

BMJ Open Common evidence gaps in point-of-care diagnostic test evaluation: a review of horizon scan reports

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ABSTRACT

Objective Since 2008, the Oxford Diagnostic Horizon Scan Programme has been identifying and summarising evidence on new and emerging diagnostic technologies relevant to primary care. We used these reports to determine the sequence and timing of evidence for new point-of-care diagnostic tests and to identify common evidence gaps in this process.

Design Systematic overview of diagnostic horizon scan reports.

Primary outcome measures We obtained the primary studies referenced in each horizon scan report (n=40) and extracted details of the study size, clinical setting and design characteristics. In particular, we assessed whether each study evaluated test accuracy, test impact or cost-effectiveness. The evidence for each point-of-care test was mapped against the Horvath framework for diagnostic test evaluation.

Results We extracted data from 500 primary studies. Most diagnostic technologies underwent clinical performance (ie, ability to detect a clinical condition) assessment (71.2%), with very few progressing to comparative clinical effectiveness (10.0%) and a cost-effectiveness evaluation (8.6%), even in the more established and frequently reported clinical domains, such as cardiovascular disease. The median time to complete an evaluation cycle was 9 years (IQR 5.5–12.5 years). The sequence of evidence generation was typically haphazard and some diagnostic tests appear to be implemented in routine care without completing essential evaluation stages such as clinical effectiveness.

Conclusions Evidence generation for new point-of-care diagnostic tests is slow and tends to focus on accuracy, and overlooks other test attributes such as impact, implementation and cost-effectiveness. Evaluation of this dynamic cycle and feeding back data from clinical effectiveness to refine analytical and clinical performance are key to improve the efficiency of point-of-care diagnostic test development and impact on clinically relevant outcomes. While the 'road map' for the steps needed to generate evidence are reasonably well delineated, we provide evidence on the complexity, length and variability of the actual process that many diagnostic technologies undergo.

INTRODUCTION

Primary care is becoming increasingly complex due to a rise in patients with multimorbidity and polypharmacy, the pressure of short consultation times and the fragmented

Strengths and limitations of this study

- This study provides the first data on evidence gaps in point-of-care diagnostic test evaluation in primary care, answering an important clinical need.
- We extracted data from multiple consistently conducted horizon scan reports.
- Our approach might ignore relevant research, but systematic evidence gaps identified suggest findings to be robust.
- Our analyses are limited by the horizon scan report publication date.

nature of primary and secondary care. Delayed or missed diagnoses are the most common reason for malpractice claims.¹ Therefore, there is a huge demand for innovations that enable efficient and accurate diagnostic assessment within a general practitioner (GP)'s consultation. Consequently, the development of point-of-care diagnostic tests is currently a hotbed of activity.² These tests have the potential to significantly improve the efficiency of diagnostic pathways, providing test results within the time frame of a single consultation, enabling them to influence immediate patient management decisions.

A potential barrier to this innovative activity however is the slow and haphazard nature of the current pathway to adoption for new healthcare technologies.³ This is particularly the case for diagnostic tests, where uptake is highly variable between settings and notable inconsistencies lie in the speed at which they are adopted.⁴ One possible cause of this inefficiency is the slow generation of evidence of efficacy relevant to the target clinical settings.

To provide an efficient means of identifying, summarising and disseminating the evidence for emerging diagnostic technologies relevant to primary care settings, the Oxford Diagnostic Horizon Scanning Programme was established in 2008 (currently funded by the National Institute

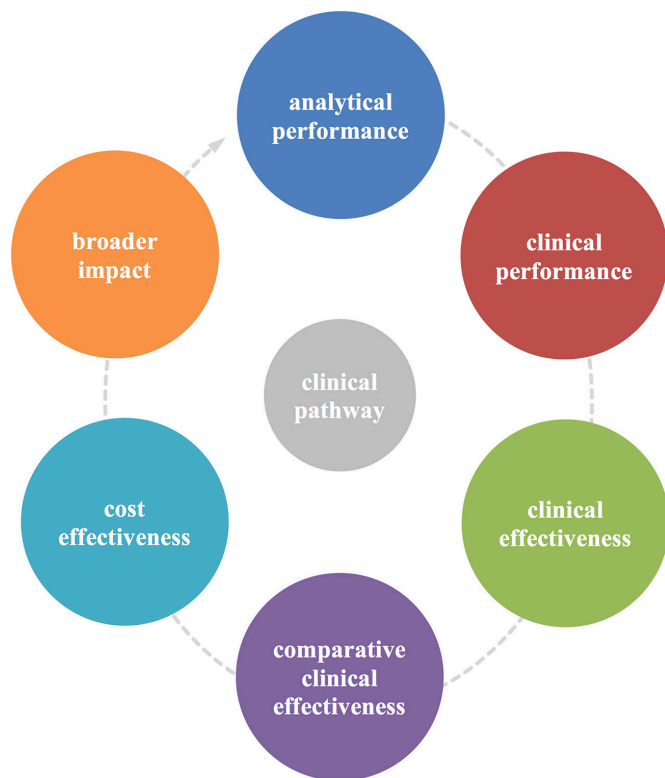


Figure 1 Horvath *et al*¹¹'s cyclical framework for the evaluation of diagnostic tests. This framework illustrates the key components of the test evaluation process. (1) Analytical performance is the aptitude of a diagnostic test to conform to predefined quality specifications. (2) Clinical performance examines the ability of the biomarker to conform to predefined clinical specifications in detecting patients with a certain clinical condition or in a physiological state. (3) Clinical effectiveness focuses on the test's ability to improve health outcomes that are relevant to an individual patient, also allowing comparison (4) of effectiveness between tests. (5) A cost-effectiveness analysis compares the changes in costs and health effects of introducing a test to assess the extent to which the test can be regarded as providing value for money. (6) Broader impact encompasses the consequences (eg, acceptability, social, psychological, legal, ethical, societal and organisational consequences) of testing beyond the above-mentioned components.

for Health Research (NIHR) Oxford Diagnostic Evidence Co-operative).⁵ New technologies are identified through systematic literature searches and interactions with clinicians and the diagnostics industry. These are then prioritised using a defined list of criteria.⁶ Evidence is gathered using systematic searches of the published literature and supplementary information obtained from manufacturer or trade websites and through web search engines, which are then used to summarise the analytical and diagnostic accuracy of the point-of-care test,⁷ impact of the test on patient outcomes and health processes, cost-effectiveness of the test and current guidelines for use within routine care in the UK. The reports, indexed in the TRIP database⁸ and freely available from the Horizon Scan Programme's website (www.oxford.dec.nihr.ac.uk), are actively disseminated to the NIHR Health Technology

Assessment Programme, the National Institute for Health and Clinical Excellence, clinical researchers and commissioners of healthcare services and highlight any further research requirements to facilitate evidence-based adoption decisions.

To date, 40 horizon scan reports have been completed, all following an identical protocol.

These horizon scan reports provide a unique opportunity to describe the evidence trajectory of new point-of-care diagnostic tests relevant to primary care settings and identify common evidence gaps.

METHODS

This is a descriptive study of all 40 horizon scan reports published to date by the Oxford Horizon Scan programme. For each horizon scan report, we extracted the year of publication and the disease area (classified per clinical domain of the International Classification of Primary Care—Revised Second Edition (ICPC2-R) 17) (see online supplementary file 1). We subsequently reviewed all studies that were included in the horizon scan reports (including systematic reviews) and extracted data on year of publication, size of the study, point-of-care test device(s) and its intended role. The intended roles were defined as 'triage', in which the new test is used at the start of the clinical pathway, 'replacement', in which the new test replaces an existing test, either as a faster equivalent test or to replace a non-point-of-care laboratory test, or 'add-on' in which the new test is performed at the end of a clinical pathway.⁹ Depending on the role, different types of evidence are required before a new point-of-care test can be adopted in routine care.¹⁰

We extracted data on study design and primary outcomes and used the dynamic evidence framework developed by Horvath *et al*,¹¹ as shown in [figure 1](#), to classify the type of evidence, defined as (1) analytical performance, (2) clinical performance, (3) clinical effectiveness, (4) comparative clinical effectiveness, (5) cost-effectiveness and (6) broader impact.

Analytical performance is the aptitude of a diagnostic test to conform to predefined quality specifications.^{12 13} Clinical performance examines the ability of the biomarker to conform to predefined clinical specifications in detecting patients with a particular clinical condition or in a physiological state.¹⁴ Clinical effectiveness focuses on the test's ability to improve health outcomes that are relevant to the individual patient.¹⁴ A cost-effectiveness analysis compares the changes in costs and in health effects of introducing a test to assess the extent to which the test can be regarded as providing value for money. Broader impact encompasses the consequences (eg, acceptability, social, psychological, legal, ethical, societal and organisational consequences) of testing beyond the above-mentioned components.

For point-of-care tests that had evidence on each of these components, we calculated the median time (in years) for a technology to complete the evaluation cycle.



We assessed whether the study was conducted in a setting that was relevant for primary care, which was defined as GP surgeries (clinics), outpatient clinics, walk-in (or urgent care) centres and emergency departments. Data extraction was piloted on 20 reports by BS and checked by JV and PT after which improvements were made to the final data extraction sheet. Three authors (JYV, BS and PJT) independently single-extracted data of the included studies.

RESULTS

We screened 40 horizon scan reports and extracted data from the 500 papers (including 41 systematic reviews) referenced by these reports (table 1). Ten horizon scan reports examined a point-of-care test relevant to cardiovascular disease, six to respiratory diseases and five to each of endocrine/metabolic diseases, digestive diseases and general/unspecified diseases. A further nine horizon scan reports examined a health problem relevant to a range of other disease areas. The intended role of the test was triage in 14 (35%), replacement in 20 (50%) and add-on in 6 (15%) of the 40 horizon scan reports.

We found a median of nine primary studies (IQR 7–15.8) per horizon scan report with a median time between first and last publication of 10 years (IQR 6.8–13.3). Across all horizon scan reports, on average, only 19% (95% CI 11.4% to 20.7%) of studies were performed in primary care (figure 2).

Of all studies, 30.4% (n=152) assessed analytical performance of the diagnostic technology, providing evidence for this component in 25 (62.5%) of the 40 horizon scan reports.

Clinical performance was evaluated in 71.2% (n=356) of all studies, while only 18.2% (n=91) of studies evaluated clinical effectiveness of the diagnostic technology. A further 10.0% (n=50) compared clinical effectiveness of two or more point-of-care tests, and only 8.6% (n=43) of the 500 papers evaluated cost-effectiveness (figure 3).

Clinical performance was often assessed earlier (in 16 tests) or together (in 12) with analytical performance and not assessed at all for 6 tests. Broader impact such as acceptability was tested before evidence on clinical effectiveness or cost-effectiveness was available in 11 horizon scan reports.

Figure 4 shows the number of years between the horizon scan report and original paper publication date for each evaluation component, split by intended role of the point-of-care test. The size of the bubbles represents the number of studies proportionate to all studies for the intended role, clearly depicting the emphasis on clinical performance and paucity of clinical effectiveness studies. Furthermore, tests acting as a triage instrument tend to spend more time on evidence generation than tests replacing an existing one or add-on tests performed at the end of the clinical pathway.

Only seven (17.5% (95% CI 7.3% to 32.8%)) horizon scan reports included evidence for all evaluation components with a median time to complete the evaluation cycle of 9 years (IQR 6–13 years). Of these, tests acting as a triage

instrument (in three reports) had a median of 15 years (IQR 10–19) while tests replacing an existing one (in four reports) had 9 years (IQR 5–11) (figure 4).

Even in the latter category of diagnostic technology replacing existing tests, where nearly half of studies (49.4%; n=247) have been performed, there was a clear imbalance between studies merely focusing on analytical or clinical performance (87.4%) and the few studies advancing to clinical effectiveness (21.1%).

The sequence of evidence generation over time for the seven horizon scan reports which had completed the evaluation cycle varied widely, as shown in figure 5. The size of the bubbles represents the proportion of studies for each evaluation component. The grey arrow shows the sequence we would expect, starting at analytical performance (at 12 o'clock) and completing at broader impact analysis (at 10 o'clock). The arrows and numbered bubbles represent the actual time sequence of evidence generation.

Very few point-of-care test evaluations seem to follow the expected sequence. In fact, only the report on point-of-care C reactive protein testing seemed to generally follow a linear temporal sequence from analytical performance towards broader impact. Some diagnostic technology, such as point-of-care international normalised ratio (INR) testing, had evidence generated for the broader impact component before any other component, suggesting that some diagnostic technologies are adopted in routine clinical care prior to any published evidence on clinical performance or effectiveness.

DISCUSSION

Main findings

Our findings suggest that most point-of-care diagnostic tests undergo clinical performance assessment, but very few progress to evaluation of their broader impact or cost-effectiveness, even in the more established and frequently reported clinical domains, such as cardiovascular disease. Some point-of-care tests even skip essential stages such as clinical effectiveness, yet are still implemented in routine care. We present a novel way to visualise the gaps in the evidence generation through bubble plots and dynamic cycle illustration.

Strengths and limitations

Our study provides novel data on common evidence gaps in the evaluation of novel point-of-care tests for a wide range of clinical conditions. The extensive library of existing horizon scan reports and the methodological rigour in which they are produced provided an ideal opportunity to review the pathway of evidence for novel point-of-care diagnostic technologies relevant to the primary care settings. The chosen topics of the reports result from a comprehensive approach to identify new or emerging diagnostic tests, including literature searches and interaction with the diagnostics industry and clinicians, prioritising technologies relevant to primary care. We have, however, no measure of how

**Table 1** Baseline characteristics of horizon scan reports by clinical domain

Diagnostic tests by clinical domain	Date report published	Studies in primary care (n)/all studies (N)	Sample sizes of primary studies (range)	Publication dates primary studies (range)	Intended role of the test	Components missing
Cardiovascular						
The D-dimer test for ruling out DVT in primary care	2009	2/9	1028–1295	1997–2009	T	I/IV/VI
Point-of-care test for INR coagulometers	2010	16/32	18–892	2000–2010	R	–
Point-of-care test for cardiac troponin	2011	0/18	20–2263	2001–2011	R	–
Point-of-care pulse wave analysis	2011	0/7	22–749	2008–2011	R	III/IV/VI
Point-of-care test for BNP	2011	1/15	150–606	2001–2011	T	IV
Handheld ECG monitors for the detection of atrial fibrillation in primary care	2013	0/9	18–605	1990–2012	R	I/IV/V
Genotyping polymorphisms of warfarin metabolism	2013	0/2	20–112	2009–2010	T	III/IV/VI
Point-of-care test for hFABP	2014	1/21	52–1074	2003–2013	A	IVI
Portable ultrasound devices	2014	1/24	3–943	1993–2013	T	VI
Point-of-care test for a panel of cardiac markers	2014	0/21	33–5201	1999–2012	T	I/III/VI
Digestive						
Point-of-care test for hepatitis C virus	2011	0/6	100–2206	1999–2011	R	IV/VI
Point-of-care test for coeliac disease	2012	1/9	87–2690	2004–2009	T	I/III/IV/V
Transcutaneous bilirubin measurement	2013	0/28	31–849	1996–2011	T	I/III/IV
Point-of-care calprotectin tests	2014	1/8	14–140	2008–2012	T	I/III/IV/VI
Point-of-care faecal occult blood testing	2014	10/30	100–85149	1990–2013	T	–
Endocrine/metabolic						
Point-of-care blood test for ketones in diabetes	2009	6/9	33–629	2000–2006	T	–
Point-of-care test for the analysis of lipid panels	2010	9/13	34–4968	1995–2010	R	–
Point-of-care test for HbA1c	2010	6/7	23–7893	2003–2010	R	IV*
Point-of-care test for thyroid-stimulating hormone	2013	0/1	64–64	1999–1999	R	I/III/IV/VI
Point-of-care HbA1c tests: diagnosis of diabetes	2016	7/46	23–6226	1996–2015	R	III/IV*
Female genital						
<i>Chlamydia trachomatis</i> testing	2009	2/8	162–2517	1998–2009	R	II/VI
General and unspecified						
Point-of-care test for total white cell count	2010	1/6	120–500	2000–2009	R	III/IV/VI
Point-of-care test for CRP	2011	6/8	20–898	1997–2011	R	–
Point-of-care test for procalcitonin	2012	0/8	54–384	2001–2010	A	IV
Non-contact infrared thermometers	2013	0/6	90–855	2005–2013	A	III/VI
Point-of-care test for malaria	2015	0/6	98–98	1999–2014	R	IV
Musculoskeletal						
Autoimmune markers for rheumatoid arthritis	2012	0/1	880–880	2008–2008	R	I/III/IV/VI
Neurological						
Point-of-care test for handheld nerve conduction measurement devices for carpal tunnel syndrome	2012	1/9	33–1190	2000–2011	R	IV/V

Continued

Table 1 Continued

Diagnostic tests by clinical domain	Date report published	Studies in primary care (n)/all studies (N)	Sample sizes of primary studies (range)	Publication dates primary studies (range)	Intended role of the test	Components missing
<i>Pregnancy</i>						
Urinalysis self-testing in pregnancy	2014	2/9	49–4 44 220	1991–2012	A	IV
Point-of-care test for quantitative blood hCG	2015	0/2	40–40	2014–2015	T	III/IV/V/VI
<i>Respiratory</i>						
Pulse oximetry in primary care	2009	1/4	114–2127	1997–2004	T	I/III/IV/V
A portable handheld electronic nose in the diagnosis of cancer, asthma and infection	2009	5/8	30–665	1999–2009	T	III/IV/V/VI
Point-of-care spirometry	2011	4/11	13–1041	1996–2009	R	V
Point-of-care automated lung sound analysis	2011	3/22	1–100	1996–2010	A	IV/V
Point-of-care tests for influenza in children	2012	8/29	73–9186	2002–2011	R	I/IV
Point-of-care tests for group A streptococcus	2015	4/10	121–892	2002–2015	R	I
<i>Skin</i>						
Dermoscopy for the diagnosis of melanoma in primary care	2009	1/7	96–3053	2001–2008	T	I/IV/V
<i>Urological</i>						
Point-of-care urine albumin–creatinine ratio test	2010	6/11	83–4968	1999–2010	R	–
Point-of-care test for creatinine	2014	1/10	20–401	2007–2013	R	IV/V/VI
Point-of-care NGAL tests	2015	0/10	100–1219	2007–2013	A	III/IV/V
Grand total		106/500				

Evaluation components: I, analytical performance; II, clinical performance; III, clinical effectiveness; IV, comparative clinical effectiveness; V, cost-effectiveness; VI, broader impact.

*Same diagnostic technology.

A, add-on; BNP, B-natriuretic peptide; CRP, C-reactive protein; DVT, deep vein thrombosis; HbA1c, glycated haemoglobin; hCG, human chorionic gonadotropin; hFABP, heart-type fatty acid-binding protein; INR, international normalised ratio; n, number; NGAL, neutrophil gelatinase-associated lipocalin; R, replacement; T, triage

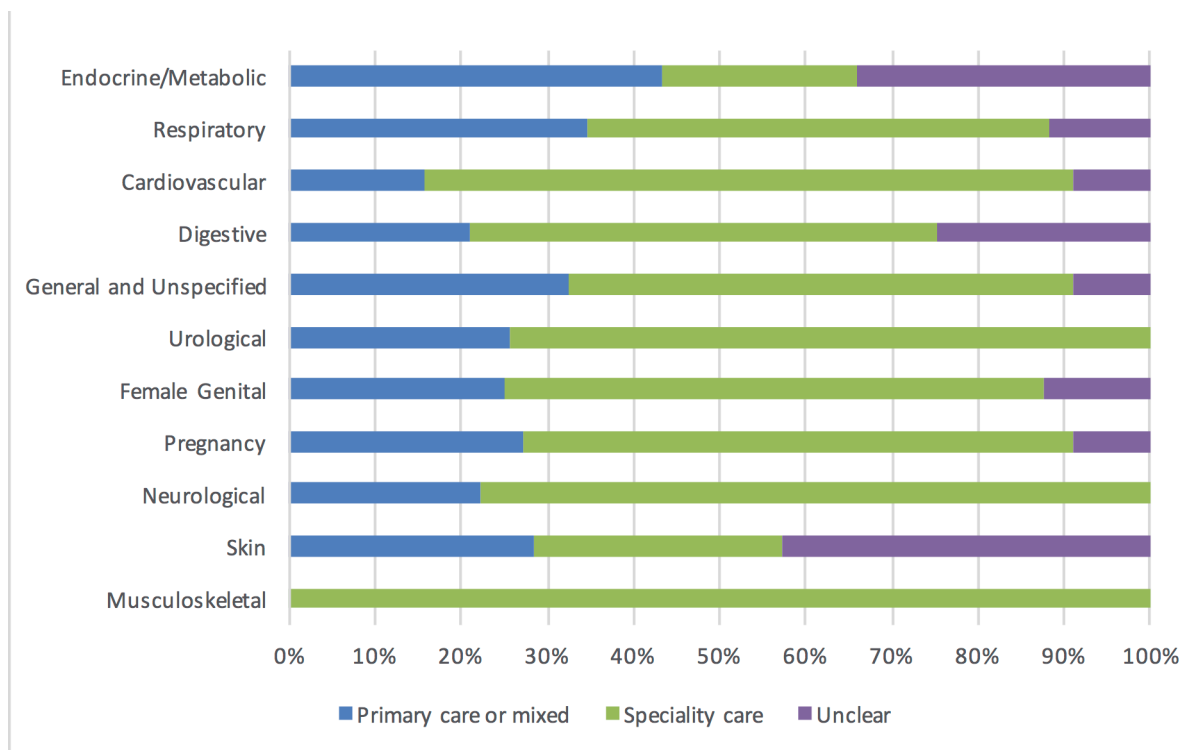


Figure 2 Setting (%) of the studies by disease area (according to the International Classification of Primary Care-Second edition coding).

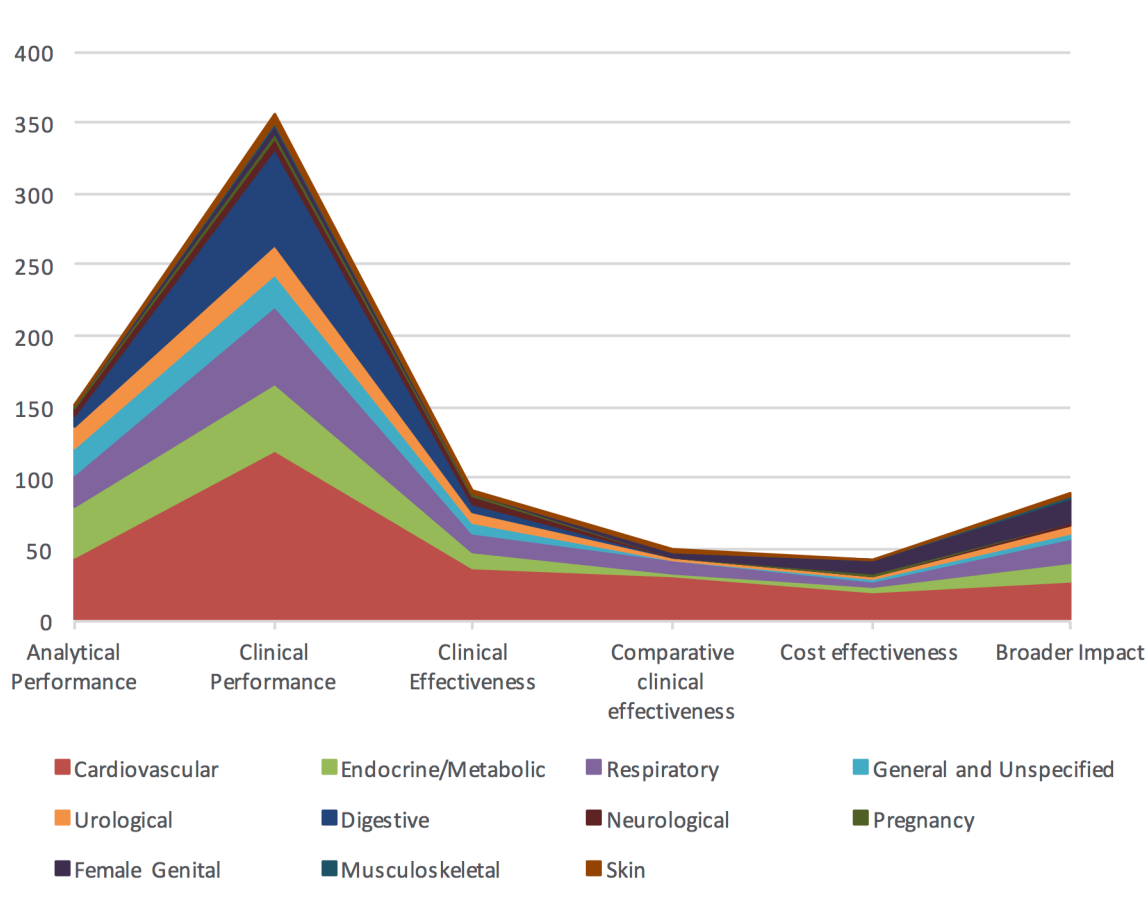


Figure 3 Test evaluation component by disease area in absolute number (n) of studies.



Figure 4 Number of years between horizon scan report and original paper publication date by the intended role for each evaluation component. Size of bubbles represents number of studies proportionate to all studies for the intended role. BNP, B-natriuretic peptide; CRP, C reactive protein; FOBT, faecal occult blood test; HbA1c, glycated haemoglobin; hCG, human chorionic gonadotropin; hFABP, heart-type fatty acid-binding protein; INR, international normalised ratio; TSH, thyroid-stimulating hormone; WBC, white cell count.

reproducible this prioritisation process is, and potentially risks greater or lesser inclusion of various disease areas. Evidence from reports on other clinical topics might provide different findings. Further to this, our review is limited to publication date of each report, thus potentially overlooking evidence generated following publication. Our approach might arguably ignore relevant (unpublished) research carried out (eg, studies performed during test development by industry), but the commonalities in the evidence gaps across the reports suggest that our findings are robust.

Three authors (JYV, BS and PJT) independently single-extracted data of the 500 included studies, making it impossible to test for inter-rater agreement.

Comparison with the existing literature

Previous evidence has shown that the adoption of a diagnostic technology is often insufficient to achieve a benefit, and in most cases, a change of care process is essential.¹⁵ The market for point-of-care tests is growing rapidly,¹⁶ and there is a clear demand from primary care clinicians for these tests to help them diagnose a

range of conditions.^{17 18} Critical appraisal of new diagnostic technologies is considered essential to facilitate implementation.¹¹ Several evaluation frameworks have been identified,¹⁹ most of which describe the evaluation process as a linear process, similar to the staged evaluation of drugs.

Considering the interactions between different evaluation components and the need for certain tests to re-enter the evaluation process after updates to the underlying technology,²⁰ it may be more realistic to assess these components in a cyclic and repetitive process.¹¹

Implications

The slow adoption of novel point-of-care tests may result from the paucity of technologies following the expected sequence of evidence generation. Specifically, there is a need to shift emphasis from examining clinical performance of point-of-care tests to comparative clinical effectiveness and broader impact assessment. We recommend using a structured dynamic approach, presenting the results in a visually appealing manner for both industry (during development and pursuit for regulatory

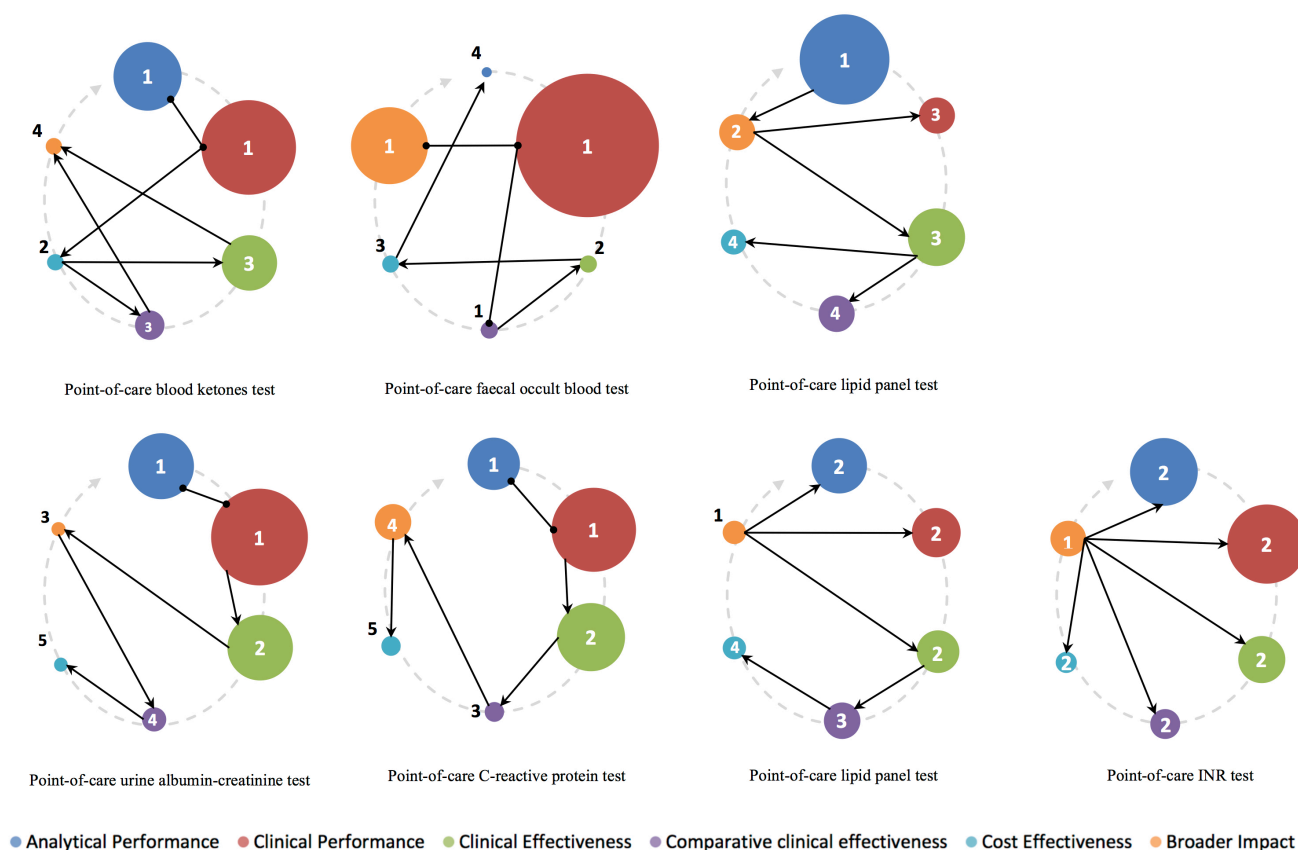


Figure 5 Sequence of evidence generation for all seven horizon scan reports completing the full evaluation cycle. INR, international normalised ratio.

approval) and research purposes. Policy-makers and guideline developers should be aware of this cyclical nature; assuming test evaluation is a linear process results in a less efficient evidence generation pathway. For example, assessing the cost-effectiveness early on in the development phase of novel point-of-care test can help determine where exactly it fits in the clinical pathway and thus ensure that the evidence subsequently generated is relevant to that population and setting.

CONCLUSIONS

Considering that evidence generation for new tests takes on an average 9 years, test developers need to be aware of the time and investment required. While the 'road map' for the steps needed to generate evidence are reasonably well delineated, we provide evidence on the complexity, length and variability of the actual process that many diagnostic technologies undergo.

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Contributors BS, MJT and AVdB conceived the study. JYV, PJT and BS did data extraction. JYV designed and performed the analyses, which were discussed with PJT, MJT, AP, CPP, BS and AVdB. JYV drafted this report and JYV, PJT, MJT, AP, CPP, BS and AVdB codrafted and commented on the final version. All authors had full access to all of the data (including statistical reports and tables) in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

JYV affirms that the manuscript is an honest, accurate and transparent account of the study being reported; that no important aspects of the study have been omitted. All authors read and approved the final manuscript.

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REFERENCES

- Wallace E, Lowry J, Smith SM, *et al*. The epidemiology of malpractice claims in primary care: a systematic review. *BMJ Open* 2013;3:e002929.
- GV ResearchIVD. *Market Analysis By Product (Tissue Diagnostics, Molecular Diagnostics, Professional Diagnostics, Diabetes Monitoring) And Segment Forecasts To 2020*. 100. Pune, India: Grand View Research, 2016.



3. Greenhalgh T, Robert G, Bate P, *et al.* *How to spread good ideas: a systematic review of the literature on diffusion, dissemination and sustainability of innovations in health service delivery and organisation.* London: NCCSDO, 2004.
4. Britain G, National Health S. *The NHS atlas of variation in diagnostic services: reducing unwarranted variation to increase value and improve quality.* Great Britain: Right Care, 2013.
5. Oxford DEC. NIHR Diagnostic evidence Cooperative Oxford. 2016 <https://www.oxford.dec.nihr.ac.uk/>
6. Plüddemann A, Heneghan C, Thompson M, *et al.* Prioritisation criteria for the selection of new diagnostic technologies for evaluation. *BMC Health Serv Res* 2010;10:109.
7. Mallett S, Halligan S, Thompson M, *et al.* Interpreting diagnostic accuracy studies for patient care. *BMJ* 2012;345:e3999.
8. Brassey J. University of Wales CoMCFRS. *TRIP database.* Cardiff: University of Wales, College of Medicine, Centre for Research Support, 2001.
9. Bossuyt PM, Irwig L, Craig J, *et al.* Comparative accuracy: assessing new tests against existing diagnostic pathways. *BMJ* 2006;332:1089–92.
10. Van den Bruel A, Cleemput I, Aertgeerts B, *et al.* The evaluation of diagnostic tests: evidence on technical and diagnostic accuracy, impact on patient outcome and cost-effectiveness is needed. *J Clin Epidemiol* 2007;60:1116–22.
11. Horvath AR, Lord SJ, StJohn A, *et al.* From biomarkers to medical tests: the changing landscape of test evaluation. *Clin Chim Acta* 2014;427:49–57.
12. Fraser CG, Kallner A, Kenny D, *et al.* Introduction: strategies to set global quality specifications in laboratory medicine. *Scand J Clin Lab Invest* 1999;59:477–8.
13. Singh S, Chang SM, Matchar DB, *et al.* Chapter 7: grading a body of evidence on diagnostic tests. *J Gen Intern Med* 2012;27(Suppl 1):47–55.
14. Global Harmonization Task Force. *Clinical Evidence for IVD medical devices – Key Definitions and Concepts:* Global Harmonization Task Force, 2012.
15. St John A. The Evidence to Support Point-of-Care Testing. *Clin Biochem Rev* 2010;31:111–9.
16. Stone Hearth News. Point of care test (POCT) market will see faster revenue growth in 2013 than tests sent to central labs for processing. 2012 <http://www.stonehearthnewsletters.com/point-of-care-test-poct-market-will-see-faster-revenue-growth-in-2013-than-tests-sent-to-central-labs-for-processing/health-care/>
17. Howick J, Cals JW, Jones C, *et al.* Current and future use of point-of-care tests in primary care: an international survey in Australia, Belgium, The Netherlands, the UK and the USA. *BMJ Open* 2014;4:e005611.
18. Turner PJ, Van den Bruel A, Jones CH, *et al.* Point-of-care testing in UK primary care: a survey to establish clinical needs. *Fam Pract* 2016;33:388–94.
19. Lijmer JG, Leeflang M, Bossuyt PM. Proposals for a phased evaluation of medical tests. *Med Decis Making* 2009;29:E13–E21.
20. Thompson M, Weigl B, Fitzpatrick A, *et al.* More Than Just Accuracy: A Novel Method to Incorporate Multiple Test Attributes in Evaluating Diagnostic Tests Including Point of Care Tests. *IEEE J Transl Eng Health Med* 2016;4:2800208:1–8.

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