Interpreting a covid-19 test result

Jessica Watson GP and National Institute for Health Research doctoral research fellow, Penny F Whiting associate professor in clinical epidemiology, John E Brush professor of internal medicine

Centre for Academic Primary Care, Bristol Medical School, University of Bristol, Bristol, UK; Sentara Healthcare and Eastern Virginia Medical School, Norfolk, VA, USA

Across the world there is a clamour for covid-19 testing, with Tedros Adhanom Ghebreyesus, director general of the World Health Organization, encouraging countries to “test, test, test.” The availability of the complete genome of covid-19 early in the epidemic facilitated development of tests to detect viral RNA. Multiple assays with different gene targets have been developed using reverse transcriptase polymerase chain reaction (RT-PCR). These viral RNA tests use samples usually obtained from the respiratory tract by nasopharyngeal swab, to detect current infections. Serology blood tests to detect antibodies indicating past infection are being developed; these will not be useful for the current infections. Serology blood tests to detect antibodies indicating past infection are being developed; these will not be useful for identifying those who are currently infectious.

A simple negative covid-19 test should not be used as a rule-out in patients with strongly suggestive symptoms. A positive RT-PCR test for covid-19 test has more weight than a negative test because of the test’s high specificity but moderate sensitivity. A single negative covid-19 test should not be used as a rule-out in patients with strongly suggestive symptoms. Clinicians should share information with patients about the accuracy of covid-19 tests.

No test gives a 100% accurate result; tests need to be evaluated to determine their sensitivity and specificity, ideally by comparison with a “gold standard.” The lack of such a clear-cut “gold-standard” for covid-19 testing makes evaluation of test accuracy challenging.

A systematic review of the accuracy of covid-19 tests reported false negative rates of between 2% and 29% (equating to sensitivity of 71-98%), based on negative RT-PCR tests which were positive on repeat testing. The use of repeat RT-PCR testing as gold standard is likely to underestimate the true rate of false negatives, as not all patients in the included studies received repeat testing and those with clinically diagnosed covid-19 were not considered as actually having covid-19.

Accuracy of viral RNA swabs in clinical practice varies depending on the site and quality of sampling. In one study, sensitivity of RT-PCR in 205 patients varied, at 93% for broncho-alveolar lavage, 72% for sputum, 63% for nasal swabs, and only 32% for throat swabs. Accuracy is also likely to vary depending on stage of disease and degree of viral multiplication or clearance. Higher sensitivities are reported depending on which gene targets are used, and whether multiple gene tests are used in combination. Reported accuracies are much higher for in vitro studies, which measure performance of primers using coronavirus cell culture in carefully controlled conditions.

The lack of a clear-cut “gold-standard” is a challenge for evaluating covid-19 tests; pragmatically, clinical adjudication may be the best available “gold standard,” based on repeat swabs, history, and contact with patients known to have covid-19, chest radiographs, and computed tomography scans. Inevitably this introduces some incorporation bias, where the test being evaluated forms part of the reference standard, and this would tend to inflate the measured sensitivity of these tests.

Disease prevalence can also affect estimates of accuracy: tests developed and evaluated in populations with high prevalence (eg, secondary care) may have lower sensitivity when applied in a lower prevalence setting (eg, primary care).
One community based study of 4653 close contacts of patients with covid-19 tested RT-PCR throat swabs every 48 hours during a 14 day quarantine period. Of 129 eventually diagnosed with covid-19 by RT-PCR, 92 (71.3%) had a positive test on the first throat swab, equating to a sensitivity of 71% in this lower prevalence, community setting.12

Further evidence and independent validation of covid-19 tests are needed.13 As current studies show marked variation and are likely to overestimate sensitivity, we will use the lower end of current estimates from systematic reviews,6 with the approximate numbers of 70% for sensitivity and 95% for specificity for illustrative purposes.

What do clinicians need to know to understand a test result?

Sensitivity and specificity can be confusing terms that may be misunderstood6 (see supplementary file ‘Definitions and formulae for calculating measures of test accuracy’). Sensitivity is the proportion of patients with disease who have a positive test, or the true positive rate. Specificity is the proportion of patients without disease who have a negative test, or true negative rate. These terms describe the operating characteristics of a test and can be used to gauge the credibility of a test result. They can be combined to calculate likelihood ratios, which are dimensionless numbers that indicate the strength of a positive or negative test result.13 For calculating probabilities, a likelihood ratio can be used as a multiplier to convert pre-test odds to post-test odds. Positive likelihood ratios greater than 1 are progressively stronger, with 10 representing a very strong positive test result. Negative likelihood ratios less than 1 are also progressively stronger, with 0.1 representing a very strong negative test result. In the case of the nasopharyngeal swab RNA test for covid-19, the positive likelihood ratio is about 14, which is excellent.15 A positive covid-19 test result should be very compelling. The negative likelihood ratio is 0.3, which is a moderate result, but not nearly as compelling as a positive result because of the moderate sensitivity (about 70%) of the covid-19 test.

Interpretation of a test result depends not only on the characteristics of the test itself but also on the pre-test probability of disease. Clinicians use a heuristic (a learned mental short cut) called anchoring and adjusting to settle on a pre-test probability (called the anchor). They then adjust this probability based on additional information. This heuristic is a useful short cut but comes with the potential for bias. When people fail to estimate the pre-test probability and only respond to a piece of additional information, they commit a fallacy called base-rate neglect. Another fallacy called anchoring is failing adequately to adjust one's probability estimate, given the strength of new information. Likelihood ratios can give a clinician an idea of how much to adjust their probability estimates. Clinicians intuitively use anchoring and adjusting thoughtfully to estimate pre- and post-test probabilities unconsciously in everyday clinical practice. However, faced with a new and unfamiliar disease such as covid-19, mental short cuts can be uncertain and unreliable and public narrative about the definitive nature of testing can skew perceptions.

Figure 1 shows how a clinician’s thinking about a patient’s probability should shift, based on either a positive or negative test result for covid-19. First, the clinician should estimate a pre-test probability, using knowledge of local rates of covid-19 infection from national6 and regional13 data and patients’ symptoms and signs,14 likelihood of alternative diagnoses, and history of exposure to covid-19. After choosing a pre-test probability on the x axis, one should then trace up to either the upper curve for a positive test result or the lower curve for a negative test result, then trace over to the y axis to read the estimate for post-test probability. The figure shows that the shift in the probability is asymmetric, with a positive test result having a greater impact than a negative test result, owing to the modest sensitivity and negative likelihood ratio of the RNA test.

The infographic (fig 2) shows the outcomes when 100 people with a pre-test probability of 80% are tested for covid-19 using natural frequencies, which are generally easier to understand. Online calculators are available which allow clinicians to adjust pre-test probability, sensitivity, and specificity to estimate post-test probability.19

What else should clinicians consider when interpreting test results?

A single negative test result may not be informative if the pre-test probability is high

A 52 year old general practitioner in London develops a cough, intermittent fever, and malaise. On day 2 of his illness he receives a nasopharyngeal swab test for covid-19, which is reported as negative. His cough and low-grade fever persist but he feels systemically well enough to return to work. What should he do?

Pre-test probability is high in someone with typical symptoms of covid-19, an occupational risk of exposure, and working in a high prevalence region, and negative test results can therefore be misleading. Table 1 shows that for a pre-test probability of 90%, someone with a negative test has a 74% chance of having covid-19; with two negative tests this risk is still around 47%. If this doctor were to return to work and subsequently the test was confirmed as a false negative, then the decision to work would potentially have significant consequences for his patients, colleagues, and everyone with whom he came into contact. It is therefore safest for this GP with strongly suggestive symptoms to self-isolate in line with guidelines for covid-19, even though his test results are negative. This case illustrates the fallacy of base-rate neglect; it can be tempting to trust the results of an “objective” test more than one’s own “subjective” clinical judgement. In general, during this pandemic, pre-test probabilities of covid-19 will be high, particularly in high prevalence secondary care settings.

A possible alternative diagnosis will reduce the pre-test probability

A 73 year old woman with severe chronic obstructive pulmonary disease (COPD) and a chronic cough develops acute shortness of breath and slight worsening of her non-productive cough. She reports no fever, has no known exposure to covid-19, and no recent travel. She presents to an emergency department where she is acutely short of breath. A chest radiograph shows possible infiltrates in the right upper and middle lung fields. She is admitted and placed in isolation on droplet precautions. She requires intubation for worsening respiratory distress. Initial nasopharyngeal covid-19 testing is negative. Should she remain in isolation on droplet precautions?

This patient has an alternative possible diagnosis: community-acquired pneumonia. Given her lack of other risk factors or clinical symptoms, and chest radiography findings we therefore estimate her pre-test probability at about 50%. One negative test reduces this risk to 24%, the patient therefore has an additional independently sampled nasopharyngeal swab RNA
test which was negative, giving a post-test probability after two negative tests of less than 10%. She is treated with antibiotics and continues to recover.

What are the implications for practice and policy?

While positive tests for covid-19 are clinically useful, negative tests need to be interpreted with caution, taking into account the pre-test probability of disease. This has important implications for clinicians interpreting tests and policymakers designing diagnostic algorithms for covid-19. The Chinese handbook of covid-19 prevention and treatment states “if the nucleic acid test is negative at the beginning, samples should continue to be collected and tested on subsequent days.” False negatives carry substantial risks; patients may be moved into non-covid-19 wards leading to spread of hospital acquired covid-19 infection, carers could spread infection to vulnerable dependents, and healthcare workers risk spreading covid-19 to multiple vulnerable individuals. Clear evidence-based guidelines on repeat testing are needed, to reduce the risk of false negatives. Clinicians should ensure that patients are counselled about the limitations of tests (box 1). Patients with a single negative test but strongly suggestive symptoms of covid-19 should be advised to self-isolate in keeping with guidelines for suspected covid-19.

Box 1: Possible phrases for explaining covid-19 testing to patients

- No test is 100% accurate
- If your swab test comes back positive for covid-19 then we can be very confident that you do have covid-19
- However, people with covid-19 can be missed by these swab tests. If you have strong symptoms of covid-19, it is safer to self-isolate, even if the swab test does not show covid-19

What is the role of serology tests?

Serology tests, which detect immunoglobulins including IgG and IgM, are under development,22-24 with the aim of detecting individuals who have had previous infection and therefore theoretically developed immunity. The time course and accuracy of serology tests are still under investigation, but the same principles of incorporating the test result with the clinical impression applies. False positive serology tests could cause false reassurance, behaviour change, and disease spread. If suitable accuracy can be established, the benefits of these antibody tests include establishing when healthcare workers are immune, helping to inform decisions about the lifting of lockdowns, and allowing the population to return to work.25

The WHO message “test, test, test” is important from a population perspective; low sensitivity can be accounted for when assessing burden of disease. However RT-PCR tests have limitations when used to guide decision making for individual patients. Positive tests can be useful to “rule-in” covid-19, a negative swab test cannot be considered definitive for “ruling out.”

How patients were involved in the creation of this article

Patients with covid-19 or possible covid-19 were not involved in the writing of this paper for practical reasons.

Education into practice

What is the protocol for covid-19 testing in your organisation?

How do you explain covid-19 test results to patients?

Reflect on a recent clinical case of suspected covid-19—what was your estimated pre-test probability? How did this alter with the results of tests?

Author contributions: JW JB and PW contributed to the conception of the work, JW ran the searches and wrote the first draft of the paper with assistance from JB. PW developed the tools for fig 2. JB, JW, and PW all contributed to the revised drafts of the paper and approved the final version for submission.

Acknowledgements: The authors would like to acknowledge Jon Deeks for helpful discussions at an early point in writing this article and Richard Lehman for suggestions and comments on a draft of this article.

Competing interests The BMJ has judged that there are no disqualifying financial ties to commercial companies. The authors declare the following other interests: JB has given Grand Rounds talks on medical reasoning and has published a book.

Funding: JW is funded by a doctoral research fellowship from the National Institute for Health Research. The views expressed in this publication are those of the authors and not necessarily those of the NHS, the National Institute for Health Research, Health Education England, or the Department of Health.

Patient consent: The cases in this article are fictitious and therefore no consent was needed.

Provenance and peer review: Commissioned, based on an idea from the author; externally peer reviewed.

who-head-out-key-message-is-test-test-test


4 Day M. Covid-19: identifying and isolating asymptomatic people who have eliminated the virus. Italian village. BMJ 2020;368:m1165. 10.1136/bmj.m1165


11 Usher-Smith JA, Sharp SJ, Griffin SJ. The effect of tests in risk prediction, screening, and diagnosis. BMJ 2016;353:i3139. 10.1136/bmj.i3139


25 Petherick A. Developing antibody tests for SARS-CoV-2. Lancet 2020;395:1101-2. 10.1016/S0140-6736(20)30788-1 32247384

Published by the BMJ Publishing Group Limited. For permission to use (where not already granted under a licence) please go to http://group.bmj.com/group/rights-licensing/permissions
Table 1 | Pre- and post- test probabilities for covid-19 RT-PCR tests, calculations based on a sensitivity of 70% and specificity of 95%

<table>
<thead>
<tr>
<th>Pre-test probability</th>
<th>Post-test probability, negative test</th>
<th>Post-test probability, two independently negative tests</th>
<th>Post-test probability positive test</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>1.6</td>
<td>0.5</td>
<td>42</td>
</tr>
<tr>
<td>15</td>
<td>5</td>
<td>2</td>
<td>71</td>
</tr>
<tr>
<td>25</td>
<td>10</td>
<td>3</td>
<td>82</td>
</tr>
<tr>
<td>50</td>
<td>24</td>
<td>9</td>
<td>93</td>
</tr>
<tr>
<td>75</td>
<td>49</td>
<td>23</td>
<td>98</td>
</tr>
<tr>
<td>90</td>
<td>74</td>
<td>47</td>
<td>99</td>
</tr>
</tbody>
</table>
Figures

**Fig 1** Leaf plot for covid-19 RT-PCR tests based on a sensitivity of 70% and specificity of 95%. The x axis gives the estimated pre-test probability of covid-19 based on the clinical details. The post-test probability is obtained by tracing up and across to the y axis from the lower curve for a negative test, or to the upper curve for a positive test result. The dashed lines illustrate pre-test probability of 90% (clinical case 1) and 50% (clinical case 2).
Fig 2 Infographic showing outcomes of 100 people who are tested for covid-19