalid on Ag test			
	Indeterminate results (inc	lex test): None reported		
	Indeterminate results (ref	erence standard): None rep	ported	
	Unit of analysis: Patients			
Comparative				
Notes	Funding: The VRD (antiger Health, Welfare and Sport		provided by the Dutch Ministry of	
	Publication status: Pre-pr	int		
	Source: medRxiv			
			ol Outbreak Management Team of the implementation of the Coro-	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	Yes			
Did the study avoid inappropriate inclusions?	Yes			
Could the selection of patients have introduced bias?		Low risk		
Are there concerns that the included patients and setting do not match the review question?			Low concern	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk		



An der Moeren 2020(a) (Continued) Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molecular te	ests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		Low risk	
/an der Moeren 2020(b)			

Study characteristics	
Patient Sampling	Study reports data for two cohorts. Van der Moeren 2020(b) relates to cohort [2] Single group study to estimate sensitivity alone: patients with a positive PCR test result at one of twp community testing facilities who were retested at home within 72h of initial positive result (n=132) see Van der Moeren 2020(a) for data related to cohort [1] Single group study to estimate sensitivity and specificity: all adults presenting at a single community test centre for COVID-19 testing (n=354)



Patient characteristics and setting Setting: Community Location: Municipal Health Service (GGD) regional test centres at Breda or Roo: Country: Netherlands Dates: Sep 28 to Oct 6 Symptoms and severity: At time of home visit: Asymptomatic 3, 2% (2/3 still PCR +we) Symptomatic 129 (123 still PCR +we) Day <7 65, 50% Day >7 57, 43% Demographics: Not stated Exposure history: Not stated Exposure history: Not stated Index tests Test name: BD Veritor System for Rapid Detection of SARS-CoV-2 Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott)	/an der Moeren 2020(b) (Continued)			
Patient characteristics and setting Setting: Community Location: Municipal Health Service (GGD) regional test centres at Breda or Roo: Country: Netherlands Dates: Sep 28 to Oct 6 Symptoms and severity: At time of home visit: Asymptomatic 3, 2% (2/2 still PCR eve) Symptomatic 129 (123 still PCR eve) Symptomatic 129 (123 still PCR eve) Day <1 65, 30% Day >7 57, 43% Demographics: Not stated Exposure history: Not stated Exposure history: Not stated Test name: BD Veritor System for Rapid Detection of SARS-CoV-2 Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitient, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Recruitment: Unclear; implies 'all' those with positive PCR invited to participate		
Location: Municipal Health Service (GGD) regional test centres at Breda or Roo. Country: Netherlands Dates: Sep 28 to Oct 6 Symptoms and severity: At time of home visit: Asymptomatic 3, 2% (2/2 still PCR +ve) Symptomatic 3, 2% (2/2 still PCR +ve) Day < 7 66, 50% Day > 7 57, 43% Demographics: Not stated Exposure history: Not stated Index tests Test name: BD Veritor System for Rapid Detection of SARS-CoV-2 Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril!"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory; tested within 6 hours after collection Test operator: trained laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Prospective or retrospective: Prospective		
Country: Netherlands Dates: Sep 28 to Oct 6 Symptoms and severity: At time of home visit: Asymptomatic 3, 2% (2/3 still PCR +ve) Symptomatic 129 (123 still PCR +ve) Day < 7 65, 50% Day < 7 57, 43% Demographics: Not stated Exposure history: Not stated Exposure history: Not stated Index tests Test name: BD Veritor System for Rapid Detection of SARS-CoV-2 Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostrill"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory tested within 6 hours after collection Test operator: trained laboratory tested within 6 hours after collection Test operator: trained laboratory tested using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated	Patient characteristics and setting	Setting: Community		
Dates: Sep 28 to Oct 6 Symptoms and severity: At time of home visit: Asymptomatic 3, 2% (2/3 still PCR +ve) Symptomatic 3, 2% (2/3 still PCR +ve) Day -7 66, 50% Day -7 57, 43% Demographics: Not stated Exposure history: Not stated Exposure history: Not stated Index tests Test name: BD Veritor System for Rapid Detection of SARS-CoV-2 Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Location: Municipal Health Service (GGD) regional test centres at Breda or Roosendaal		
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Index tests Test name: BD Veritor System for Rapid Detection of SARS-CoV-2 Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Asymptomatic 3, 2% (2/3 still PCR +ve) Symptomatic 129 (123 still PCR +ve) Day <7 66, 50%		
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Manufacturer: Becton Dickinson Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Exposure history: Not stated		
Antibody: NP Antigen target: Not stated Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated	Index tests	Test name: BD Veritor System for Rapid Detection of SARS-CoV-2		
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Test method: LFA; no further detail Samples used: NOP? "specimen from the throat and the superficial nasal cavitieral, 2.5 cm proximal from the nostril)"; collected by GGD employee Transport media: Direct testing Sample storage: stored dry in sterile test tubes and stored and transported on until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Antibody: NP		
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until processing at the laboratory; tested within 6 hours after collection Test operator: trained laboratory technicians Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Transport media: Direct testing		
Definition of test positivity: reported using Analyzer (included in main analysis view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Sample storage: stored dry in sterile test tubes and stored and transported on dry ice until processing at the laboratory; tested within 6 hours after collection		
view), and by naked eye inspection alone Blinding reported: Not stated Timing of samples: Not reported; on presentation Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Test operator: trained laboratory technicians		
Timing of samples: Not reported; on presentation Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Definition of test positivity: reported using Analyzer (included in main analysis for review), and by naked eye inspection alone		
Target condition and reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott). Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Blinding reported: Not stated		
Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Timing of samples: Not reported; on presentation		
Definition of non-COVID cases: n/a Genetic target(s): E- and RDRP-gene (Roche) or E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated		Reference standard: RT-PCR; either Cobas 6800 (Roche) or the m2000 (Abbott).		
E-gene and N-gene (Abbott) Samples used: NOP; specimen from the throat and nasal cavity up to the nasal Timing of reference standard: As for index test Blinded to index test: Not stated	dard(s)	Definition of non-COVID cases: n/a		
Timing of reference standard: As for index test Blinded to index test: Not stated				
Blinded to index test: Not stated		Samples used: NOP; specimen from the throat and nasal cavity up to the nasal bridge		
		Timing of reference standard: As for index test		
Incorporated index test: No		Blinded to index test: Not stated		
		Incorporated index test: No		
Flow and timing Time interval between index and reference tests: Paired	Flow and timing	Time interval between index and reference tests: Paired		



Van der Moeren 2020(b) (Continued)			
	All patients received same reference standard: Yes; different assays		
	Missing data: Review team excluded 7 no longer PCR+ at time of home visit (1 asymptomatic, 6 symptomatic) - VRD result for 1 asymptomatic PCR- is given (VRD-)		
	Uninterpretable results: None reported		
	Indeterminate results (index test): None reported		
	Indeterminate results (re	ference standard): None re	ported
	Unit of analysis: Patients		
Comparative			
Notes	Funding: The VRD (antige Health, Welfare and Spor		provided by the Dutch Ministry of
	Publication status: Pre-p	rint	
	Source: medRxiv		
			al Outbreak Management Team of the implementation of the Coro-
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled?	Unclear		
Was a consecutive or random sample of	Unclear		
Was a consecutive or random sample of patients enrolled?			
Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclu-	No		
Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Did the study avoid inappropriate inclu-	No Yes	High risk	
Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Did the study avoid inappropriate inclusions? Could the selection of patients have in-	No Yes	High risk	High
Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Did the study avoid inappropriate inclusions? Could the selection of patients have introduced bias? Are there concerns that the included patients and setting do not match the	No Yes	High risk	High
Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Did the study avoid inappropriate inclusions? Could the selection of patients have introduced bias? Are there concerns that the included patients and setting do not match the review question?	No Yes	High risk	High



an der Moeren 2020(b) (Continued)		
Could the conduct or interpretation of the index test have introduced bias?	Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?	High	
DOMAIN 2: Index Test (Rapid molecular to	ests)	
DOMAIN 3: Reference Standard		
s the reference standards likely to cor- rectly classify the target condition?	No	
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear	
Reference standard does not incorporate result of index test?	Yes	
Could the reference standard, its conduct, or its interpretation have introduced bias?	High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?	High	
DOMAIN 4: Flow and Timing		
Was there an appropriate interval be- tween index test and reference standard?	Yes	
Did all patients receive the same reference standard?	Yes	
Nere all patients included in the analysis?	No	
Did all participants receive a reference standard?	Yes	
Were results presented per patient?	Yes	
Could the patient flow have introduced bias?	High risk	
eyrenche 2020		
Study characteristics		
Patient Sampling	Two group study estimating sensitivity and specificity: [1] PCR+ hospital inpatients (n=45) [2] pre-pandemic samples from 'patients' (not otherwise specified) (n=20)	_



/eyrenche 2020 (Continued)	Recruitment: Not stated; appears to be convenience as equal numbers per Ct value
	subgroup
	Prospective or retrospective: Retrospective
Patient characteristics and setting	Setting: Inpatient
	Location: Montpellier University hospitals (Centre Hospitalier Universitaire de Montpellier, Montpellier)
	Country: France
	Dates: 14 March to 11 April
	Symptoms and severity: 27/45, 60% cases 'severe' according to WHO guideline (similar numbers per Ct subgroup)
	Demographics: Median age: Ct<=25 - 66 (IQR 48, 84) Ct 25-35 - 63 (50, 76) Ct>=35 - 58 (49-67) Controls 64 (35, 93); 32/45, 71% male, all controls were male
	Exposure history: Not stated
Index tests	Test name: Coris COVID-19 Ag Respi-Strip
	Manufacturer: BioConcept®, Gembloux, Belgium
	Antibody: NP
	Antigen target: monoclonal ab
	Test method: CGIA
	Samples used: NP; collection not described
	Transport media: Yes; "swabs were collected in various transport media (eSwab™ COPAN Amies 1 ml, Σ-Transwab® liquid Amies, viral transport medium tube VTM-№ 2.0ml)."
	Sample storage: Unclear; RT-PCR conducted prospectively within a few hours but not reported for Ag testing
	Test operator: Not stated; presume lab staff
	Definition of test positivity: Visual, as per manufacturer
	Blinding reported: Not stated
	Timing of samples: day 1 to 20 pso, median Ct<=25 - 7 (4, 10; presume this is IQR but could be range - is described as SD in paper) Ct 25-35 - 8 (4, 12) Ct>=35 - 11 (7, 15)
Target condition and reference standard(s)	Reference standard: RT-PCR; Allplex™ 2019-nCoV Assay (Seegene, Seoul, South Korea)
	Definition of non-COVID cases: pre-pandemic
	Genetic target(s): RdRp, N, E
	Samples used: NP; as for index



Veyrenche 2020 (Continued)				
	Timing of reference stand	dard: As for index		
	Blinded to index test: Yes	, conducted first		
	Incorporated index test: No			
Flow and timing	Time interval between index and reference tests: Simultaneous; same swab All patients received same reference standard: No Missing data: None reported, no participant flow diagram reported Uninterpretable results: None reported			
	Indeterminate results (in	dex test): None reported	d	
	Indeterminate results (re	ference standard): None	e reported	
	Unit of analysis: Patients			
Comparative				
Notes	Funding: supported by Grants from Montpellier University Hospital and Montpell er University (MUSE).			
	Publication status: pre-p	rint		
	Source: medRxiv			
	Author COI: The authors declare that there are no conflicts of interest			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	No			
Did the study avoid inappropriate exclusions?	Unclear			
Did the study avoid inappropriate inclusions?	Unclear			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (Antigen tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre-specified?	Yes			



Veyrenche 2020 (Continued)			
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Unclear
DOMAIN 2: Index Test (Rapid molecular tests)			
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	No		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	

Weitzel 2020 [A]

Study characteristics	
Patient Sampling	Single-group study to estimate sensitivity and specificity: - samples from patients with respiratory symptoms and/or fever attending a private hospital ED
	Recruitment: convenience with deliberate sampling of positive cases to ensure a 2:1 distribution reported (5276 samples processed during study period)



Weitzel 2020 [A] (Continued)

Prospective or retrospective: retrospective

Number of samples (samples with confirmed SARS-CoV-2): 111 (80)

*17 samples included in Porte 2020a

Patient characteristics and setting

Setting: ED (private hospital)

Location: Clínica Alemana de Santiago

Country: Chile

Dates: 16 March-26 April 2020

Symptoms and severity: respiratory symptoms and/or fever; no further detail

Demographics: median age 40 years; 50, 45% male (median age 38 years, 43% male for all sam-

ples tested during period)

Exposure history: none reported

Index tests

Weitzel 2020 [A] entry is for test [A] in the list below

Test name:

[A] Biocredit COVID-19 Ag One Step SARS-CoV-2 Antigen Test (RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea)

[B] COVID-19 Antigen Rapid Test Device StrongStep COVID-19 Antigen Test (Liming Bio-Products Co., Jiangsu, China)

[C] Huaketai New Coronavirus (SARS-CoV-2) N Protein Detection Kit (Fluorescence immunochromatography) (Savant Biotechnology Co., Beijing, China),

[D] Diagnostic Kit for 2019-Novel Coronavirus (2019-nCoV) Ag Test (Fluorescence Immunochromatographic Assay) (Bioeasy Biotechnology Co., Shenzhen, China).

Manufacturer:

[A] RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea

[B] Liming Bio-Products Co., Jiangsu, China

[C] Savant Biotechnology Co., Beijing, China

[D] Bioeasy Biotechnology Co., Shenzhen, China

Antigen target: not reported in study

Antibody: not reported in study

Test method: [A] and [B] CGIA

[C] and [D] FIA

Samples used: NOP swabs in 3 mL UTM

Transport media: UTM-RT System (Copan Diagnostics, Murrieta, CA, USA)

Sample storage: stored at -80 °C; index tests applied on 28 and 29 April 2020

Test operator: single, trained laboratory technician under BSL2 cabinet; visual outputs read by 2 independent observers with referral to third if needed

Definition of test positivity: as per manufacturer; Beijing Savant test required use of manufacturer supplied UV torch due to unavailability of reader device in Chile

Blinding reported: yes; blinding stated

Timing of samples: median 2 days (IQR 1-5 days); 88% (96/109) during the first week of symptoms



Weitzel 2020 [A] (Continued)

Target condition and reference standard(s)

Reference standard: RT-PCR; COVID-19 Genesig Real-Time PCR assay (Primerdesign Ltd., Chan-

dler's Ford, UK). Ct ≤ 40 considered positive

Definition of non-COVID cases: single PCR negative

Genetic target(s): RdRp

Samples used: NOP swabs; as for index

Timing of reference standard: as for index test; median 2 days (IQR 1-5 days)

Blinded to index test: yes; prior to index

Incorporated index test: no

Flow and timing

Time interval between index and reference tests: same samples; index tests conducted after

frozen storage

All participants received same reference standard: yes

Missing data: none reported; evaluation of Liming test was discontinued after initial poor per-

formance (zero TP)

Uninterpretable results: 2 tests had invalid results due to insufficient liquid migration (2 results

excluded for each test)

Indeterminate results (index test): visual interpretation of the Beijing Savant assay (using manufacturer supplied UV torch) was reportedly difficult under daylight conditions; manufacturer's

fluorescence reader not available in Chile.

Indeterminate results (reference standard): none reported

Unit of analysis: participants

Comparative

Notes Funding: study authors report that the work received no funding; Savant Biotechnology Co.

provided test kits free of charge

Publication status: preprint

Source: medRxiv

Author COI: all authors declare no competing interests

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		



Weitzel 2020 [A] (Continued)			
Did the study avoid inappropriate inclusions?	Yes		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen test	es)		
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it prespecified?	Yes		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid molec	cular tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		



Weitzel 2020 [A] (Continued)	
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis?	No
Did all participants receive a reference standard?	Yes
Were results presented per patient?	Yes
Could the patient flow have introduced bias?	High risk

Weitzel 2020 [B]

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Patient Sampling See Weitzel 2020 [A] for full study details and QUADAS entries

Patient characteristics and setting

Index tests

Weitzel 2020 [B] entry is for test [B] in the list below; see Weitzel 2020 [A] for full study details and QUADAS entries

Test name:

[A] Biocredit COVID-19 Ag One Step SARS-CoV-2 Antigen Test (RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea)

[B] COVID-19 Antigen Rapid Test Device StrongStep COVID-19 Antigen Test (Liming Bio-Products Co., Jiangsu, China)

[C] Huaketai New Coronavirus (SARS-CoV-2) N Protein Detection Kit (Fluorescence immunochromatography) (Savant Biotechnology Co., Beijing, China),

[D] Diagnostic Kit for 2019-Novel Coronavirus (2019-nCoV) Ag Test (Fluorescence Immunochromatographic Assay) (Bioeasy Biotechnology Co., Shenzhen, China).

Manufacturer:

- [A] RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea
- [B] Liming Bio-Products Co., Jiangsu, China
- [C] Savant Biotechnology Co., Beijing, China
- [D] Bioeasy Biotechnology Co., Shenzhen, China

Antigen target: not reported in study

Antibody: not reported in study

Test method: [A] and [B] CGIA

[C] and [D] FIA

Samples used: NOP swabs in 3 mL UTM

Transport media: UTM-RT System (Copan Diagnostics, Murrieta, CA, USA)

Sample storage: stored at -80 °C; index tests applied on 28 and 29 April 2020



Weitze	l 2020	[B]	(Continued)
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Test operator: single, trained laboratory technician under BSL2 cabinet; visual outputs read by 2 independent observers with referral to third if needed

Definition of test positivity: as per manufacturer; Savant test required use of manufacturer supplied UV torch due to unavailability of reader device in Chile

Blinding reported: yes; blinding stated

Timing of samples: median 2 days (IQR 1-5 days); 88% (96/109) during the first week of symptoms

Target condition and reference standard(s)

See Weitzel 2020 [A] for full study details and QUADAS entries

Flow and timing

See Weitzel 2020 [A] for full study details and QUADAS entries

Comparative

Notes

Weitzel 2020 [C]

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Stuay	characte	eristics

Patient Sampling See Weitzel 2020 [A] for full study details and QUADAS entries

Patient characteristics and setting

See Weitzel 2020 [A] for full study details and QUADAS entries

Index tests

Weitzel 2020 [C] entry is for test [C] in the list below; see Weitzel 2020 [A] for full study details and QUADAS entries

Test name:

[A] Biocredit COVID-19 Ag One Step SARS-CoV-2 Antigen Test (RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea)

[B] COVID-19 Antigen Rapid Test Device StrongStep COVID-19 Antigen Test (Liming Bio-Products Co., Jiangsu, China)

[C] Huaketai New Coronavirus (SARS-CoV-2) N Protein Detection Kit (Fluorescence immunochromatography) (Savant Biotechnology Co., Beijing, China),

[D] Diagnostic Kit for 2019-Novel Coronavirus (2019-nCoV) Ag Test (Fluorescence Immunochromatographic Assay) (Bioeasy Biotechnology Co., Shenzhen, China).

Manufacturer:

[A] RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea

[B] Liming Bio-Products Co., Jiangsu, China

[C] Savant Biotechnology Co., Beijing, China

[D] Bioeasy Biotechnology Co., Shenzhen, China

Antigen target: not reported in study

Antibody: not reported in study

Test method: [A] and [B] CGIA

[C] and [D] FIA

Samples used: NOP swabs in 3 mL UTM

Transport media: UTM-RT System (Copan Diagnostics, Murrieta, CA, USA)



Weitzel 2020 [C] (Continued)

Sample storage: stored at -80 °C; index tests applied on 28 and 29 April 2020

Test operator: single, trained laboratory technician under BSL2 cabinet; visual outputs read by 2 independent observers with referral to third if needed

Definition of test positivity: as per manufacturer; Savant test required use of manufacturer supplied UV torch due to unavailability of reader device in Chile

Blinding reported: yes; blinding stated

Timing of samples: median 2 days (IQR 1-5 days); 88% (96/109) during the first week of symptoms

Target condition and reference standard(s) See Weitzel 2020 [A] for full study details and QUADAS entries

Flow and timing

See Weitzel 2020 [A] for full study details and QUADAS entries

Comparative

Notes

Weitzel 2020 [D]

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Patient Sampling See Weitzel 2020 [A] for full study details and QUADAS entries

Patient characteristics and setting

See Weitzel 2020 [A] for full study details and QUADAS entries

Index tests

Weitzel 2020 [D] entry is for test [D] in the list below; see Weitzel 2020 [A] for full study details and QUADAS entries

Test name:

[A] Biocredit COVID-19 Ag One Step SARS-CoV-2 Antigen Test (RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea)

[B] COVID-19 Antigen Rapid Test Device StrongStep® COVID-19 Antigen Test (Liming Bio-Products Co., Jiangsu, China)

[C] Huaketai New Coronavirus (SARS-CoV-2) N Protein Detection Kit (Fluorescence immunochromatography) (Savant Biotechnology Co., Beijing, China),

[D] Diagnostic Kit for 2019-Novel Coronavirus (2019-nCoV) Ag Test (Fluorescence Immunochromatographic Assay) (Bioeasy Biotechnology Co., Shenzhen, China).

Manufacturer:

[A] RapiGEN Inc., Anyang-si, Gyeonggi-do, Republic of Korea

[B] Liming Bio-Products Co., Jiangsu, China

[C] Savant Biotechnology Co., Beijing, China

[D] Bioeasy Biotechnology Co., Shenzhen, China

Antigen target: not reported in study

Antibody: not reported in study

Test method: [A] and [B] CGIA

[C] and [D] FIA

Samples used: NOP swabs in 3 mL UTM



Weitzel 2020 [D] (Continued)

Transport media: UTM-RT System (Copan Diagnostics, Murrieta, CA, USA)

Sample storage: stored at -80°C; index tests applied on 28 and 29 April 2020

Test operator: single, trained laboratory technician under BSL2 cabinet; visual outputs read by 2 independent observers with referral to third if needed

Definition of test positivity: as per manufacturer; Savant test required use of manufacturer supplied UV torch due to unavailability of reader device in Chile

Blinding reported: yes; blinding stated

Timing of samples: median 2 days (IQR 1-5 days); 88% (96/109) during the first week of symptoms

Target condition and reference standard(s)

See Weitzel 2020 [A] for full study details and QUADAS entries

Flow and timing

See Weitzel 2020 [A] for full study details and QUADAS entries

Comparative

Notes

Wolters 2020

Study characteristics			
Patient Sampling	2-group study to estimate sensitivity and specificity for diagnosis of active disease: - samples selected from laboratories on the basis of presence/absence of 2 genetic targets on RT-PCR: SARS-CoV-2 E-gene +/RdRp gene + (n = 30); SARS-CoV-2 E-gene +/RdRp gene - (n = 28); SARS-CoV-2 E-gene -/RdRp gene (n = 30) (A separate set of samples were tested in triplicate at all 3 laboratories to determine limits of detection and analytical specificity)		
	Recruitment: not stated; deliberate sampling used		
	Prospective or retrospective: retrospective		
	Sample size (cases): 88 (58)		
Patient characteristics and set-	Setting: not stated; 3 laboratories		
ting	Location: Radboud UMC in Nijmegen, PAMM in Veldhoven and the RIVM in Bilthoven		
	Country: The Netherlands		
	Dates: January-March 2020		
	Symptoms and severity: not stated		
	Demographics: not stated		
	Exposure history: not stated		
Index tests	Test name: Cepheid Xpert Xpress SARS-CoV-2 (product code not reported)		
	Manufacturer: Cepheid Europe		
	Antigen target: E-gene (sarbeco-specific) and N2-gene (SARS-CoV-2-specific)		



Wolters 2020 (Continued)

Antibody: N/A

Test method: not stated (it should be automated PCR)

Samples used: NP or mid-turbinate, and OP swabs

Transport media: UTM or GLY medium; no further details

Test operator: not stated; presume laboratory staff

Definition of test positivity: as per manufacturer; reported E-gene-only positive specimens as presumptive positive but no re-testing with Xpert Xpress was reported. N2-only positives were considered positive (but re-tested with RT-PCR)

Blinding reported: not stated (see comment section)

Timing of samples: not stated

Target condition and reference standard(s)

Reference standard: in-house RT-PCR:

Radboud UMC Lab: MagNApure 96 (Roche) (isolation platform); MagNApure 96 DNA and Viral NA Small Volume (extraction kit); Roche LC480 II (PCR platform); Life Technologies Taqman FastVirus 1-step mastermix (RT-PCR mastermix)

PAMM Lab: Roche cobas 4800 (isolation platform); CT/NG extraction protocol (extraction kit); Roche LC480 II (PCR platform); Roche LightCycler Multiplex RNA Virus Master (RT-PCR mastermix):

RIVM Lab: BioMérieux NucliSens (isolation platform); easyMAG EasyMAG extraction reagents (extraction kit); Thermo Fisher QuantStudio 6 (PCR platform); Life Technologies Taqman FastVirus 1-step mastermix (RT-PCR mastermix)

Definition of non-COVID cases: yes (performed prior to index test)

Genetic target(s): Radboud UMC lab: E-gene and RdRp-gene

PAMM Lab: started with E-gene and RdRp-gene and mid-March moved on to E-gene testing only RIVM Lab: started with E-gene and RdRp-gene and at the beginning of April moved on to E-gene and CDC N1-gene primer and probes

Samples used: as for index test

Timing of reference standard: as for index test

Blinded to index test: storage prior to freezing was not reported; samples were analysed at or near time of collection ("processed ... in the routine diagnostic procedure using the locally implemented RT-PCR")

Incorporated index test: no

Flow and timing

Time interval between index and reference tests: same samples used; index text seems to have been conducted after frozen storage

Missing data: none reported, no participant flow diagram reported

Uninterpretable results: none reported

Indeterminate results (index test): 1 sample was positive only on N2 gene (positive according to IFU) and 1 was positive only on E gene (presumptive positive, requires re-testing according to IFU). Both samples were re-tested on RT-PCR only

Indeterminate results (reference standard): re-testing of the two 'FN' samples (one TP and 1 presumptive positive according to IFU definition) with RT-PCR found both samples to be disease-negative (reclassed as 1 TN and 1 FP); study authors note that the viral loads of these samples are at the limit of detection for Xpert Xpress and that multiple freeze-thaw steps of samples could have had a significant impact on detection.



Nolters 2020 (Continued)	Unit of analysis: not stated;	only samples reported	
Comparative			
Notes	Funding: not stated		
	Publication status: accepted	d manuscript	
	Source: Journal of Clinical V	irology	
	Author COI: the study autho	rs declare no COI present	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen te	sts)		
DOMAIN 2: Index Test (Rapid mole	ecular tests)		
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	No		
Could the conduct or interpre- tation of the index test have in- troduced bias?		High risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard		,	



Wolters 2020 (Continued)			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Unclear		
Could the patient flow have introduced bias?		Unclear risk	

Wong 2020

Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: - samples submitted for routine testing from patients with suspected COVID-19 infection presenting at A&E (n=93), in-patient (n=47) or outpatient n=18) (total n=158 providing 162 samples)
	Recruitment: Not stated
	Prospective or retrospective: Both retrospective (n=74) and prospective (n=88)
Patient characteristics and setting	Setting: Mixed; A&E, inpatient and outpatient



Wong 2020 (Continued)

Location: Prince of Wales Hospital, Hong Kong

Country: China

Dates: Not stated

Symptoms and severity: Not stated

Demographics: Median age 46 (IQR: 35(28-63); males = 69 (44%)

Exposure history: Not stated

Index tests Test name: Xpert Xpress

Manufacturer: Cepheid Inc

Antibody: E and N2
Antigen target: n/.a

Test method: Automated RT-PCR

Samples used: deep throat saliva (DTS) (n=120), or lower respiratory tract (LRT) (n=42; 35 sputum, 6 tracheal aspirate 1 BAL)

Transport media: None; collected in plain sterile container.

Prior to testing, PBS was added to was added into neat DTS specimens (ratio 1:1) and vortexed for homogenization and allowed to settle for 5- 10 min. 2mL of homogenized sample transferred to another vial for centrifugation at 2000 g for 5 min. 1mL of LRT specimens added to 3 mL of in-house prepared Maintenance Medium (MM) (10X Minimum Essential Medium (MEM), 200 mM glutamine, 1 M HEPES, 7.5 % NaHCO3, 12 mg gentamicin, 0.5 mg amphotericin B, 10,000 units penicillin, 10 mg streptomycin, pH 7.1–7.4); mixture was emulsified by pipetting up and down, followed by centrifugation at 2000 g for 5 min. Supernatant was used for testing as per manufacturer's instructions for both RT-PCR and Xpert Xpress

Sample storage: transported to laboratory on the same day and tested promptly

Test operator: Lab staff

Definition of test positivity: As per manufacturer; presumptive positives mentioned only

in Introduction section

Blinding reported: Not stated

Timing of samples: Not stated

Target condition and reference standard(s)

Reference standard: RT-PCR; TIB-Molbiol LightMix® SarbecoV E-gene assay; all positive cases confirmed by reference laboratory of Hong Kong (Public Health Laboratory Ser-

vice Branch, PHLSB).

Definition of non-COVID cases: As for cases (single negative)

Genetic target(s): Not stated

Samples used: DTS or LRT; as per index test

Timing of reference standard: Not stated

Blinded to index test: Yes; conducted first (upon receipt, all samples were screened with

our standard-of-care assay)

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Simultaneous (Same samples)



Wong 2020 (Continued)			
	All patients received same	reference standard: Yes	
	Missing data: None report	ed	
	Uninterpretable results: N	one reported	
	Indeterminate results (ind	ex test): None reported	
	Indeterminate results (ref	erence standard): None re	ported
	Unit of analysis: Samples	(162/158)	
Comparative			
Notes	Funding: This research dic public, commercial, or no		rant from funding agencies in the
	Publication status: Publisl	ned	
	Source: J Clin Virol		
	Author COI: The authors re	eport no declarations of in	terest.
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			Unclear
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular to	ests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	



Wong 2020 (Continued) Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?	Н	ligh risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	Unclear		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	No		

bias?

Could the patient flow have introduced

Young 2020	
Study characteristics	
Patient Sampling	Single group study to estimate sensitivity and specificity: - Patients with one or more symptoms of COVID-19 (within <=7 days post symptom onset) at 21 study sites (n=260) [Second cohort of 361 samples from COVID suspects <=5 days p.s.o. also evaluated to compare BD Veritor with Quidel Sofia® 2 SARS Antigen FIA but excluded from review as only discrepant results on the two Ag assays underwent RT-PCR]
	Recruitment: Not stated
	Prospective or retrospective: Prospective

High risk



Young 2020 (Continued)

Patient characteristics and setting

Setting: Mixed; drive-through/tent (n=42), outpatient clinic (n=74), research clinic (n=72), or skilled nursing facility (n=66)

Location: Unclear; 21 geographically diverse study sites [Author institutions BD Life Sciences, Louisiana State University Health Sciences Center, Tricore Reference Laboratory)

Country: USA

Dates: June 5-11, 2020

Symptoms and severity: 110 (43%) cough, 98 (39%) muscle pain, 95 (37%) headache, 90 (35%) sore throat, 90 (35%) sore throat, 78 (31%) fever.

Of those at <=6 days p.s.o (n=245): 94 (38%) with one symptom, 151 (62%) with >= 2 symptoms

Demographics: median age 43 (range 18 to 90); 91 (36%) male

Exposure history:

Index tests

Test name: BD Veritor SARS-CoV-2 antigen test (no product codes)

Manufacturer: Becton, Dickinson and Company, BD Life Sciences—Integrated Diagnostic Solutions, San Diagn. CA

tions, San Diego, CA

Antibody: NP

Antigen target: not stated

Test method: Not stated; chromatographic immunoassay with analyser

Samples used: Nasal; clinician collected from both nostrils (same swab)

Transport media: dry nasal swabs

Sample storage: Swabs were shipped for testing on dry ice (-70°C);

Test operator: Not stated; Veritor testing was performed internally at BD (San Diego, CA, USA)

Definition of test positivity: As per manufacturer

Blinding reported: Yes; all personnel blinded to all other test results

Timing of samples: All <=7 days p.s.o; median 3.0 d, mean 3.2 d. 38 (15%) 1 day p.s.o, 57 (23%) 2 days, 54 (22%) 3 days, 40 (16%) 4 days, 37 (15%) 5 days, 19 (8%)

6 days, 6 (2%) 7 days

Target condition and reference standard(s)

Reference standard: Lyra® SARS-CoV-2 PCR Assay (Quidel Corporation. Athens, OH); BD MAX™ real time SARS-CoV-2 PCR assay used for discordant testing

Definition of non-COVID cases: As for cases (single negative)

Genetic target(s): Not stated

Samples used: NP (n=217) or OP (n=34); clinician collected (if an NP swab was collected as part of SOC, the participant had the option of having an OP study swab taken in lieu of a second NP swab)

Timing of reference standard: Swabs taken prior to any study swabs (potential for contamination of nasal cavity)

Blinded to index test: Yes; performed at TriCore Reference Laboratories. "All testing was conducted with all personnel blinded to all other test results"

Incorporated index test: No

Flow and timing

Time interval between index and reference tests: Simultaneous (paired)



Young 2020 (Continued)			
	All patients received same	reference standard: Yes	
	Missing data: 9 excluded; (2 on RT-PCR and 1 labelling)		ria and 3 had invalid specimens/results
	Uninterpretable results: 3	invalid on at least one assay	
	Indeterminate results (ind	ex test): None reported	
		onfirmed FN (BD MAX +ve ar	rted. Re-test of 9 'FN' results with BD and sero +ve), 6 were BD Max -ve (incl 1
	Unit of analysis: Patients		
Comparative			
Notes			Company; BD Life Sciences—Integrated ved research funds as part of this work
	Publication status: Pre-pri	nt	
	Source: medRxiv		
			ees of Becton, Dickinson and Company; , None; RA, CEO and PI of Comprehen-
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
Item DOMAIN 1: Patient Selection	Authors' judgement	Risk of bias	Applicability concerns
	Authors' judgement Unclear	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection Was a consecutive or random sam-		Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled? Was a case-control design avoid-	Unclear	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate	Unclear	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions?	Unclear Yes Yes	Risk of bias Unclear risk	Applicability concerns
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Did the study avoid inappropriate inclusions?	Unclear Yes Yes		Applicability concerns Unclear
DOMAIN 1: Patient Selection Was a consecutive or random sample of patients enrolled? Was a case-control design avoided? Did the study avoid inappropriate exclusions? Did the study avoid inappropriate inclusions? Could the selection of patients have introduced bias? Are there concerns that the included patients and setting do	Ves Yes Yes		



Young 2020 (Continued)			
If a threshold was used, was it prespecified?	Yes		
Could the conduct or interpreta- tion of the index test have intro- duced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 2: Index Test (Rapid mole	cular tests)		
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		
Did all participants receive a reference standard?	Yes		
Were results presented per patient?	Yes		
Could the patient flow have introduced bias?		High risk	



Zhen 2020 [A]

Study characteristics	
Patient Sampling	2-group study to estimate sensitivity and specificity: - samples from symptomatic patients of all ages and gender
	Recruitment: not stated; specimens selected to represent the true positivity rate at authors' institution (50% to 60%), and to span low and high viral loads
	Prospective or retrospective: mixed; included frozen samples (n = 88) and prospectively tested (n = 20)
	Number of samples (samples with confirmed SARS-CoV-2):108 (58)
Patient characteristics and setting	Setting: not stated; selected from laboratory
	Location: not stated; authors' institutions were Northwell Health Laboratories, and Dept Pathology and Laboratory Medicine, The Donald and Barbara Zucker School of Medicine
	Country: USA
	Dates: March-April 2020
	Symptoms and severity: "symptomatic"; no further details
	Demographics: not stated (all ages and genders)
	Exposure history: not stated
Index tests	Zhen 2020 [A] is the entry for test [A] from the list below
	Test name:
	[A] Xpert® Xpress SARS-CoV-2 [B] ID NOW COVID-19 (no product codes reported)
	Manufacturer: [A] Cepheid, [B] Abbott
	Antigen target: [A] N2, E; [B] RdRp
	Antibody: N/A
	Test method: rapid PCR
	Samples used: NP swabs
	Transport media: UTM (various manufacturers)
	Sample storage: on collection, stored at 2-8 $^{\rm o}{\rm C}$ for up to 72 h; after routine testing, stored at –80 $^{\rm o}{\rm C}$
	88 samples tested using ePlex on collection, then frozen prior to testing with ID NOW, Xpert Xpress and Hologic RT-PCR; 20 samples tested prospectively after collection on all systems
	Test operator: not stated; presume laboratory staff
	Definition of test positivity: not stated; states "testing was performed according to the manufacturer's instructions" but no presumptive positives reported
	Blinding reported: not stated
	Timing of samples: not stated



Zhen 2020 [A] (Continued)	Study also evaluates [C] G	enMar kePlex® SARS-CoV-2	Test (not eligible for this review)
Target condition and reference standard(s)	Reference standard: RT-PCR; Hologic Panther Fusion SARS-CoV-2 assay, performed cording to manufacturer's IFU		SARS-CoV-2 assay, performed ac-
	Definition of non-COVID ca	ases: single RT-PCR	
	Genetic target(s): 2 region	s of ORF1ab; either positive	
	Samples used: NP swabs;	same as for index test	
	Timing of reference stands	ard: not stated	
	Blinded to index test: not	stated	
	Incorporated index test: n	0	
Flow and timing	tween index and reference		stated in exact terms; delay bese 88 samples tested at time of collecter assays.
	All participants received sa	ame reference standard: ye	S
	Missing data: none reporte	ed, no participant flow diag	ram reported
	Uninterpretable results: 1 dataset	specimen with invalid resul	lt on ID NOW was excluded from that
	Indeterminate results (ind	ex test): none reported; no	re-testing conducted
	Indeterminate results (refe	erence standard): none repo	orted; no re-testing conducted
	Unit of analysis: not stated	l only refers to samples	
Comparative			
Notes	Funding: none stated; stud	dy authors thank Cepheid fo	or providing the reagents used
	Publication status: accept	ed manuscript	
	Source: Journal of Clinical	Microbioloby	
	Author COI: Gregory Berry and Hologic, Inc. and has i		tion seminars for Abbott, Cepheid,
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	Unclear		
Did the study avoid inappropriate inclusions?	Yes		



Zhen 2020 [A] (Continued)			
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (Antigen tests)			
DOMAIN 2: Index Test (Rapid molecular	tests)		
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			High
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	No		
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Unclear		
Reference standard does not incorporate result of index test?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis?	No		



Zhen 2020 [<i>F</i>	(Continued)
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Did all participants receive a reference Yes standard?

Were results presented per patient? Unclear

Could the patient flow have introduced bias?

High risk

Zhen 2020 [B]

Study cho	racteristics
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Patient Sampling See Zhen 2020 [A] for full study details and QUADAS entries

Patient characteristics and setting See Zhen 2020 [A] for full study details and QUADAS entries

Index tests Zhen 2020 [B] is the entry for test [B] from the list below, see Zhen 2020 [A] for full study details and QUADAS

entries

Test name:

[A] Xpert® Xpress SARS-CoV-2

[B] ID NOWCOVID-19

(no product codes reported)

Manufacturer: [A] Cepheid, [B] Abbott

Antigen target: [A] N2, E; [B] RdRp

Antibody: N/A

Test method: isothermal amplification test

Samples used: NP swabs

Transport media: UTM (various manufacturers)

Sample storage: on collection, stored at 2-8 °C for up to 72 h; after routine testing, stored at −80 °C 88 samples tested using ePlex on collection, then frozen prior to testing with ID NOW, Xpert Xpress and Holog-

ic RT-PCR; 20 samples tested prospectively after collection on all systems

Test operator: not stated; presume laboratory staff

Definition of test positivity: not stated; states "testing was performed according to the manufacturer's instruc-

tions" but no presumptive positives reported

Blinding reported: not stated

Timing of samples: not stated

Study also evaluates [C] GenMar kePlex® SARS-CoV-2 Test (not eligible for this review)

Target condition and reference standard(s)

See Zhen 2020 [A] for full study details and QUADAS entries

Flow and timing See Zhen 2020 [A] for full study details and QUADAS entries



Zhen 2020 [B] (Continued)

Comparative

Notes Funding: none stated; study authors thank Cepheid for providing the reagents used

Publication status: accepted manuscript
Source: Journal of Clinical Microbioloby

Author COI: Gregory Berry has previously given education seminars for Abbott, Cepheid, and Hologic, Inc. and

has received Honorariums

BAL: bronchoalveolar lavage; CDC: Center for Disease Control; CGIA: colloidal gold immunoassay; COI: conflict of interest; Ct: cycle threshold; ED: Emergency Department; EUA: emergency use authorisation; FIA: fluorescence immunochromatographic; FN: false negative; FP: false positive; GLY: Glucose-Lactalbumin-Yeast; HCW: healthcare worker; ICU: intensive care unit; IFU: instructions for use; IQR: interquartile range; LDT: laboratory-developed test; N/A: not applicable; NAAT: nucleic acids amplification test; NIH: National Institutes of Health; NOP: naso-oropharyngeal; NP: nasopharyngeal; OP: oropharyngeal; PCR: polymerase chain reaction; PHE: Public Health England; qRT-PCR: quantitative reverse transcription polymerase chain reaction; RNA: ribonucleic acid; RT-PCR: reverse transcription polymerase chain reaction; SD: standard deviation; TA: tracheal aspirate; TN: true negative; TP: true positive; UTM: universal transport medium; UV: ultraviolet; UW: University of Washington; VTM: viral transport medium;

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Ai 2020	Ineligible index test
Anahtar 2020	Ineligible index test
Ar Gouilh 2020	Ineligible index test
Arizti-Sanz 2020	Ineligible index test
Arumugam 2020	Ineligible index test
Avetyan 2020	Ineligible index test
Azhar 2020	Ineligible index test
Azzi 2020	Ineligible index test
Baek 2020	Ineligible index test
Barra 2020	Ineligible study design
Basu 2020	Ineligible reference standard
Behrmann 2020	Accuracy data cannot be extracted
Bokelmann 2020	Ineligible index test
Bordi 2020	Ineligible index test
Brandsma 2020	Ineligible index test
Broughton 2020	Ineligible index test



Study	Reason for exclusion
Bull 2020	Ineligible index test
Bulterys 2020	Ineligible index test
Callahan 2020a	Accuracy data cannot be extracted
Callahan 2020b	Ineligible index test
Chandler-Brown 2020	Ineligible study design
Chen 2020b	Ineligible index test
Chow 2020	Ineligible index test
CNR 2020	Insufficient details in study report
CNR 2020a	Insufficient details in study report
Colson 2020	Inadequate sample size
Comar 2020	Ineligible reference standard
Comer 2020	Ineligible population
Crone 2020	Ineligible index test
Curti 2020	Ineligible study design
Davda 2020	Ineligible index test
Ding 2020a	Ineligible study design
Ding 2020b	Ineligible index test
Dohla 2020	Ineligible index test
Dong 2020	Ineligible index test
El-Tholoth 2020	Ineligible study design
Farfan 2020	Ineligible study design
FIND 2020f	Superseded by Kruger 2020(a)
Fowler 2020	Ineligible index test
Francis 2020	Ineligible study design
Freire-Paspuel 2020a	Ineligible study design
Freire-Paspuel 2020b	Ineligible index test
Ganguli 2020	Ineligible population
Giamarellos-Bourboulis 2020	Ineligible study design



Study	Reason for exclusion
Gonzalez-Gonzalez 2020a	Ineligible study design
Gonzalez-Gonzalez 2020b	Ineligible population
Grant 2020	Ineligible index test
Hass 2020	Ineligible target condition
Herrera 2020	Ineligible reference standard
Hirotsu 2020	Ineligible index test
Hogan 2020a	Ineligible index test
Howson 2020	Ineligible study design
Hu 2020	Ineligible index test
Huang 2020	Ineligible index test
Huang 2021	Ineligible study design
James 2020	Ineligible index test
Jiang 2020	Ineligible index test
Joung 2020	Ineligible index test
Joung 2020a	Ineligible index test
Kalikiri 2020	Ineligible index test
Kashiwagi 2020	Inadequate sample size
Kim 2019	Ineligible study design
Kim 2020	Ineligible index test
Konrad 2020	Ineligible study design
Kurstjens 2020	Ineligible index test
Kyosei 2020	Ineligible study design
Lalli 2020	Inadequate sample size
Lamb 2020	Ineligible study design
Landry 2020	Ineligible index test
Lee 2020	Ineligible index test
Le Hingrat 2020	Ineligible index test
Li 2020	Ineligible index test



Study	Reason for exclusion
Lin 2020	Ineligible population
Liotti 2020a	Ineligible index test
Lowe 2020	Ineligible index test
Lu 2020	Ineligible study design
Lu 2020a	Ineligible index test
Lubke 2020	Ineligible index test
Mahari 2020	Ineligible study design
Marais 2020	Ineligible index test
Marzinotto 2020	Accuracy data cannot be extracted
McCormick-Baw 2020	Ineligible reference standard
McDonald 2020	Ineligible reference standard
McRae 2020	Ineligible index test
Mei 2020	Ineligible index test
Meyerson 2020	Ineligible index test
Michel 2020	Ineligible index test
Mlcochova 2020	Ineligible index test
Mohon 2020	Ineligible index test
Moses 2020	Ineligible index test
Mostafa 2020	Ineligible study design
Muraoka 2020	Ineligible study design
Nachtigall 2020	Ineligible index test
Newman 2020	Ineligible index test
Noerz 2020	Ineligible index test
Ogawa 2020	Inadequate sample size
Osterdahl 2020	Ineligible index test
Paden 2020	Ineligible study design
Patchsung 2020	Ineligible index test
Pellanda 2020	Ineligible index test



Study	Reason for exclusion
Peto 2020	Ineligible index test
Pfefferle 2020	Ineligible study design
Pollock 2020a	Ineligible index test
Qian 2020	Ineligible index test
Rabe 2020	Ineligible population
Rauch 2020	Ineligible index test
Rodel 2020	Ineligible index test
Rodriguez-Manzano 2020	Ineligible index test
Seo 2020	Accuracy data cannot be extracted
Shirato 2020	Ineligible index test
Singh 2020a	Ineligible index test
Singh 2020b	Ineligible index test
Smyrlaki 2020	Ineligible index test
St Hilaire 2020	Ineligible index test
Tan 2020	Ineligible study design
Tanida 2020	Ineligible index test; also preselected on cycle threshold (only < 34 cycle threshold included)
Tibbetts 2020	Ineligible index test
Tran 2020	Ineligible population
Visseaux 2020	Ineligible index test
Wang 2020a	Ineligible index test
Wang 2020b	Accuracy data cannot be extracted
Wang 2020c	Ineligible index test
Wee 2020	Ineligible study design
Wu 2020	Ineligible index test
Xue 2020	Ineligible index test
Yan 2020	Ineligible index test
Yang 2020b	Ineligible index test



Study	Reason for exclusion	
Yu 2020a	Ineligible index test	
Yu 2020b	Ineligible index test	
Yu 2020c	Ineligible index test	
Zamecnik 2020	Ineligible index test	
Zeng 2020	Ineligible study design	
Zhang 2020	Ineligible index test	
Zhao 2020	Ineligible study design	
Zhu 2020	Ineligible index test	

ADDITIONAL TABLES

Table 1. Description of studies

		No. of studies (%)	
Participants		Antigen tests	Rapid molecular
Number of studies		48	29
Sample size (by test type)	Median (IQR)	291.5 (155 to 502.5)	104 (75 to 172)
	Range	56 to 1676	19 to 524
Number of COV- ID-19 cases (by test type)	Median (IQR)	99.5 (45.5 to 128.5)	50 (20 to 88)
	Range	0,951	6, 220
Setting	COVID-19 test centre	22 (46)	0 (0)
	Contacts	4 (8)	0 (0)
	Hospital A&E	3 (6)	3 (10)
	Hospital inpatient	2 (4)	2 (7)
	Laboratory-based	11 (23)	20 (69)
	Mixed	4 (8)	4 (14)
	Unclear	2 (4)	0 (0)
Symptom status	Asymptomatic	3 (6)	0 (0)



• • •	n of studies (Continued) Symptomatic	16 (33)	12 (41)
		11 (23)	0 (0)
	Mainly symptomatic ^a		
	Mixed	8 (17)	3 (10)
	Not reported	10 (21)	14 (48)
Study design			
Recruitment struc- ture	Single group – sensitivity and specificity	29 (60)	17 (59)
	Two or more groups - sensitivity and specificity	10 (21)	7 (24)
	Unclear	2 (4)	2 (7)
	Single group – sensitivity only	6 (13)	3 (10)
	Single group – specificity only	1 (2)	0 (0)
Reference standard for COVID-19 cases	All RT-PCR-positive	47 (98)	29 (100)
		No. of studies = 42	No. of studies = 26
Reference standard for non-COVID-19	COVID suspects (single RT-PCR-negative)	39 (93)	24 (92)
	COVID suspects (double+ RT-PCR-negative)	1 (2)	1 (4)
	Current other disease (RT-PCR-negative)	0 (0)	1 (4)
	Pre-pandemic (not described)	1 (2)	0 (0)
	Pre-pandemic other disease	1 (2)	0 (0)
Tests		No. of evaluations (%	s)
Total number of test evaluations		58	32
Number of tests per study	1	44 (92)	26 (90)
	2	1 (2)	3 (10)
	3	1 (2)	0 (0)
	4	1 (2)	0 (0)
		1 (2)	0 (0)
	5	<u> </u>	0 (0)
Test method	5 CGIA	41 (71)	0 (0)



Table 1. Descripti	on of studies (Continued)		
	LFA (alkaline phosphatase labelled)	2 (3)	0 (0)
	LFA (not otherwise specified)	6 (10)	0 (0)
	Automated RT-PCR	0 (0)	18 (56)
	Isothermal amplification	0 (0)	13 (41)
	Other molecular (PCR + LFA)	0 (0)	1 (3)
Sample type	NP alone	30 (52)	16 (50)
	NP + OP combined	12 (21)	2 (6)
	Nasal alone	2 (3)	2 (6)
	OP alone	1 (2)	1 (3)
	Two or more of NP, or nasal or OP	8 14)	8 (25)
	Saliva	1 (2)	1 (3)
	Other	3 (5)	0 (0)
	Mixed (including lower respiratory)	4 (7)	1 (3)
	Not specified	0 (0)	1 (3)
Sample storage	Direct	28 (48)	7 (22)
	VTM	20 (35)	12 (38)
	Saline	1 (2)	0 (0)
	Direct or VTM	0 (0)	1 (3)
	VTM or PBS	1 (2)	0 (0)
	VTM or other	0 (0)	6 (19)
	Not specified	8 (14)	6 (19)
Sample collection	HCW	15 (26)	2 (6)
	Trained non-HCW	3 (5)	0 (0)
	Self-collected	6 (10)	0 (0)
	HCW or self-collection	0	1 (3)
	Not specified	34 (59)	29 (91)
Sample testing	HCW (on-site)	13 (22)	0
	Trained non-HCW (on-site)	3 (5)	0



Table 1. Description of studies (Con

	HCW or on-site laboratory personnel	0 (0)	1 (3)
	Not specified (on-site testing)	5 (9)	1 (3)
	Laboratory staff	12 (21)	4 (13)
	Not stated (laboratory setting)	15 (26)	16 (50)
IFU compliance	No	16 (28)	16 (50)
	Yes	29 (50)	9 (28)
	Unclear	13 (22)	7 (22)

A&E: accident and emergency department; **CGIA:** colloidal gold immunoassay; **CI:** confidence intervals; **DRW:** Diagnostics for the Real World; **FIA:** fluorescent immunoassay; **HCW:** healthcare worker; **IFU:** instructions for use; **IQR:** inter-quartile range; **LFA:** lateral flow assay; **NP:** nasopharyngeal; **OP:** oropharyngeal; **PBS:** phosphatase-buffered saline; **RT-PCR:** reverse transcription polymerase chain reaction; **VTM:** viral transport medium

Table 2. Antigen tests: summary of sensitivity and specificity analyses

Subgroup	Test	Evalua- tions	Samples	Cases	Average sensitivity, % (95% CI)	Average specificity, % (95% CI)
Overall anal	ysis					
Evaluations r and specificit	eporting both sensitivity y	51	21,614	6136	68.9 (61.8 to 75.1)	99.6 (99.0 to 99.8)
Evaluations r data ^a	eporting sensitivity	57	22,605	7127	67.7 (60.8 to 74.0)	N/A
Evaluations r data ^a	eporting specificity	52	22,152	6136	N/A	99.5 (99.0 to 99.8)
Subgroup an	alyses (with sensitivity	analyses restr	ricting to direc	t comparisor	ns)	
Symptom status (all)	Symptomatic	37	15,530	4410	72.0 (63.7 to 79.0)	99.5 (98.5 to 99.8)
	Asymptomatic	12	1581	295	58.1 (40.2 to 74.1)	98.9 (93.6 to 99.8)
	Difference				-13.8 (-33.1 to 5.4)	-0.6 (-2.6 to 1.4)
					P = 0.159	P = 0.551
	Symptomatic: direct comparison	9	2437	890	68.0 (51.4 to 81.1)	99.2 (83.9 to 100)
	Asymptomatic: direct comparison	9	1182	213	53.6 (35.0 to 71.3)	99.2 (85.5 to 100)

a'mainly' symptomatic indicates \geq 75% of included participants reported as symptomatic.



	Difference				-14.4 (-38.8 to 10.0)	-0.01 (-3.2 to
					P = 0.246	3.2),
						P = 0.995
	Mixed symptoms or not reported	19	6220	2392	63.0 (52.2 to 72.6)	98.4 (98.0 to 98.8)
Time post- symptom	Week 1	26	5769	2320	78.3 (71.1 to 84.1) ^a	N/A
onset	Week 2	22	935	692	51.0 (40.8 to 61.0) ^a	N/A
(sensitivity only)	Difference				-27.3 (-32.8 to -21.9)	
•					P < 0.0001	
	Week 1: direct comparison	22	4978	2164	76.6 (68.2 to 83.4) <i>a</i>	N/A
	Week 2: direct comparison	22	935	692	48.8 (37.9 to 59.8) ^a	N/A
	Difference				-27.9 (-33.3 to -22.5)	
					P < 0.0001	
Ct value (sensitivity	Higher viral load (< or ≤ 25 Ct threshold) ^b	36	2613	2613	94.5 (91.0 to 96.7) ^a	N/A
only)	Lower viral load (> or >= 25 Ct threshold) ^b	36	2632	2632	40.7 (31.8 to 50.3) ^a	N/A
	Difference				-53.8 (-63.6 to -44.1)	
					P < 0.0001	
	Higher viral load (≤ 32 or33 Ct threshold) ^c	15	2127	2127	82.5 (74.0 to 88.6) ^a	N/A
	Lower viral load (> 32 or 33 Ct threshold) ^c	15	346	346	8.9 (3.3 to 21.7) <i>a</i>	N/A
	Difference				-73.5 (-84.7 to -62.4)	
					P < 0.0001	
Study de- sign	Single group: sensitivity and specificity	29	15,336	3536	72.1 (64.8 to 78.3)	99.6 (99.1 to 99.8)
	Two or more groups: sensitivity and speci- ficity	20	5729	2396	64.1 (48.5 to 77.2)	97.3 (96.7 to 97.8)
					-8.0 (-24.2 to 8.2)	-2.3 (-2.9 to −1.
					P = 0.334	P < 0.0001

Table 2. Antigen tests: summary of sensitivity and specificity analyses (Continued)

98.8)

P = 0.113

97.1)

96.0 (94.5 to

100 (96.4 to 100)

15.6 (2.6 to 28.5)

78.0 (46.0 to 93.7)

80.6 (68.6 to 89.6)

P = 0.019

-1.3 (-3.0 to 0.3)



	Unclear	2	549	204	65.2 (39.6 to 84.3)	96.3 (88.0 to 98.9)
Test method	CGIA	36	17,448	5085	64.0 (55.7 to 71.6)	99.0 (98.8 to 99.2)
	FIA	9	2820	712	79.6 (67.5 to 88.0)	97.7 (95.3 to

ALP: alkaline phosphatase labelled; CGIA: colloidal gold immunoassay; CI: confidence intervals; Ct: cycle threshold; FIA: fluorescent immunoassay; LFA: lateral flow assay; N/A: not applicable

277

62

1184

162

LFA (not otherwise

Difference

specified)

LFA (ALP)

5

1

Table 3. Antigen tests: summary data by test brand and compliance with manufacturers' instructions for use

Test	All		IFU-complia	IFU-compliant			
	Number of evalua- tions; sam- ples (cas- es)	Average sensitiv- ity, % (95% CI)	Average specificity, % (95% CI)	Number of evalua- tions; sam- ples (cas- es)	Average sensitivity, % (95% CI)	Average specificity, % (95% CI)	
AAZ - COVID-VIRO (2 studies not pooled)	1; 632 (295)	61.7 (55.9 to 67.3)	100 (98.9 to 100)				
, ,	1; 248 (101)	96.0 (90.2 to 98.9)	86.4 (79.8 to 91.5)	1; 248 (101)	96.0 (90.2 to 98.9)	86.4 (79.8 to 91.5)	
Abbott - Panbio Covid-19 Ag	10; 5509 (1849)	72.0 (60.6 to 81.1)	99.3 (99.0 to 99.6)	5; 1776 (362)	72.0 (56.5 to 83.5)	99.2 (98.5 to 99.5)	
including sensitivity-only cohort	11; 2031 (2031)	72.8 (62.6 to 81.0) ^a		6; 544 (544)	73.5 (61.1 to 83.0) ^a		
Becton Dickinson - BD Veritor	2; 602 (55)	82.3 (62.1 to 93.0)	99.5 (98.3 to 99.8)				
including sensitivity-only cohort	3; 180 (180)	79.4 (72.9 to 84.7) ^a					
BIONOTE - NowCheck COVID-19 Ag	1; 400 (102)	89.2 (81.5 to 94.5)	97.3 (94.8 to 98.8)	1; 400 (102)	89.2 (81.5 to 94.5)	97.3 (94.8 to 98.8)	

^aSeparate pooling of sensitivity or specificity, or both.

 $[^]b$ threshold for 'higher' viral load was < 25 Ct in 18 evaluations and \leq 25 Ct in 18 evaluations

^c threshold for 'higher' viral load ≤ 33 Ct in 13 evaluations and < 32 in 2 evaluations



Se (Continued) Biosynex - Biosynex COVID-19 Ag BSS	1; 634 (297)	59.6 (53.8 to 65.2)	100 (98.9 to 100)			
Coris Bioconcept - COVID-19 Ag Respi-Strip	7; 1781 (707)	39.7 (31.3 to 48.7)	98.3 (97.4 to 98.9)	7; 1781 (707)	39.7 (31.3 to 48.7)	98.3 (97.4 to 98.9)
E25Bio - DART (N-based)	1; 190 (100)	80.0 (70.8 to 87.3)	91.1 (83.2 to 96.1)			
Fujirebio - ESPLINE SARS-CoV-2 (2 studies not pooled)	1; 162 (62)	80.6 (68.6 to 89.6)	100 (96.4 to 100)			
(2 stadies not pooted)	1; 103 (103)	11.6 (6.2 to 19.5)				
Innova Medical Group - Innova SARS-CoV-2 Ag	3; 2945 (596)	47.9 (34.3 to 61.8)	99.8 (99.5 to 99.9)	1; 1676 (372)	57.5 (52.3 to 62.6)	99.6 (99.1 to 99.9)
including sensitivity-only cohorts	5; 1017	59.0 (43.4 to 73.0)a		3; 793	69.1 (58.3 to 78.2)a	
including specificity-only cohort	4; 2887		99.8 (99.5 to 99.9)a	2; 1842		99.7 (99.3 to 99.9)a
Liming Bio-Products - StrongStep® COVID-19 Ag	1; 19 (9)	0 (0 to 33.6)	90.0 (55.5 to 99.7)			
Quidel Corporation - SOFIA SARS Ag	1; 64 (32)	93.8 (79.2 to 99.2)	96.9 (83.8 to 99.9)			
RapiGEN - BIOCREDIT COVID-19 Ag	5; 2010 (310)	63.3 (45.7 to 78.0)	99.5 (99.1 to 99.8)	3; 1828 (189)	73.0 (57.4 to 84.4)	99.8 (99.4 to 99.9)
including sensitivity-only cohort	6; 470 (470)	57.7 (39.8 to 73.8) ^a				
Roche - SARS-CoV-2	1; 73 (42)	88.1 (74.4 to 96.0)	19.4 (7.5 to 37.5)			
Savant Biotech - Huaketai SARS-CoV-2 N Protein	1; 109 (78)	16.7 (9.2 to 26.8)	100 (88.8 to 100)			
SD Biosensor - STANDARD F COVID-19 Ag	4; 1552 (295)	72.6 (54.0 to 85.7)	97.5 (96.4 to 98.2)	2; 1129 (159)	75.5 (68.2 to 81.5)	97.2 (96.0 to 98.1)
SD Biosensor - STANDARD Q COVID-19 Ag	6; 3480 (821)	79.3 (69.6 to 86.6)	98.5 (97.9 to 98.9)	4; 2522 (421)	85.8 (80.5 to 89.8)	99.2 (98.2 to 99.6)
Shenzhen Bioeasy Biotech - 2019-nCoV Ag	3; 965 (177)	86.2 (72.4 to 93.7)	93.8 (91.9 to 95.3)	1; 727 (15)	66.7 (38.4 to 88.2)	93.1 (91.0 to 94.9)
development-phase publication	1; 239 (208)	67.8 (61.0 to 74.1)	100 (88.8 to 100)			

 a Separate pooling of sensitivity or specificity.

 $^{^{\}mathrm{b}}2\mathrm{x}2$ tables combined prior to calculating estimates.



Table 4. Antigen tests: summary data by symptom status, test brand and compliance with manufacturers' instructions for use

	All			IFU-complia	nt	
	Number of evalua- tions; sam- ples (cas- es)	Average sensitiv- ity, % (95% CI)	Average specificity, % (95% CI)	Number of evalua- tions; sam- ples (cas- es)	Average sensitivity, % (95% CI)	Average specificity, % (95% CI)
SYMPTOMATIC participants by te	st					
AAZ - COVID-VIRO	1; 632 (295)	61.7 (55.9 to 67.3)	100 (98.9 to			
(2 studies not pooled)			100)			
	1; 248 (101)	96.0 (90.2 to 98.9)	86.4 (79.8 to 91.5)	1; 248 (101)	96.0 (90.2 to 98.9)	86.4 (79.8 to 91.5)
Abbott - Panbio Covid-19 Ag	8; 3699 (1162)	74.1 (60.8 to 84.0)	99.8 (99.5 to 99.9)	3; 1094 (252)	75.1 (57.3 to 87.1)	99.5 (98.7 to 99.8)
including sensitivity-only cohort	9; 1344 (1344)	74.8 (63.4 to 83.6) ^a		4; 434 (434)	76.2 (63.6 to 85.4) ^a	
Becton Dickinson - BD Veritor	2; 602 (55)	82.3 (62.1 to 93.0)	99.5 (98.3 to 99.8)			
including sensitivity-only cohort	3; 180 (180)	79.4 (72.9 to 84.7) ^a				
BIONOTE - NowCheck COVID-19 Ag	1; 400 (102)	89.2 (81.5 to 94.5)	97.3 (94.8 to 98.8)	1; 400 (102)	89.2 (81.5 to 94.5)	97.3 (94.8 to 98.8)
Biosynex - Biosynex COVID-19 Ag BSS	1; 634 (297)	59.6 (53.8 to 65.2)	100 (98.9 to 100)			
Coris Bioconcept - COVID-19 Ag Respi-Strip	3; 780 (414)	34.1 (29.7 to 38.8) ^a	100 (99.0 to 100)a,b	3; 780 (414)	34.1 (29.7 to 38.8) ^a	100 (99.0 to 100) ^{a,b}
Fujirebio - ESPLINE SARS-CoV-2	1; 88 (88)	11.4 (5.6 to 19.9)				
Innova Medical Group - Innova SARS-CoV-2 Ag	2; 2794 (550)	56.2 (52.0 to 60.3)	99.8 (99.5 to 99.9)	1; 1676 (372)	57.5 (52.3 to 62.6)	99.6 (99.1 to 99.9)
including sensitivity-only cohorts	4; 971 (971)	65.5 (54.8 to 74.9)†		3; 793 (793)	69.1 (58.3 to 78.2)†	
Liming Bio-Products - StrongStep® COVID-19 Ag	1; 19 (9)	0 (0 to 33.6)	90.0 (55.5 to 99.7)			
Quidel Corporation - SOFIA SARS Ag	1; 64 (32)	93.8 (79.2 to 99.2)	96.9 (83.8 to 99.9)			
RapiGEN - BIOCREDIT COVID-19 Ag	3; 608 (206)	58.4 (36.3 to 77.5)	96.4 (82.8 to 99.3)	1; 476 (117)	74.4 (65.5 to 82.0)	98.9 (97.2 to 99.7)



Table 4. Antigen tests: summary data by symptom status, test brand and compliance with manufacturers' instructions for use (Continued)

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Roche - SARS-CoV-2	1; 23 (10)	100 (69.2 to 100)	7.7 (0.2 to 36.0)			
Savant Biotech - Huaketai SARS-CoV-2 N Protein	1; 109 (78)	16.7 (9.2 to 26.8)	100 (88.8 to 100)			
SD Biosensor - STANDARD F COVID-19 Ag	3; 1193 (191)	78.0 (71.6 to 83.3)	97.2 (96.0 to 98.1)	2; 1129 (159)	75.5 (68.2 to 81.5)	97.2 (96.0 to 98.1)
SD Biosensor - STANDARD Q COVID-19 Ag	5; 2760 (731)	80.1 (68.5 to 88.1)	98.1 (97.4 to 98.6)	3; 1947 (336)	88.1 (84.2 to 91.1)	99.1 (97.8 to 99.6)
Shenzhen Bioeasy Biotech - 2019-nCoV Ag	3; 965 (177)	86.2 (72.5 to 93.7)	93.8 (91.9 to 95.3)	1; 727 (15)	66.7 (38.4 to 88.2)	93.1 (91.0 to 94.9)
ASYMPTOMATIC participants by t	est					
Abbott - Panbio Covid-19 Ag	6; 1097 (190)	58.1 (41.7 to 72.9)	98.4 (92.2 to 99.7)	2; 474 (47)	48.9 (35.1 to 62.9)	98.1 (96.3 to 99.1)
Coris Bioconcept - COVID-19 Ag Respi-Strip	1; 45 (14)	28.6 (8.4 to 58.1)	100 (88.8 to 100)	1; 45 (14)	28.6 (8.4 to 58.1)	100 (88.8 to 100)
Fujirebio - ESPLINE SARS-CoV-2	1; 15 (15)	13.3 (1.7 to 40.5)	N/A			
RapiGEN - BIOCREDIT COVID-19 Ag	2; 140 (60)	63.2 (21.7 to 91.4)	98.9 (82.9 to 99.9)	1; 113 (47)	85.1 (71.7 to 93.8)	100 (94.6 to 100)
Roche - SARS-CoV-2	1; 27 (13)	84.6 (54.6 to 98.1)	14.3 (1.8 to 42.8)			
SD Biosensor - STANDARD Q COVID-19 Ag	2; 272 (18)	61.1 (37.9 to 80.2)	99.6 (97.3 to 99.9)	1; 127 (13)	69.2 (38.6 to 90.9)	99.1 (95.2 to 100)

Ag: antigen; CI: confidence interval; N: nucleoprotein; N/A: not applicable

Table 5. Molecular tests: summary of sensitivity and specificity analyses

Test or subgroup	Evalua- tions	Samples	Cases	Average sensitivity, % (95% CI)	Average specificity, % (95% CI)
Overall analysis					
Evaluations reporting both sensitivity and specificity	29	4351	1787	95.1 (90.5 to 97.6)	98.8 (98.3 to 99.2)
Evaluations reporting sensitivity data ^a	32	4537	1973	95.5 (91.5 to 97.7)	N/A
Subgroup analyses (with sensitivity ar	alyses restric	ting to direct c	omparisons)		
Viral load High viral load (≤ 30 Ct)	6	204	204	100 (98.2 to 100) ^{a,b}	N/A

^aseparate pooling of sensitivity or specificity.

b2x2 tables combined prior to calculating estimates.



(sensitivity only)	Low viral load (> 30 Ct)	6	149	149	95.6 (55.7 to 99.7)	N/A
By study design	Single group – sensitivity and specificity	18	2899	976	93.2 (85.5 to 97.0)	99.4 (98.4 to 99.8)
	Two or more groups - sensitivity and specificity	9	1265	718	97.2 (90.7 to 99.2)	99.3 (96.5 to 99.8)
	Difference				4.0 (-2.2 to 10.1)	-0.2 (-1.3 to 1.0)
					P = 0.211	P = 0.771
	Unclear designs	2	187	93	93.2 (71.0 to 98.7) ^a	100 (96.2 to 100) ^{a,b}
Test brand	Abbott – ID NOW	12	1853	634	78.6 (73.7 to 82.8)	99.8 (99.2 to 99.9)
	Cepheid – Xpert Xpress	13	1691	911	99.1 (97.7 to 99.7)	97.9 (94.6 to 99.2)
	Difference				19.8 (14.9 to 24.7)	-1.9 (-3.8 to -0.1)
					P < 0.0001	P = 0.036
	Abbott – ID NOW (including sensitivity only cohort)	13	1949	730	81.5 (75.2 to 86.5) ^a	N/A
	Cepheid – Xpert Xpress (including sensitivity only cohorts)	15	1781	1001	99.1 (97.8 to 99.6) ^a	N/A
	DNANudge – COVID Nudge	1	386	71	94.4 (86.2 to 98.4)	100 (98.8 to 100)
	Diagnostics for the Real World – SAMBA II	2	321	121	96.0 (81.1 to 99.3)	97.0 (93.5 to 98.6)
	Mesa Biotech – Accula	1	100	50	68.0 (53.3 to 80.5)	100 (92.9 to 100)
Test brand	Abbott – ID NOW	4	812	222	73.0 (66.8 to 78.4)	99.7 (98.7 to 99.9)
(restrict- ed to IFU-	Cepheid – Xpert Xpress	2	100	29	100 (88.1 to 100) ^a	97.2 (89.4 to 99.3) ^a
compliant)	DRW – SAMBA II	1	149	33	87.9 (71.8 to 96.6)	97.4 (92.6 to 99.5)
	DNANudge – COVID Nudge	1	386	71	94.4 (86.2 to 98.4)	100 (98.8 to 100)
Discrepant analysis	Before discrepant analysis	6	1533	623	97.9 (88.1 to 99.7)	97.8 (96.6 to 98.6)
	After discrepant analysis	6	1533	632	99.2 (93.6 to 99.9)	99.6 (98.8 to 99.8)
	Difference				1.3 (-2.8 to 5.4)	1.8 (0.7 to 2.8)
					P = 0.528	P = 0.001



Table 5. Molecular tests: summary of sensitivity and specificity analyses (Continued)

CI: confidence interval; Ct: cycle threshold; IFU: [manufacturers'] instructions for use; N/A: not applicable

^aSeparate pooling of sensitivity or specificity.

b2x2 tables combined prior to calculating estimates.

WHAT'S NEW

Date	Event	Description
15 April 2021	Amended	Clarification in Appendices that isothermal amplification is not a RT-PCR test.

HISTORY

Review first published: Issue 8, 2020

Date	Event	Description
24 March 2021	Amended	Amendment to PLS title
24 March 2021	Amended	Correction of typo in abstract
9 March 2021	New citation required and conclusions have changed	This review has been updated and the conclusions have changed
30 September 2020	New search has been performed	We have updated our review and now include 64 study reports in 78 study cohorts, evaluating 16 antigen and 5 molecular assays

CONTRIBUTIONS OF AUTHORS

JD was the contact person with the editorial base.

JDI co-ordinated contributions from the co-authors and wrote the final draft of the review.

JJD, JDi, YT, CD, STP, IH, AA, LFR, MP, MT, JDr, SB screened papers against eligibility criteria.

RS conducted the literature searches.

JDi, MT and AA appraised the quality of papers.

JDi, MT and AA extracted data for the review and sought additional information about papers.

JDi entered data into Review Manager 2020.

 $\ensuremath{\mathsf{JDi}},\ensuremath{\mathsf{JJD}},\ensuremath{\mathsf{YT}}$ and $\ensuremath{\mathsf{SB}},$ analysed and interpreted data.

JJD, JDi, YT, CD, STP, RS, ML, LH, AVB, DE, SD, JC worked on the methods sections and commented on the draft review.

JJD and JDi responded to the comments of the referees.

JJD is the guarantor of the update.

DECLARATIONS OF INTEREST

Jonathan J Deeks: JD has published or been quoted in opinion pieces in scientific publications, and in the mainstream and social media related to diagnostic testing. JD was the statistician on the Birmingham evaluation of the Innova test which is mentioned in the discussion of the paper. There was no funding for this evaluation of the Innova test. JD is a member of the Royal Statistical Society (RSS) COVID-19 taskforce steering group, and co-chair of the RSS Diagnostic Test Advisory Group. He is a consultant adviser to the WHO Essential Diagnostic List. JD receives payment from the BMJ as their Chief Statistical advisor.

Jacqueline Dinnes: none known

Yemisi Takwoingi: none known



Clare Davenport: none known

Mariska MG Leeflang: none known

René Spijker: none known Lotty Hooft: none known

Ann Van den Bruel: none known

Devy Emperador: is employed by FIND with funding from DFID and KFW. FIND is a global non-for profit product development partnership and WHO Diagnostic Collaboration Centre. It is FIND's role to accelerate access to high-quality diagnostic tools for low-resource settings and this is achieved by supporting both R&D and access activities for a wide range of diseases, including COVID-19. FIND has several clinical research projects to evaluate multiple new diagnostic tests against published Target Product Profiles that have been defined through consensus processes. These studies are for diagnostic products developed by private sector companies who provide access to know-how, equipment/reagents, and contribute through unrestricted donations as per FIND policy and external SAC review.

Sabine Dittrich: is employed by FIND with funding from DFID and Australian Aid. FIND is a global non-for profit product development partnership and WHO Diagnostic Collaboration Centre. It is FIND's role to accelerate access to high-quality diagnostic tools for low-resource settings and this is achieved by supporting both R&D and access activities for a wide range of diseases, including COVID-19. FIND has several clinical research projects to evaluate multiple new diagnostic tests against published Target Product Profiles that have been defined through consensus processes. These studies are for diagnostic products developed by private sector companies who provide access to know-how, equipment/reagents, and contribute through unrestricted donations as per FIND policy and external SAC review.

Ada Adriano: none known

Sophie Beese: none known

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Sian Taylor-Phillips: none known

Sarah Berhane: none known

Jane Cunningham: none known

SOURCES OF SUPPORT

Internal sources

- · Liverpool School of Tropical Medicine, UK
- University of Birmingham, UK

External sources

· Department for International Development, UK

Project number: 300342-104

- National Institute for Health Research (NIHR), UK
- NIHR Birmingham Biomedical Research Centre at the University Hospitals Birmingham NHS Foundation Trust and the University of Birmingham, UK

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We planned to check the following websites for eligible index tests, however these did not prove to be very accessible or easy to use and, after initial review, were not further considered:

- National Institute for Health Research (NIHR) Innovation Observatory (www.io.nihr.ac.uk/)
- www.rapidmicrobiology.com/test-method/testing-for-the-wuhan-coronavirus-a-k-a-covid-19-sars-cov-2-and-2019-ncov



We planned to check the following evidence repository for additional eligible studies however, the EPPI-Centre and Norwegian Institute of Public Health resources proved to be more accessible therefore we decided to prioritise our other sources of evidence.

• Meta-evidence (meta-evidence.co.uk/the-role-of-evidence-synthesis-in-covid19/)

We intended for two authors to independently perform data extraction, however one review author extracted study characteristics, and a second author checked them. Contingency table data were extracted independently by two review authors as planned.

We planned to evaluate the effect of additional sources of heterogeneity, including reference standard and sample type. However, additional formal investigations using meta-regression were not possible because of lack of variability across the studies in these features.

We planned to conduct a sensitivity analysis excluding studies that are solely published as preprints. We have inadequate study numbers to allow this at present but will reconsider for the next update.

INDEX TERMS

Medical Subject Headings (MeSH)

Antigens, Viral [*analysis]; Asymptomatic Infections; Bias; Cohort Studies; COVID-19 [*diagnosis]; COVID-19 Nucleic Acid Testing; COVID-19 Serological Testing [*methods] [standards]; False Negative Reactions; False Positive Reactions; Molecular Diagnostic Techniques [*methods] [standards]; *Point-of-Care Systems; Predictive Value of Tests; Reference Standards; SARS-CoV-2 [*immunology]; Sensitivity and Specificity

MeSH check words

Adult; Child; Humans