Horses for courses? Assessing the potential value of a surrogate, point-of-care test for SARS-CoV-2 epidemic control

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Abstract
Point-of-care tests (POCTs) offer considerable potential for improving clinical and public health management of COVID-19 by providing timely information to guide decision-making, but data on real-world performance are in short supply. Besides SARS-CoV-2-specific tests, there is growing interest in the role of surrogate (non-specific) tests such as FebriDx, a biochemical POCT which can be used to distinguish viral from bacterial infection in patients with influenza-like illnesses. This short report assesses what is currently known about FebriDx performance across settings and populations by comparison with some of the more intensively evaluated SARS-CoV-2-specific POCTs. While FebriDx shows some potential in supporting triage for early-stage infection in acute care settings, this is dependent on SARS-CoV-2 being the most likely cause for influenza-like illnesses, with reduction in discriminatory power when COVID-19 case numbers are low, and when co-circulating viral respiratory infections become more prevalent during the autumn and winter. Too little is currently known about its performance in primary care and the community to support use in these contexts, and further evaluation is needed. Reliable SARS CoV2-specific POCTs—when they become available—are likely to rapidly overtake surrogates as the preferred option given the greater specificity they provide.

KEYWORDS
antibody, COVID-19, diagnostic, molecular, rapid diagnostic test, SARS-CoV-2

1 | BACKGROUND

The COVID-19 pandemic presents health policymakers and clinicians with difficult decisions under conditions of uncertainty, facing an infection that in its early stages mimics many other influenza-like illnesses (ILIs). The range of diagnostic tests available for SARS-CoV-2 is growing, but they have important limitations. RT-PCR is the reference standard but depends on advanced laboratory facilities that are not available in primary care or community settings, and long turn-around times for results increase the risk of nosocomial transmission. Performance profiles for RT-PCR tests for SARS-CoV-2 are also sub-optimal, with reported false-negative rates of up to 30%.

Point-of-care tests (POCTs) could transform epidemic control by better guiding triage decisions to reduce risks of nosocomial and...
community transmission. However, reliable POCT options in the period before a detectable antibody response has been mounted are few. There could therefore be a role for surrogate tests (those that do not directly diagnose SARS-CoV-2) for screening. These include FebriDx, a low-cost POCT used to distinguish bacterial from viral infections and originally designed to support rational antibiotic prescribing. A number of evaluations are ongoing to consider the value of FebriDx in acute and primary care settings during the COVID-19 pandemic.

This short report explores the place of surrogate POCTs in the diagnostic mix in high-income settings, by considering the performance of FebriDx and SARS-CoV-2-specific POCTs on the market.

2 | POCT MODALITIES AND SARS-CoV-2 IN THEORY

There are four POC testing approaches for SARS-CoV-2 with different decision-making implications:

- **Class I:** POCTs for the presence of SARS-CoV-2 antigens to identify active infection.
- **Class II:** molecular POCTs to identify active infection but with shorter turn-around times than conventional RT-PCR.
- **Class III:** serological antibody tests (IgG and/or IgM) for confirmation of infection. This includes most SARS-CoV-2-specific POCTs.
- **Class IV:** surrogate tests (including FebriDx) that do not directly diagnose SARS-CoV-2 but can be used as screening tests.

Point-of-care test utility varies according to time from symptom onset. PCR-based tests offer better sensitivity in the first few days, but performance declines from as early as day 5. Class III POCTs likely offer value only from day 7 to 8 because of the time taken to generate a detectable antibody response. All POCTs are, for now, used to guide interim decision-making pending final diagnosis via RT-PCR.

While class I, II and III modalities test for SARS-CoV-2-specific antigens, RNA or antibodies respectively, FebriDx is a composite tool that detects both myxovirus resistance protein A (MxA—elevated in acute viral infections) and C-reactive protein (CRP—elevated in either viral or bacterial infections). The test uses fixed thresholds for "positive" MxA and CRP levels; any positive MxA result, with or without positive CRP, indicates viral infection. These markers are used because: (a) CRP production is stimulated by interleukin-6 (IL-6), produced in higher quantities in bacterial infections; (b) MxA expression is exclusively driven by type 1 interferons (IFN-1), secreted in response to detection of viral signatures by host intracellular receptors. IFN-1 and consequently MxA are thought to be specific to viral infection.

There may be grounds for caution in interpreting FebriDx results at the extremes of the clinical course. Early SARS-CoV infection appears not to activate MxA and IFN-1 transcription in the normal way, and if SARS-CoV-2 similarly attenuates early MxA expression, FebriDx may have low sensitivity in at least the first few days following infection. The low MxA threshold for a positive result (40 ng/mL) means FebriDx may nevertheless be positive even with weak MxA activation.

3 | COMPARATIVE PERFORMANCE OF SURROGATE AND SPECIFIC TESTS FOR SARS-CoV-2

The key risk in using any POCT is of a false-negative result leading to inappropriate management of SARS-CoV-2 infection. For FebriDx, an additional risk is that a positive result cannot exclude the possibility of another virus as the cause of infection.

Studies evaluating FebriDx mostly consider performance in distinguishing viral from bacterial causes of acute respiratory illnesses in toto, in secondary care. Reported sensitivities range from 64% to 90%, and specificities from 78% to 88%.

2-16 However, two recent studies from the UK evaluate FebriDx specifically for screening for SARS-CoV-2 in hospital: a small-scale pilot, and a study nested within a non-randomised clinical trial of molecular POCTs. These studies report impressive sensitivities and specificities of 100% and 93%, and 100% and 86%, respectively. However, interpretation is limited by (a) the use of clinical diagnosis as reference standard rather than RT-PCR in the first study, (b) evaluation in single secondary care centres in England in both cases and (c) the inclusion of patients in the range 2-7 days from symptom onset only in the first study. Finally, test performance in both studies may have been artificially boosted because they were conducted at times when the range of co-circulating respiratory viruses was lower than in the autumn and winter.

A comprehensive assessment of POCT field performance is beyond the scope of this paper, and available data indicate large context-dependent variations even for the same platform. However, based on data covering the first 14 days from symptom onset collated by FIND, SARS-CoV2-specific tests perform comparably to FebriDx (Figure 1). Caveats to this assessment are that: (a) the majority of tests are antibody-based and therefore only reactive some time after symptom onset, and (b) most studies used patient samples collected in clinical settings only. Evidence on the performance of FebriDx or any other POCT for diagnosis of SARS-CoV-2 infection in key workers, care home residents or other high-risk populations is in very short supply.

4 | IMPLICATIONS FOR APPLICATIONS OF FEBRIDX

What can be said about the utility of FebriDx by comparison with other POCTs given the limited available data? Preliminary judgements are possible depending on the (a) timing and location of testing, and (b) the broader epidemiological context.

FebriDx offers greatest value early in the clinical course, given the absence of reliable antigen POCTs and scarcity of molecular
POCTs. However, reported sensitivity ranges for COVID and non-
COVID patients in healthcare settings are wide. The test
should therefore be positioned within a diagnostic algorithm, along-
side other clinical and radiological markers, to optimise performance.
FebriDx may have a role in triage in the community where access to
advanced diagnostics is limited, but there are no performance data
for the test in this context, and only a handful of studies evaluate any
SARS-CoV-2 POCTs in the community.

Secondly, surrogate tests such as FebriDx will likely only have
value around the peak of the epidemic curve, in a situation where
widespread community transmission is probable. Here, a positive
result in a patient with an ILI could be interpreted as indicative of
COVID-19 pending laboratory confirmation. Discriminatory power
will likely decline during the autumn and winter months where
other viral diagnoses are equally or more likely. Here, the clinical
and social cost of imposing infection control measures such as iso-
lation on the basis of a presumptive diagnosis alone may become
unjustifiable.

5 | CONCLUSION

Although a growing number of specific and surrogate POCTs are now
available for SARS-CoV-2, test performance is variable across the clin-
ical course. Given existing data, its ease of use and low cost, FebriDx
shows promise as a screening tool for early-stage COVID-19 infection
in hospital settings. However, not enough is yet known about its value
in primary care or community settings. Reliable, SARS-CoV2-specific
tests—when they become available—are likely to be the preferred
option especially during the autumn and winter months when the in-
cidence of other, co-circulating respiratory viral infections will affect
the discriminating power of surrogate tests like FebriDx.

CONFLICT OF INTEREST

All authors declare that they have no conflicts of interest in relation
to this work.

AUTHOR CONTRIBUTIONS

Sharif A. Ismail: Conceptualization (equal); investigation (equal);
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FIGURE 1 Scatter plot of sensitivity
and specificity values for antigen- and
antibody-based POCTs, molecular rapid
diagnostic tests and scores for FebriDx
from the two studies that specifically
evaluate performance in screening for
COVID-19 published thus far. Bubble
sizes indicate the size of the population
involved in each study; the centre point
of each bubble gives the sensitivity
and specificity point estimate [Source:
FIND Dx open-source data on test
performance]
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