



Accuracy of line probe assays for the diagnosis of pulmonary and multidrug-resistant tuberculosis: a systematic review and meta-analysis

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Line probe assays have high accuracy for detection of RIF resistance and INH resistance
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ABSTRACT Only 25% of multidrug-resistant tuberculosis (MDR-TB) cases are currently diagnosed. Line probe assays (LPAs) enable rapid drug-susceptibility testing for rifampicin (RIF) and isoniazid (INH) resistance and *Mycobacterium tuberculosis* detection. Genotype MTBDRplusV1 was WHO-endorsed in 2008 but newer LPAs have since been developed.

This systematic review evaluated three LPAs: Hain Genotype MTBDRplusV1, MTBDRplusV2 and Nipro NTM+MDRTB. Study quality was assessed with QUADAS-2. Bivariate random-effects meta-analyses were performed for direct and indirect testing. Results for RIF and INH resistance were compared to phenotypic and composite (incorporating sequencing) reference standards. *M. tuberculosis* detection results were compared to culture.

74 unique studies were included. For RIF resistance (21225 samples), pooled sensitivity and specificity (with 95% confidence intervals) were 96.7% (95.6–97.5%) and 98.8% (98.2–99.2%). For INH resistance (20954 samples), pooled sensitivity and specificity were 90.2% (88.2–91.9%) and 99.2% (98.7–99.5%). Results were similar for direct and indirect testing and across LPAs. Using a composite reference standard, specificity increased marginally. For *M. tuberculosis* detection (3451 samples), pooled sensitivity was 94% (89.4–99.4%) for smear-positive specimens and 44% (20.2–71.7%) for smear-negative specimens.

In patients with pulmonary TB, LPAs have high sensitivity and specificity for RIF resistance and high specificity and good sensitivity for INH resistance. This meta-analysis provides evidence for policy and practice.

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Introduction

Tuberculosis causes 10.4 million cases and 1.8 million deaths annually and it is estimated that 4.3 million cases go undiagnosed each year [1]. The emergence of multidrug and extensively drug-resistant tuberculosis (MDR-TB and XDR-TB, respectively) is a major threat to global tuberculosis control [2]. Culture and drug-susceptibility testing (DST) using solid media can take up to 8–12 weeks for results [3] and faster liquid-based culture techniques still take 4–6 weeks [4]. The delays associated with DST lead to prolonged periods of ineffective therapy and ongoing tuberculosis transmission. The development of rapid molecular diagnostic tests for the identification of *Mycobacterium tuberculosis* and drug resistance has consequently become a research and implementation priority [5].

Line probe assays (LPAs) are rapid molecular diagnostics that can detect *M. tuberculosis* and drug resistance. Although LPAs are more technically complex (designed for reference or regional laboratory settings) and take longer to perform than the Xpert MTB/RIF assay (Cepheid, Sunnyvale, CA, USA), they have the ability to detect isoniazid (INH) resistance in addition to rifampicin (RIF) resistance unlike Xpert MTB/RIF [6]. LPAs detect RIF and INH resistance by identifying mutations in the *rpoB*, *katG*, and *inhA* genes. By targeting mutations in the 81-base pair “core region” of the *rpoB* gene, more than 95% of all RIF resistant strains can be detected [7]. In comparison, the mutations that cause INH resistance are located in several genes and regions [8, 9]. Although mutations in *katG* and *inhA* account for approximately 80–90% of INH-resistant strains [10], an additional 5–10% of INH-resistant strains have mutations in the *ahpC-ohyR* intergenic region, often in conjunction with *katG* mutations outside of codon 315 [11].

The World Health Organization (WHO) approved LPAs for the diagnosis of *M. tuberculosis* and RIF resistance in smear-positive tuberculosis in 2008 [12], guided by a systematic review evaluating two first-generation LPAs: INNO-LiPA Rif.TB assay (Innogenetics, Ghent, Belgium) and GenoType MTBDR assay (Hain Lifescience GmbH, Nehren, Germany) [13], both of which assays are no longer used in clinical practice. Newer versions of the LPA technology have been developed [14–17] and additional studies have been published. This systematic review was commissioned by the WHO to guide a policy update on the use of molecular diagnostics. We evaluated the diagnostic accuracy of three LPAs (appendix A in the supplementary material): GenoType MTBDRplus V1 (subsequently referred to as “Hain V1”), GenoType MTBDRplus V2 (subsequently referred to as “Hain V2”) and Nipro NTM+MDRTB Detection Kit 2 (subsequently referred to as “Nipro”), for the detection of RIF and INH resistance and detection of *M. tuberculosis*.

Methods

We followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines and methods for systematic reviews and meta-analyses of diagnostic test accuracy [18, 19]. We prepared a protocol for the literature search, article selection, data extraction, assessment of methodological quality and synthesis of results.

Search methods

We performed a comprehensive search of the following databases (PubMed, EMBASE, BIOSIS, Web of Science, LILACS, Cochrane) for relevant citations (full search strategy reported in appendix C in the supplementary material). Our search was restricted to the time period January 2004 to August 2015, since the first Hain LPA was introduced in October 2004. In addition, we contacted laboratory experts and the test manufacturers for additional published studies. We also searched reference lists from included studies and previous meta-analyses [13]. No language restriction was initially applied but at the full-text review stage we restricted studies to English, French and Spanish. Abstracts or conference proceedings were not included.

Study selection and data extraction

Two review authors (R.R. Nathavitharana and P.G.T. Cudahy) independently assessed titles and abstracts (screen 1). Any citation identified by either review author during screen 1 was selected for full-text review. The same two review authors (R.R. Nathavitharana and P.G.T. Cudahy) independently assessed the full-text articles for inclusion (screen 2). In screen 2, any discrepancies were resolved by discussion between the review authors or by arbitration by a third review author (C.M. Denking). Two review authors (R.R. Nathavitharana and P.G.T. Cudahy) extracted data from the included studies with a pre-piloted standardised form and crosschecked to ensure accuracy. Disagreement between review authors on data extraction was resolved by discussion or by a third reviewer (C.M. Denking). Studies without extractable sensitivity and specificity data were excluded if no further information was acquired after three attempts to contact the study authors.

Selection criteria

We included cross-sectional, case-control, cohort studies or randomised controlled trials comparing LPAs to a reference standard test (see below), if at least 25 samples were tested. Patients of all age groups with

suspected or confirmed pulmonary tuberculosis or MDR-TB were included, regardless of setting or country. Specimen types were limited to sputum. Patients who were already on therapy were excluded from analyses of *M. tuberculosis* detection (since dead bacilli not detected by culture could be detected by LPAs leading to false positive results) but were included in analyses for RIF and INH resistance detection.

The reference standard test for the detection of *M. tuberculosis* was a positive solid or liquid culture for *M. tuberculosis*. The reference standard test for the detection of RIF and INH resistance detection was phenotypic DST for our primary analysis for all studies. Where data were available, LPA results were also compared with a composite reference standard, which combined the results from targeted genetic sequencing and phenotypic DST results (see appendix B in the supplementary material for details).

Outcome measures

Our outcome measures for all questions were sensitivity and specificity. Indeterminate results were excluded from the analyses for determination of sensitivity and specificity and were reported separately (further details in appendix B in the supplementary material).

Assessment of methodological quality

We used the Quality Assessment of Studies of Diagnostic Accuracy included in Systematic Reviews-2 (QUADAS-2) instrument, a validated tool for diagnostic studies, to assess study quality [20]. The information needed to answer QUADAS-2 questions was incorporated in the data extraction sheet. A description of the QUADAS-2 items and the interpretation in the study context can be found in appendices D1 and D2 in the supplementary material.

Statistical analysis and data synthesis

We performed statistical analyses using STATA (version 13; STATA corporation, College Station, TX, USA). The studies were grouped by type of index test and reference standard used. Our QUADAS-2 analysis was performed using Excel (version 14.5.4; Microsoft, Seattle, WA, USA).

Meta-analysis

Meta-analysis was performed for each index test if at least four studies were available for the same index test and if there was limited heterogeneity between studies. Bivariate random effects meta-analyses were performed [21, 22] using the metandi package in STATA for index tests that included enough data to calculate sensitivity and specificity, with 95% confidence intervals. Summary and individual estimates were also presented graphically with the 95% confidence intervals and prediction region. Several studies did not contribute to both sensitivity and specificity but only to one of the two. In order to make complete use of the data for these studies we performed a univariate random effects meta-analysis of the sensitivity and/or specificity estimates separately. Where there were fewer than four studies available or if substantial heterogeneity precluded meta-analysis, a descriptive analysis was performed. Forest plots were visually assessed for heterogeneity among the studies within each index test. Using summary plots, we examined the variability in estimates and the width of the prediction region, with a wider prediction region suggesting more heterogeneity. We anticipated that studies included in the meta-analysis would be fairly heterogeneous and thus sub-groups for analysis were pre-specified as LPA type, specimen type, specimen conditions and smear status.

Results

Characteristics of included studies

From the literature search, we identified 1650 citations and reviewed 218 full-text articles. 74 studies were included in this systematic review (figure 1) [15, 17, 23–94]. 16 of these studies contributed data to more than one analysis, resulting in a total of 94 datasets. A list of excluded studies and the reasons for exclusion is presented in appendix E in the supplementary material. Tables 1 and 2 demonstrate the characteristics of the 94 datasets that provided data on RIF and INH (of note, four of these datasets only provided data on RIF but not INH) and the six datasets that provided data on *M. tuberculosis* detection respectively. The majority of datasets were cross-sectional in design and almost all were performed in either a regional or national reference laboratory setting. 48 datasets evaluated LPA for direct testing on sputum specimens [15, 17, 24, 25, 27, 29–32, 35–39, 41–43, 46, 47, 49, 50, 52, 57, 59, 62–69, 71, 72, 74, 77–82, 84, 87, 91, 93] and 46 datasets evaluated LPA for indirect testing on culture isolates [17, 23, 25–28, 33–35, 37, 39, 40, 44, 45, 48, 51–58, 60–62, 68, 70, 72, 73, 75, 76, 78, 79, 83, 85, 86, 88–90, 92, 94]. 83 datasets evaluated Hain V1, five datasets evaluated Hain V2 [15, 30, 36, 72] and six datasets evaluated Nipro [17, 72, 79]. Very few datasets recorded demographic data or HIV status due to the use of anonymised samples.

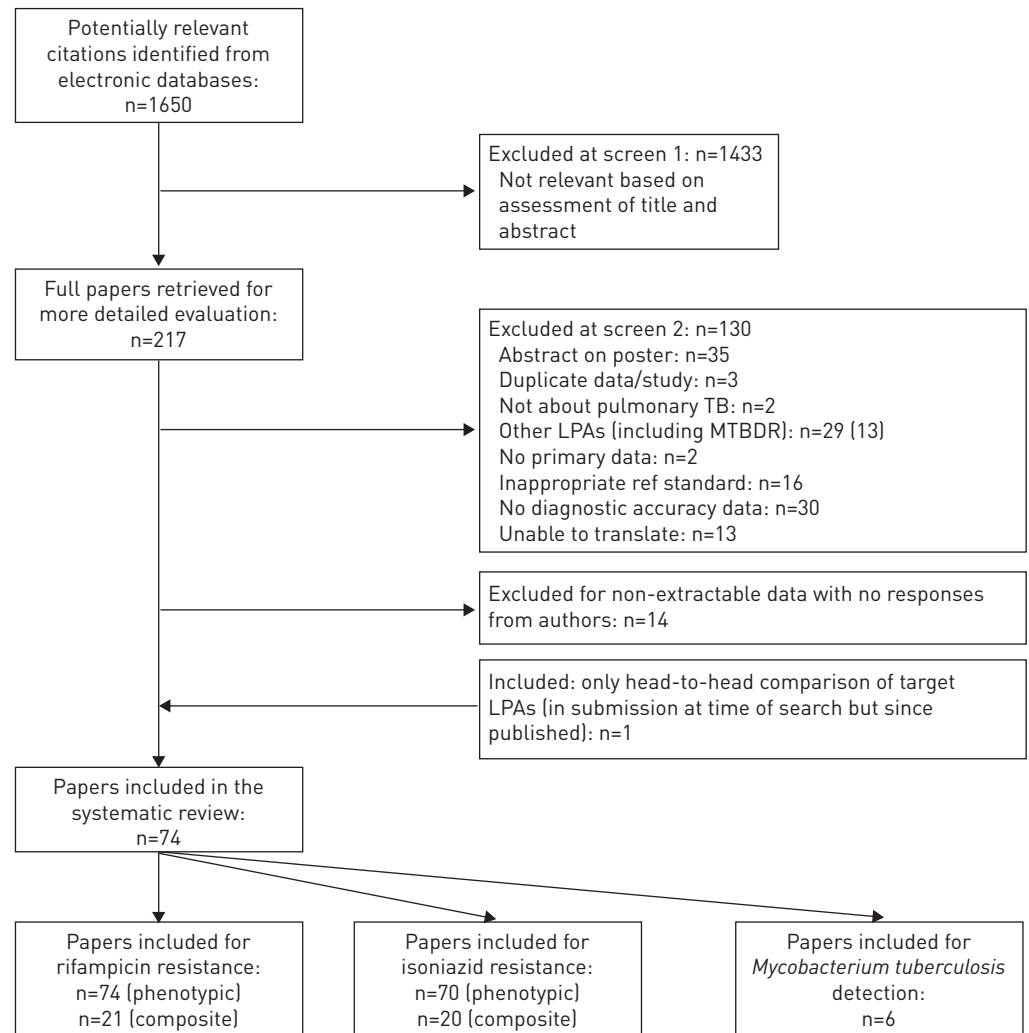


FIGURE 1 Preferred Reporting Items for Systematic Reviews and Meta-Analyses diagram of included studies. TB: tuberculosis; LPA: line probe assay.

Methodological quality

The methodological quality across all included studies is summarised in figure 2 and presented for each individual study in appendix D3 in the supplementary material. Many studies did not report all factors that could affect methodological quality. For the “patient selection” domain, there was unclear risk of bias for 56 out of 94 datasets for RIF and INH resistance and five out of six datasets for *M. tuberculosis* detection predominantly because the method of sampling of patients was not defined. Applicability concerns were unclear in 18 out of 94 datasets for RIF and INH resistance and one out of six datasets for *M. tuberculosis* detection that did not specify the type of patients tested or laboratory setting. For the “index test” domain, there was unclear risk of bias for 66 out of 94 datasets for RIF and INH resistance and two out of six datasets for *M. tuberculosis* detection because it was not stated whether the person performing the index test was blinded to the results of the reference standard testing. Applicability concerns in this domain were high risk in eight out of 94 datasets that reported variations in test processing that were not according to the manufacturer’s recommendations. For the “reference test” domain, there was unclear risk of bias for many datasets (68 out of 94) for RIF and INH resistance and three out of six datasets for *M. tuberculosis* detection because it was not stated whether the person performing the reference test was blinded to the results of the index tests. Applicability concerns were low. In the “flow and timing” domain, the majority of datasets (78 out of 94 and six out of six, respectively) were judged to have a low risk of bias.

Indeterminate result and culture contamination rates

30 datasets reported indeterminate results for directly tested specimens with a median of 5.3% and range of 1.0–14.5% for rifampicin and 5.6% and 0.9–14.5% for isoniazid (appendix F, table S1 in the

TABLE 1 Characteristics of included studies for rifampicin (RIF) and isoniazid (INH) resistance detection, grouped alphabetically by index test type

First author [ref.]	Country (income category)	Study design	Laboratory setting	Population	Number tested: RIF	Number tested: INH	Direct or indirect	Smear status	Condition of specimen	Phenotypic reference standard
Hain Genotype MTBDRplus V1										
AL-MUTAIRI [23] [#]	Kuwait (A)	Case-control	National reference	Culture positives	125	125	Indirect	N/A	Unknown	Radiometric BacTec 460
ALBERT [24]	Uganda (B)	Cross-sectional	National reference	At risk for MDR-TB	97	97	Direct	Positive	Frozen	Non-radiometric BacTec MGIT
ANEK-VORAPONG [25]	Thailand (B)	Unclear	Regional	Culture positives	50	50	Indirect	N/A	Frozen	Non-radiometric BacTec MGIT
ANEK-VORAPONG [25] [#]	Thailand (B)	Cross-sectional	Regional	Smear positives	164	164	Direct	Positive	Frozen	Non-radiometric BacTec MGIT
ASANTE-POKU [26] [#]	Ghana (B)	Cross-sectional	Regional	Smear positives	113	113	Indirect	Positive	Frozen	Proportion method
ASENCIOS [27]	Peru (B)	Cross-sectional	National reference	Culture positives	95	95	Indirect	N/A	Unknown	Proportion method
ASENCIOS [27]	Peru (B)	Unclear	National reference	Smear positives	100	100	Direct	Positive	Unknown	Proportion method
AUNG [28] [#]	Myanmar (B)	Cross-sectional	National reference	Smear positives	189	189	Indirect	Positive	Unknown	Proportion method
AURIN [29]	Bangladesh (B)	Cross-sectional	National reference	At risk for MDR-TB	277	277	Direct	Positive	Fresh	Proportion method
BANU [31]	Bangladesh (B)	Cross-sectional	Regional	At risk for MDR-TB	79	87	Direct	Positive	Unknown	Proportion method
BARNARD [32]	South Africa (B)	Cross-sectional	Regional	Smear positives	484	479	Direct	Positive	Unknown	Proportion method
BROSSIER [33] [#]	France (A)	Unclear	National reference	Unspecified	113	113	Indirect	N/A	Frozen	Proportion method
BWANGA [34]	Uganda (B)	Unclear	National reference	Unspecified	31	31	Indirect	N/A	Unknown	Proportion method
CABIBBE [35]	Uganda (B)	Unclear	Regional	Unspecified	91	91	Indirect	N/A	Unknown	Non-radiometric BacTec MGIT
CABIBBE [35]	Uganda (B)	Unclear	Regional	Unspecified	49	49	Direct	Both	Unknown	Non-radiometric BacTec MGIT
CAUSSE [37]	Spain (A)	Unclear	Regional	Smear positives	41	41	Indirect	Positive	Unknown	Non-radiometric BacTec MGIT
CAUSSE [37]	Spain (A)	Unclear	Regional	Smear positives	18	18	Direct	Positive	Frozen	Non-radiometric BacTec MGIT
CHEN [38]	China (B)	Cross-sectional	Regional	Smear positives	326	326	Direct	Positive	Frozen	Proportion method
CHRYSSANTHOU [39]	Sweden (A)	Cross-sectional	Regional	Culture positives	477	477	Indirect	N/A	Fresh	Non-radiometric BacTec MGIT
CHRYSSANTHOU [39]	Sweden (A)	Cross-sectional	Regional	Culture positives	90	90	Direct	Both	Fresh	Non-radiometric BacTec MGIT

Continued

TABLE 1 Continued

First author [ref.]	Country (income category)	Study design	Laboratory setting	Population	Number tested: RIF	Number tested: INH	Direct or indirect	Smear status	Condition of specimen	Phenotypic reference standard
DAUM [40]	South Africa (B)	Cross-sectional	Regional	Culture positives	26	26	Indirect	N/A	Frozen	Non-radiometric BacTec MGIT
DORMAN [41] [#]	South Africa (B)	Cross-sectional	Regional	Prior screened	223	223	Direct	Both	Fresh	Non-radiometric BacTec MGIT
DUBOIS CAUWELAERT [42]	Madagascar (B)	Cross-sectional	Regional	Culture positives	254	254	Direct	Positive	Unknown	Proportion method
ELISEEV [43]	Russia (A)	Cross-sectional	Regional	Smear positives	211	211	Direct	Positive	Unknown	Non-radiometric BacTec MGIT
EVANS [44]	South Africa (B)	Unclear	Regional	Culture positives	223	223	Indirect	N/A	Unknown	Non-radiometric BacTec MGIT
FABRE [45] ^{#,¶}	Multiple (C)	Unclear	Regional	Culture positives	144		Indirect	N/A	Frozen	Radiometric BacTec 460
FAROOQI [46] [#]	Pakistan (B)	Cross-sectional	Regional	Smear positives	105	105	Direct	Positive	Fresh	Proportion method
FELKEL [47] [#]	Nigeria (B)	Cross-sectional	Regional	Known MDR-TB	32	32	Direct	Unclear	Frozen	Non-radiometric BacTec MGIT
FERRO [48]	Colombia (B)	Cross-sectional	National reference	Unspecified	221	222	Indirect	N/A	Frozen	Proportion method
FRIEDRICH [49]	South Africa (B)	Cross-sectional	Regional	Prior screened	94	94	Direct	Both	Fresh	Non-radiometric BacTec MGIT
GAUTHIER [50]	Haiti (B)	Cross-sectional	National reference	Smear positives	221	221	Direct	Positive	Unknown	Non-radiometric BacTec MGIT
GITTI [51]	Greece (A)	Cross-sectional	Regional	Culture positives	221	221	Indirect	N/A	Unknown	Proportion method
HILLEMANN [52] [#]	Germany (A)	Case-control	National reference	Culture positives	125	125	Indirect	N/A	Unknown	Mixed
HILLEMANN [52] [#]	Germany (A)	Unclear	National reference	Smear positives	72	72	Direct	Positive	Unknown	Mixed
HUANG [53] [#]	China (B)	Cross-sectional	Regional	Smear positives	215	215	Indirect	Positive	Unknown	Proportion method
HUANG [54] [#]	Taiwan (B)	Unclear	Regional	Culture positives	272	272	Indirect	N/A	Unknown	Mixed
HUANG [55] [#]	Taiwan (B)	Unclear	National reference	Culture positives	324	324	Indirect	N/A	Unknown	Proportion method
HUYEN [56]	Vietnam (B)	Case-control	Regional	Culture positives	110	110	Indirect	Positive	Frozen	Proportion method
IMPERIALE [57]	Argentina (B)	Unclear	National reference	Culture positives	30	30	Indirect		Frozen	Mixed
IMPERIALE [57] [#]	Argentina (B)	Unclear	National reference	Smear positives	70	70	Direct	Positive	Frozen	Mixed

Continued

TABLE 1 Continued

First author [ref.]	Country (income category)	Study design	Laboratory setting	Population	Number tested: RIF	Number tested: INH	Direct or indirect	Smear status	Condition of specimen	Phenotypic reference standard
JIN [58] [#]	China (B)	Unclear	Regional	Culture positives	237	237	Indirect	N/A	Unknown	Absolute concentration
KAPATA [59]	Zambia (B)	Cross-sectional	National reference	Smear positives	598	594	Direct	Positive	Frozen	Proportion method
KHADKA [60]	Nepal (B)	Cross-sectional	Unknown	Culture positives	207	207	Indirect	N/A	Unknown	Absolute concentration
KUMAR [61]	India (B)	Unclear	National reference	Culture positives	141	141	Indirect	N/A	Unknown	Non-radiometric BacTec MGIT
LACOMA [62] [#]	Spain (A)	Unclear	Unknown	Culture positives	62	62	Indirect	N/A	Frozen	Radiometric BacTec 460
LACOMA [62]	Spain (A)	Unclear	Unknown	Unspecified	53	53	Direct	Both	Frozen	Radiometric BacTec 460
LI [63] [#]	China (B)	Cross-sectional	Unknown	Smear positives	1370	1370	Direct	Positive	Unknown	Proportion method
LUETKEMEYER [64]	Multiple (B)	Cross-sectional	Regional	HIV positives	303	301	Direct	Both	Fresh	Non-radiometric BacTec MGIT
LYU [65]	South Korea (B)	Cross-sectional	Regional	Smear positives	168		Direct	Both	Unknown	Absolute concentration
MACEDO [66]	Portugal (A)	Cross-sectional	National reference	Smear positives	68	68	Direct	Positive	Frozen	Radiometric BacTec 460
MASCHMANN RDE [67] [#]	Brazil (B)	Cross-sectional	Regional	At risk for MDR-TB	66	66	Direct	Positive	Fresh	Proportion method
MIOTTO [68]	Italy (A)	Cross-sectional	Regional	Culture positives	206	206	Indirect	N/A	Frozen	Non-radiometric BacTec MGIT
MIOTTO [68]	Italy (A)	Cross-sectional	Regional	Culture positives	78	78	Direct	Both	Unknown	Non-radiometric BacTec MGIT
MIOTTO [69]	Burkina Faso (B)	Cross-sectional	National reference	At risk for MDR-TB	31	31	Direct	Both	Frozen	Non-radiometric BacTec MGIT
MIOTTO [69]	Burkina Faso (B)	Cross-sectional	National reference	At risk for MDR-TB	11	11	Direct	Both	Frozen	Non-radiometric BacTec MGIT
MIRONOVA [70]	Multiple (C)	Cross-sectional	National reference	Unspecified	243	243	Indirect	N/A	Unknown	Non-radiometric BacTec MGIT
MIRONOVA [70]	Multiple (C)	Cross-sectional	National reference	Unspecified	74	74	Indirect	N/A	Unknown	Proportion method
N'GUESSAN [71]	Cote D'Ivoire (B)	Cross-sectional	National reference	Smear positives	120	120	Direct	Positive	Fresh	Non-radiometric BacTec MGIT
NATHAVITHARANA [72] [#]	Multiple (C)	Case-control	National reference	Culture positive	376	378	Indirect	N/A	Frozen	Mixed
NATHAVITHARANA [72]	Multiple (C)	Cross-sectional	National reference	At risk for MDR-TB	455	462	Direct	Both	Fresh	Mixed

Continued

TABLE 1 Continued

First author [ref.]	Country (income category)	Study design	Laboratory setting	Population	Number tested: RIF	Number tested: INH	Direct or indirect	Smear status	Condition of specimen	Phenotypic reference standard
NIEHAUS [73]	South Africa (B)	Cross-sectional	Regional	Unspecified	994	994	Indirect	N/A	Unknown	Proportion method
NIKOLAYEVSKYY [74]	Russia (A)	Cross-sectional	Regional	Smear positives	163	163	Direct	Positive	Fresh	Mixed
NWOFOR [75]	Nigeria (B)	Cross-sectional	National reference	Culture positives	97	97	Indirect	N/A	Unknown	Proportion method
OCHERETINA [76] ^{#,¶}	Haiti (B)	Unclear	National reference	Culture positives	153		Indirect	N/A	Unknown	Non-radiometric BacTec MGIT
RAIZADA [77]	India (B)	Cross-sectional	Regional	At risk for MDR-TB	267	267	Direct	Positive	Fresh	Proportion method
RAVEENDRAN [78]	India (B)	Cross-sectional	Regional	Culture positives	69	69	Indirect	N/A	Fresh	Non-radiometric BacTec MGIT
RAVEENDRAN [78]	India (B)	Cross-sectional	Regional	Smear positives	16	16	Direct	Positive	Fresh	Non-radiometric BacTec MGIT
RIGOUTS [80]	Tanzania (B)	Cross-sectional	National reference	Smear positives	303	303	Direct	Positive	Unknown	Proportion method
RUFAL [81] [¶]	India (B)	Cross-sectional	National reference	Smear positives	23		Direct	Positive	Fresh	Non-radiometric BacTec MGIT
SANGSAYUNH [82]	Thailand (B)	Cross-sectional	Regional	At risk for MDR-TB	18	19	Direct	Both	Fresh	Proportion method
SCHON [83] [¶]	Sweden (A)	Case-control	Regional	Culture positives	95		Indirect	N/A	Frozen	Absolute Concentration
SCOTT [84]	South Africa (B)	Cross-sectional	Unknown	All-comers	89	89	Direct	Both	Frozen	Non-radiometric BacTec MGIT
SHUBLADZE [85]	Georgia (B)	Cases only	National reference	Known MDR-TB	634	634	Indirect	N/A	Unknown	Proportion method
SIMONS [86]	Netherlands (A)	Cross-sectional	National reference	Unspecified	2649	2649	Indirect	N/A	Unknown	Non-radiometric BacTec MGIT
SINGHAL [87]	India (B)	Cross-sectional	National reference	Smear positives	120	120	Direct	Positive	Unknown	Non-radiometric BacTec MGIT
TESSEMA [88]	Ethiopia (B)	Cross-sectional	Regional	Smear positives	260	260	Indirect	Positive	Unknown	Non-radiometric BacTec MGIT
THO [89]	Vietnam (B)	Case-control	National reference	Culture positives	150	150	Indirect	N/A	Frozen	Proportion method
TOLANI [90]	India (B)	Cross-sectional	Unknown	Smear positives	88	88	Indirect	Positive	Unknown	Radiometric BacTec 460
TOLANI [90]	India (B)	Cross-sectional	Unknown	Smear positives	67	67	Indirect	Positive	Unknown	Radiometric BacTec 460
TUKVADZE [91]	Georgia (B)	Cross-sectional	National reference	Smear positives	474	474	Direct	Positive	Frozen	Mixed
VJDEA [92] [#]	Denmark, Lithuania (C)	Case-control	Supra-national reference	Culture positives	115	115	Indirect	N/A	Unknown	Radiometric BacTec 460
YADAV [93]	India (B)	Cross-sectional	Regional	At risk for MDR-TB	242	242	Direct	Positive	Fresh	Proportion method
YORDANOVA [94]	Bulgaria (B)	Cases only	National reference	Known MDR-TB	66	66	Indirect	N/A	Unknown	Non-radiometric BacTec MGIT

Continued

TABLE 1 Continued

First author [ref.]	Country (income category)	Study design	Laboratory setting	Population	Number tested: RIF	Number tested: INH	Direct or indirect	Smear status	Condition of specimen	Phenotypic reference standard
Hain Genotype MTBDRplus V2										
BABLISHVILI [30]	Georgia (B)	Cross-sectional	National reference	Smear positives	350	350	Direct	Positive	Fresh	Mixed
CATANZARO [36]	Moldova, India, South Africa (B)	Cross-sectional	Regional	At risk for MDR-TB	914	914	Direct	Positive	Unknown	Non-radiometric BacTec MGIT
CRUDU [15]	Moldova (B)	Cross-sectional	National reference	All-comers	156	156	Direct	Both	Fresh	Proportion method
NATHAVITHARANA [72] [#]	Multiple (C)	Case-control	National reference	Culture positive	376	378	Indirect	N/A	Frozen	Mixed
NATHAVITHARANA [72]	Multiple (C)	Cross-sectional	National reference	At risk for MDR-TB	452	452	Direct	Both	Fresh	Mixed
Nipro NTM+MDR Detection Kit 2										
MITARAI [17] [#]	Japan (A)	Cross-sectional	National reference	Unspecified	314	314	Indirect	N/A	Unknown	Unknown
MITARAI [17]	Japan (A)	Cross-sectional	National reference	Unspecified	55	52	Direct	Both	Frozen	Proportion method
RIENTHONG [79] [#]	Thailand (B)	Case-control	Unknown	Culture positives	260	260	Indirect	N/A	Frozen	Non-radiometric BacTec MGIT
RIENTHONG [79]	Thailand (B)	Cross-sectional	Unknown	Unspecified	127	127	Direct	Both	Fresh	Non-radiometric BacTec MGIT
NATHAVITHARANA [72] [#]	Multiple (C)	Case-control	National reference	Culture positive	378	378	Indirect	N/A	Frozen	Mixed
NATHAVITHARANA [72]	Multiple (C)	Cross-sectional	National reference	At risk for MDR-TB	475	474	Direct	Both	Fresh	Mixed

MDR-TB: multidrug-resistant tuberculosis. [¶]: these studies only contributed data to PICO A1a and 1b (Rifampicin resistance detection); [#]: these studies contributed data from which a composite reference standard could be derived.

TABLE 2 Characteristics of included studies for *M. tuberculosis* detection, grouped alphabetically by index test type

Author, year	Country (income category)	Study design	Laboratory setting	Population	Number tested	Direct or indirect	Smear status	Condition of specimen	Processed per manufacturer's instructions	Culture reference standard
Hain Genotype MTBDRplus V1										
DORMAN [41]	South Africa (B)	Cross-sectional	Regional	Prior screened	223	Direct	Both	Fresh	Yes	Liquid: MGIT 960
FELKEL [47]	Nigeria (B)	Cross-sectional	Regional	Known MDR-TB	110	Direct	Unclear	Frozen	Yes	Liquid: MGIT 960
FRIEDRICH [49]	South Africa (B)	Cross-sectional	Regional	Prior screened	126	Direct	Both	Fresh	Yes	Liquid: MGIT 960
LUETKEMEYER [64]	Multiple (B)	Cross-sectional	Regional	HIV positives	595	Direct	Both	Fresh	Yes	Liquid: MGIT 960
SCOTT [84]	South Africa (B)	Cross-sectional	Unknown	All-comers	177	Direct	Both	Frozen	Yes	Liquid: MGIT 960
Hain Genotype MTBDRplus V2										
CRUDU [15]	Moldova (B)	Cross-sectional	National reference	All-comers	336	Direct	Both	Fresh	Yes-GenoLyse and GeneXtract	Liquid: MGIT 960

MDR-TB: multidrug-resistant tuberculosis.

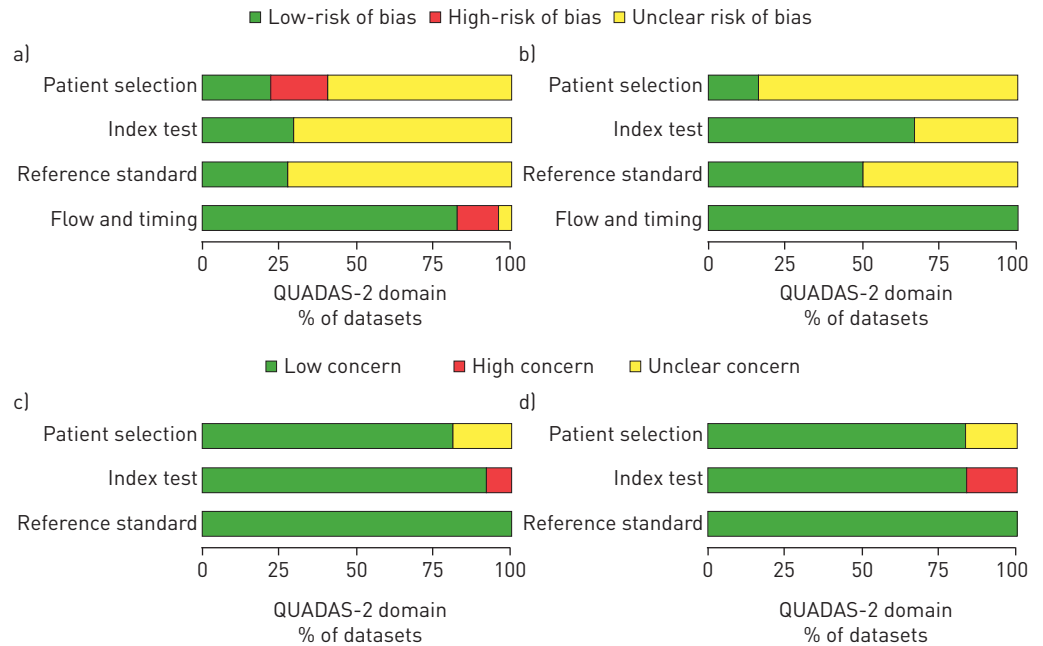


FIGURE 2 QUADAS-2 summaries. a and c) Risk of Bias and Applicability Concerns summary about each QUADAS-2 domain presented as percentages across the 94 included datasets for rifampicin (RIF) and isoniazid (INH) resistance compared with phenotypic culture-based reference standard (of note, four datasets only contributed to RIF). The summaries for the datasets for RIF and INH compared with composite reference standard are not displayed separately since these datasets are a subset of the 94 datasets displayed below and thus the figures displayed are thought to be accordingly representative. b and d) Risk of Bias and Applicability Concerns summary about each QUADAS-2 domain presented as percentages across the six included datasets for *Mycobacterium tuberculosis* detection compared with a culture-based reference standard.

supplementary material). Only five datasets reported indeterminate results for indirectly tested isolates but these percentages were lower than for direct testing, with a median of 1.0% and range of 0.5–2.1% for rifampicin and 0.5% and 0.5–1.0% for isoniazid. Only three datasets that performed direct testing for *M. tuberculosis* detection reported indeterminate results, with a median of 1.0% and range of 0.7–1.7%. Data on smear grade were limited. Studies did not typically report whether repeat testing was performed on indeterminate results. For comparison purposes, four datasets reported the number of contaminated cultures obtained using the culture reference standard, with a median of 7.6% and range of 2.8–17.2% of the total specimens.

Analysis of primary outcomes of interest

Diagnosis of RIF resistance using a phenotypic reference standard

Pooled Analysis for all LPAs on all specimen types

91 datasets were included in the bivariate analysis, with a total of 21 225 samples that included 6789 (32%) confirmed RIF-resistant tuberculosis cases. Meta-analysis revealed a pooled sensitivity of 96.7% (95% CI 95.6–97.5%) and specificity of 98.8% (95% CI 98.2–99.2%) (table 3). Results were largely homogenous, with a small proportion of studies being outliers. Pooled analysis stratified by LPA (appendix F, table S2 in the supplementary material) demonstrated a slightly lower sensitivity for Hain V2 and Nipro (95.0% and 94.3% compared with 97.1% for Hain V1) although confidence intervals overlapped and specificity was similar (98.3%, 98.1% and 98.9% respectively).

Direct testing

48 datasets tested RIF resistance detection with LPA directly from specimens, with a total of 10 560 samples that included 2876 (27%) confirmed RIF-resistant tuberculosis cases. The pooled sensitivity was 96.3% (95% CI 94.6–97.5%) and specificity was 98.2% (95% CI 97.2–98.8%) (table 3, figure 3a). Outliers with lower sensitivity and specificity were predominantly datasets with limited numbers of resistant specimens (<10) and thus accompanied by very wide confidence intervals (figure 4).

Indirect testing

43 datasets tested RIF resistance detection with LPA indirectly from isolates, with a total of 10 696 samples that included 3913 (37%) confirmed RIF-resistant tuberculosis cases. The pooled sensitivity was 96.9%

TABLE 3 Diagnostic accuracy of line probe assays for all three assays combined for rifampicin (RIF) and isoniazid (INH) resistance and multidrug-resistant tuberculosis (MDR-TB) detection

Reference standard	Test	Direct or indirect	Smear status	Datasets (samples) n	Sensitivity (95% CI)	Specificity (95% CI)
Phenotypic drug susceptibility testing	RIF	Both	All	91 (21 225)	96.7% (95.6–97.5)	98.8% (98.2–99.2)
	RIF	Direct	All	48 (10 560)	96.3% (94.6–97.5)	98.2% (97.2–98.8)
	RIF	Indirect	All	43 (10 696)	96.9% (95.4–98.0)	99.3% (98.6–99.6)
Composite drug susceptibility testing	RIF	Both	All	23 (5483)	95.3% (93.4–96.6)	99.5% (98.6–99.8)
Phenotypic drug susceptibility testing (same samples as composite drug susceptibility testing)	RIF	Both	All	23 (5484)	95.2% (93.2–96.7)	98.9% (98.0–99.4)
Phenotypic drug susceptibility testing	INH	Both	All	87 (20 954)	90.2% (88.2–91.9)	99.2% (98.7–99.5)
	INH	Direct	All	46 (10 472)	89.2% (85.8–91.9)	98.4% (97.5–98.9)
	INH	Indirect	All	41 (10 462)	91.0% (88.6–93.0)	99.7% (99.3–100)
Composite drug susceptibility testing	INH	Both	All	24 (4516)	85.1% (80.8–88.6)	99.9% (99.6–99.9)
Phenotypic drug susceptibility testing (same samples as composite drug susceptibility testing)	INH	Both	All	24 (4520)	85.0% (80.5–88.6)	99.5% (99.1–99.8)
Phenotypic drug susceptibility testing	MDR-TB	Both	All	57 (13 033)	92.9% (90.2–94.7)	99.3% (98.7–99.6)
Composite drug susceptibility testing	MDR-TB	Both	All	12 (2745)	86.6% (81.9–90.3)	99.6% (98.9–99.9)
Phenotypic drug susceptibility testing (same samples as composite drug susceptibility testing)	MDR-TB	Both	All	12 (2745)	86.9% (82.1–90.7)	99.5% (97.9–99.9)

(95% CI 95.5–98.0%) and specificity was 99.3% (95% CI 98.6–99.6%) (table 3; and appendix F, figure S8a in the supplementary material). Point estimates for sensitivity for individual studies were even more homogenous than those for direct testing (appendix F, figure S4 in the supplementary material). The reasons for the outlier studies with lower sensitivities were unclear as the populations tested (all-comers *versus* those with MDR-TB risk) differed in the respective studies [60, 73, 90]. One outlier demonstrated a lower specificity (78.3%, 95% CI 63.6–89.1%) for specimens tested by solid (Löwenstein–Jensen) rather than liquid (Mycobacteria Growth Indicator Tube (MGIT)) culture [70].

Diagnosis of RIF resistance using a composite reference standard

Pooled Analysis for all LPAs on all specimen types

23 datasets contained data comparing LPA with a composite reference standard (using the results from targeted sequencing of either the RIF-resistance determining region or *rpoB* gene and phenotypic DST), with a total of 5483 samples that included 2091 (38%) RIF-resistant *M. tuberculosis* cases [17, 23, 26, 41, 45–47, 52–58, 62, 63, 67, 79, 92]. Most studies only performed sequencing on discrepant results, thus results from this analysis may be potentially biased in favour of the LPAs. Bivariate meta-analysis of these studies revealed a pooled sensitivity of 95.3% (95% CI 93.4–96.6%) and specificity of 99.5% (95% CI 98.6–99.8%) (table 3).

Specificity increased when a composite standard was used as 37 LPA “false-positive results” based on comparison to phenotypic DST (from 11 datasets) were reclassified as true positives as sequencing confirmed the presence of known resistance-conferring mutations (appendix F, table S3a in the supplementary material). Of note, the sensitivity was lower in this subset of datasets for which data on a composite reference standard could be derived compared with the overall dataset, which we hypothesise may be due to some selection bias in the studies that performed targeted sequencing alongside phenotypic DST.

Heterogeneity across studies was limited (appendix F, figure S5 in the supplementary material). MASCHMANN RDE *et al.* [67] demonstrated a sensitivity of 82.8% and stated that two out of the five specimens incorrectly classified had insertions in codons 516–517 which may have caused hybridisation of the corresponding wild-type probe (wt3 for codons 517–520) and the other three were wild-type on sequencing, suggesting that resistance may be driven by mutations outside of the *rpoB* hotspot.

Diagnosis of INH resistance using a phenotypic reference standard

Pooled analysis for all LPAs on all specimen types

87 datasets were included in the bivariate analysis, with a total of 20 954 samples that included 8135 (39%) confirmed INH-resistant tuberculosis cases. Meta-analysis revealed a pooled sensitivity of 90.2% (95% CI 88.2–91.9%) and specificity of 99.2% (95% CI 98.7–99.5%) (table 3). Results were moderately heterogeneous

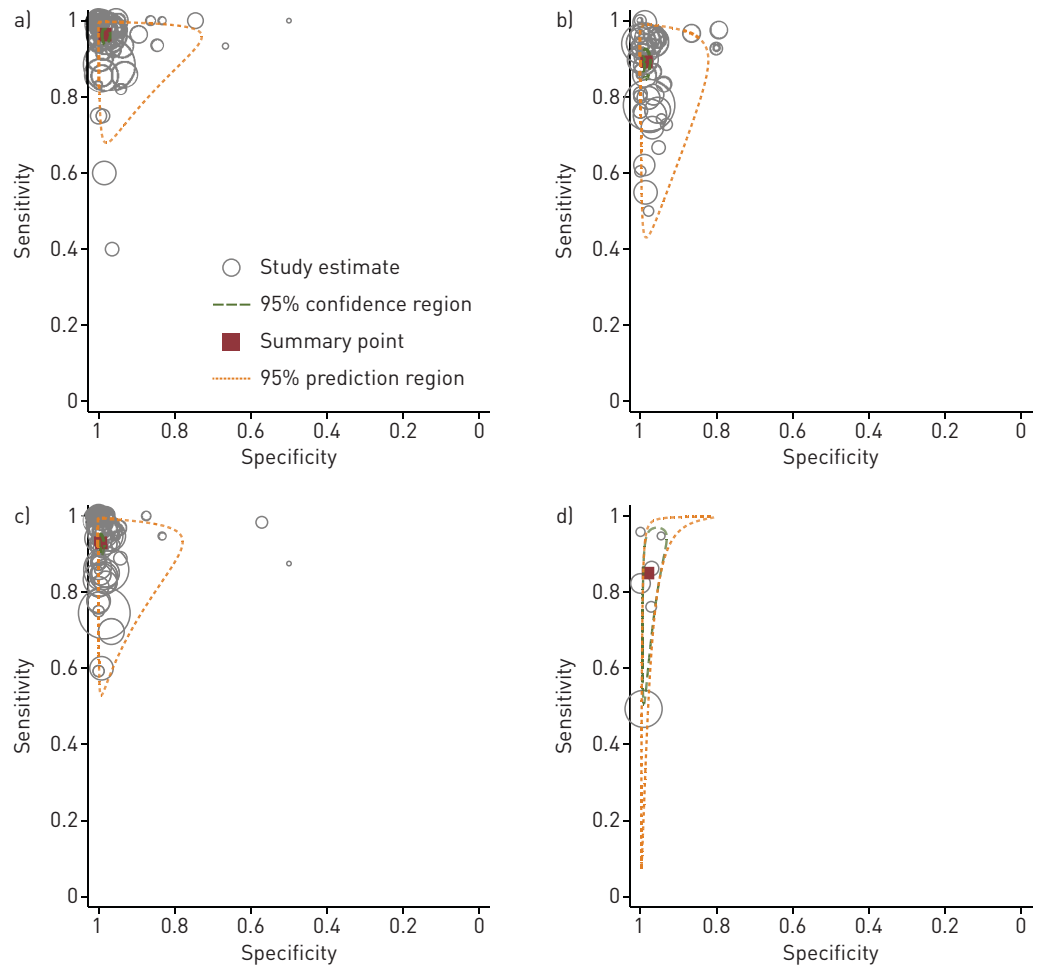


FIGURE 3 Hierarchical summary receiver operating characteristic graphs of summary estimates. Bivariate analysis of the sensitivity and specificity for all line probe assays for the diagnosis of drug resistance detection compared with a phenotypic reference standard for specimens tested directly for a) rifampicin resistance, b) isoniazid resistance, c) multi-drug resistance and d) the detection of *Mycobacterium tuberculosis* compared to a culture reference standard. In the plots below, the red squares represent the pooled summary estimates, the dashed red lines represent the 95% confidence region and the dashed green lines represent the 95% prediction region. The individual circles represent each study and the size of the circle is proportional to the total sample size.

for sensitivity, whereas specificity estimates were more homogeneous. Pooled analysis stratified by LPA (appendix F, table S2 in the supplementary material) demonstrated a lower sensitivity for Nipro (86.9%) and higher sensitivity for Hain V2 (93.6%) compared with Hain V1 (90.2%) although specificity was similar (99.1%, 99.1% and 99.2%) respectively.

Direct testing

46 datasets tested INH-resistance detection with LPA directly from specimens against a phenotypic reference standard, with a total of 10472 samples that included 3576 (34%) confirmed INH-resistant tuberculosis cases. The pooled sensitivity across studies was 89.2% (95% CI 85.8–91.9%) and specificity was 98.4% (95% CI 97.5–98.9%) (table 3, figure 3b). Greater heterogeneity was noted for INH-sensitivity compared with RIF for sensitivity (figure 5). Several outliers had limited numbers of resistant specimens (<10) and were thus accompanied by very wide confidence intervals [17, 31, 82, 84]. Explanations for outlier results included the known geographic variation of mutations and heteroresistance.

Indirect testing

40 datasets tested INH resistance detection with LPA indirectly from isolates against a phenotypic reference standard, with a total of 10462 samples that included 4559 (44%) confirmed INH-resistant tuberculosis cases. The pooled sensitivity across studies was 91.0% (95% CI 88.6–93.0%), which was higher than seen with direct testing, as was the case for specificity, which was 99.7% (95% CI 99.3–100.0%) (table 3; and

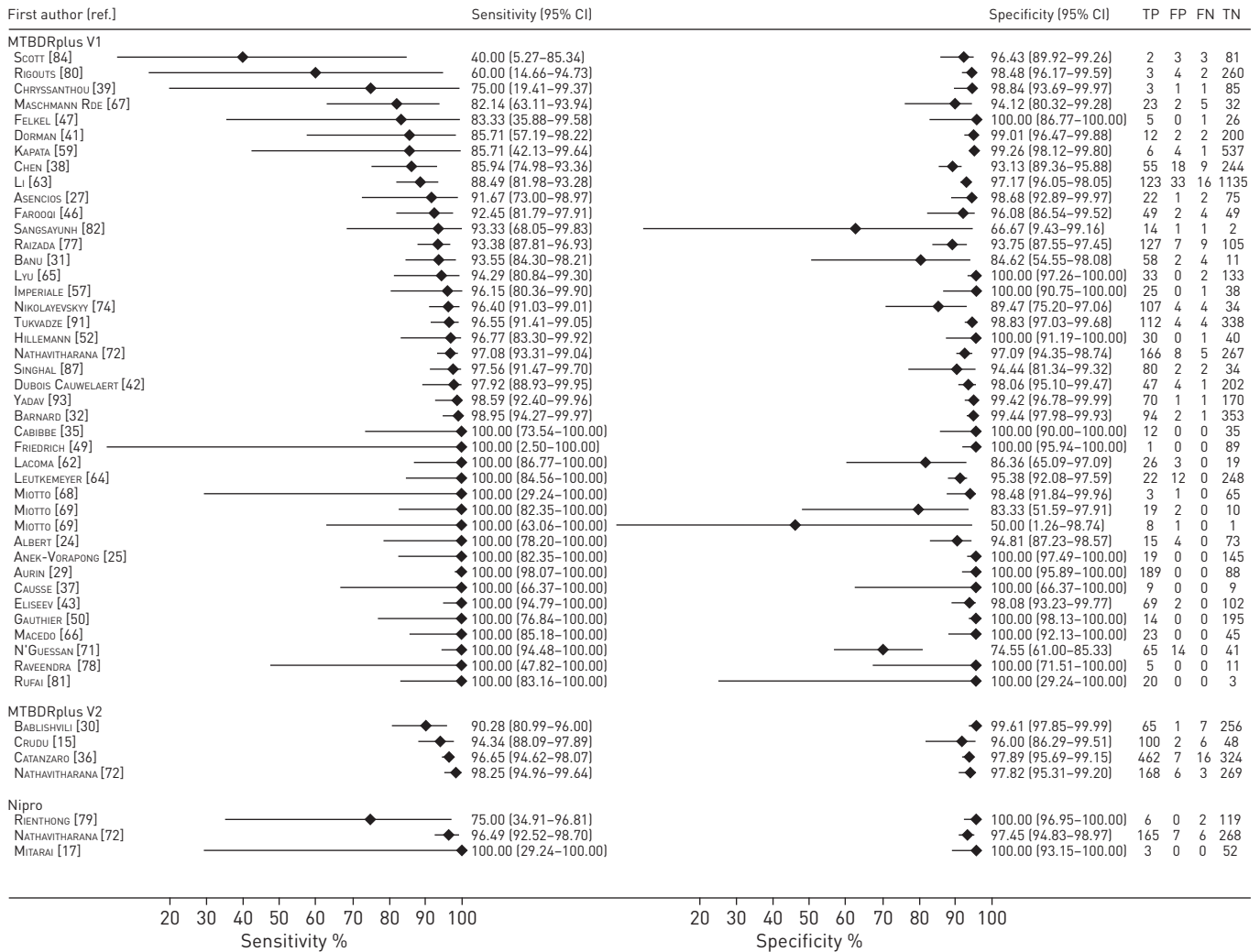


FIGURE 4 Forest plots demonstrating the sensitivity and specificity of all line probe assays for rifampicin resistance-detection for sputum specimens tested directly compared with phenotypic drug susceptibility testing. TP: true positive; FP: false positive; FN: false negative; TN: true negative.

appendix F, figure S8b in the supplementary material). Several studies were outliers for sensitivity but specificity was largely homogeneous (appendix F, figure S6 in the supplementary material). Reasons for lower sensitivity include the use of different types of phenotypic DST within a study [17], the presence of less common resistance mutations due to geographic variation and difficulty detecting low-level INH resistance [62]. The outlier for specificity only contained three INH-sensitive strains [90].

Diagnosis of INH resistance using a composite reference standard

Pooled analysis for all LPAs on all specimen types

24 datasets contained data comparing LPA with a composite reference standard, with a total of 4516 samples that included 2346 (52%) INH-resistant *M. tuberculosis* cases [17, 23, 25, 26, 28, 33, 41, 46, 47, 52-55, 58, 62, 67, 68, 79, 92]. Bivariate meta-analysis of these studies revealed a pooled sensitivity of 85.1% (95% CI 80.8-88.6%) and specificity of 99.9% (95% CI 99.6-100.0%) (table 3). Bivariate analysis of the same 24 datasets compared to phenotypic DST revealed a pooled sensitivity of 85.0% (95% CI 80.5-88.6%) and specificity of 99.5% (95% CI 99.1-99.8%).

Sequencing also revealed resistance mutations that were not detected by LPA (appendix F, table S3b in the supplementary material). For example, 10 of the 11 strains with a rarer *katG* mutation S315N were not detected in the study by JIN *et al.* [58] due to the lack of the appropriate mutation probe in the Hain V1 assay and because the wild-type band also failed to disappear. Although seven LPA false-positive results (from six datasets) were reclassified as true positives in total based on sequencing confirming a known resistance mutation (four *katG* S315T mutations and three *inhA* c-15t mutations), specificity barely increased when a composite standard was used.

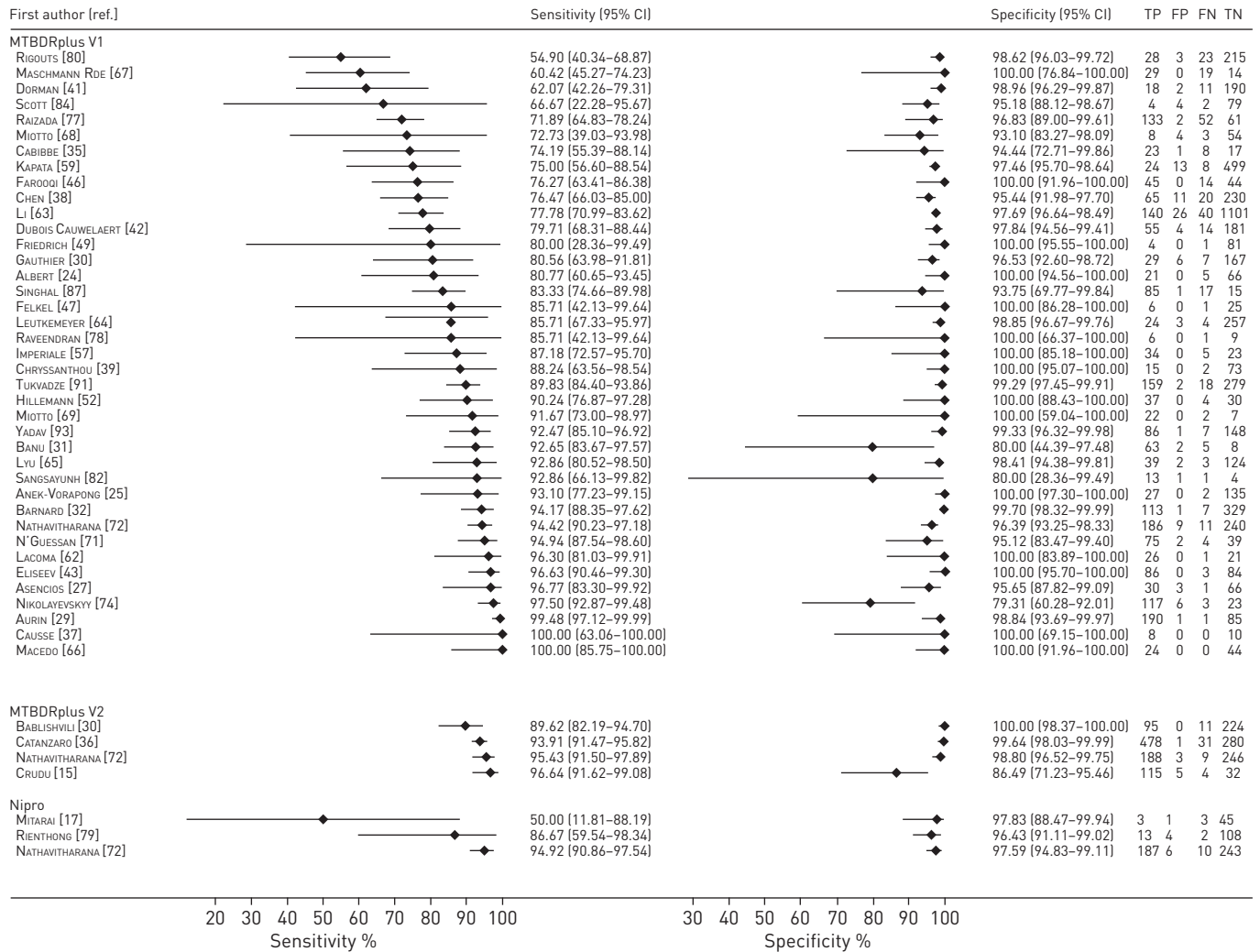


FIGURE 5 Forest plots demonstrating sensitivity and specificity of all line probe assays for isoniazid-resistance detection for sputum specimens tested directly compared with phenotypic drug susceptibility testing. TP: true positive; FP: false positive; FN: false negative; TN: true negative.

Heterogeneity assessment (appendix F, figure S7 in the supplementary material) demonstrated homogenous results for specificity, which was largely also the case for sensitivity aside from a few outliers. MITARAI *et al.* [17] demonstrated a specificity of 61.6% (95% CI 52.9–69.7%). Of the 53 isolates incorrectly identified as sensitive by LPA, 24 had a range of rare *katG* mutations not identified by any of the *katG* probes, 17 had *fabG1 inhA* mutations and 12 were identified as wild-type by sequencing. MASCHMANN RDE *et al.* [67] demonstrated a sensitivity of 60.4% (95% CI 45.3–74.2%) and reported that all 19 strains misclassified as susceptible on LPA were found to have wild-type *katG* and *inhA* genes according to targeted sequencing, indicating that there may have been mutations in other genes associated with INH resistance or efflux systems that could not be detected by the LPA.

Diagnosis of multidrug resistance

Pooled analysis for all LPAs on all specimen types

57 datasets included data on the diagnostic accuracy of LPA for MDR-TB detection, with a total of 13033 samples that included 4248 (33%) confirmed MDR-TB cases [23–29, 32, 34, 35, 37, 38, 41–44, 48, 50, 52–56, 58–64, 66–69, 71–73, 75, 78, 80, 82, 84, 88, 90, 91, 93]. Bivariate meta-analysis of these datasets revealed a pooled sensitivity of 92.9% (95% CI 90.4–94.8%) and specificity of 99.3% (95% CI 98.7–99.6%) (table 3 and figure 3c). Figure 6 demonstrates homogenous results for specificity aside from a few outliers in which the number of sensitive (non-MDR strains) was <15, which was largely also the case for these sensitivity outliers.

Comparison of diagnostic accuracy from direct versus indirect testing

Based on the analysis of all data, the estimates for sensitivity of LPA for RIF and INH resistance were almost identical for LPA performed directly on sputum specimens and indirectly on culture isolates (96.3% and

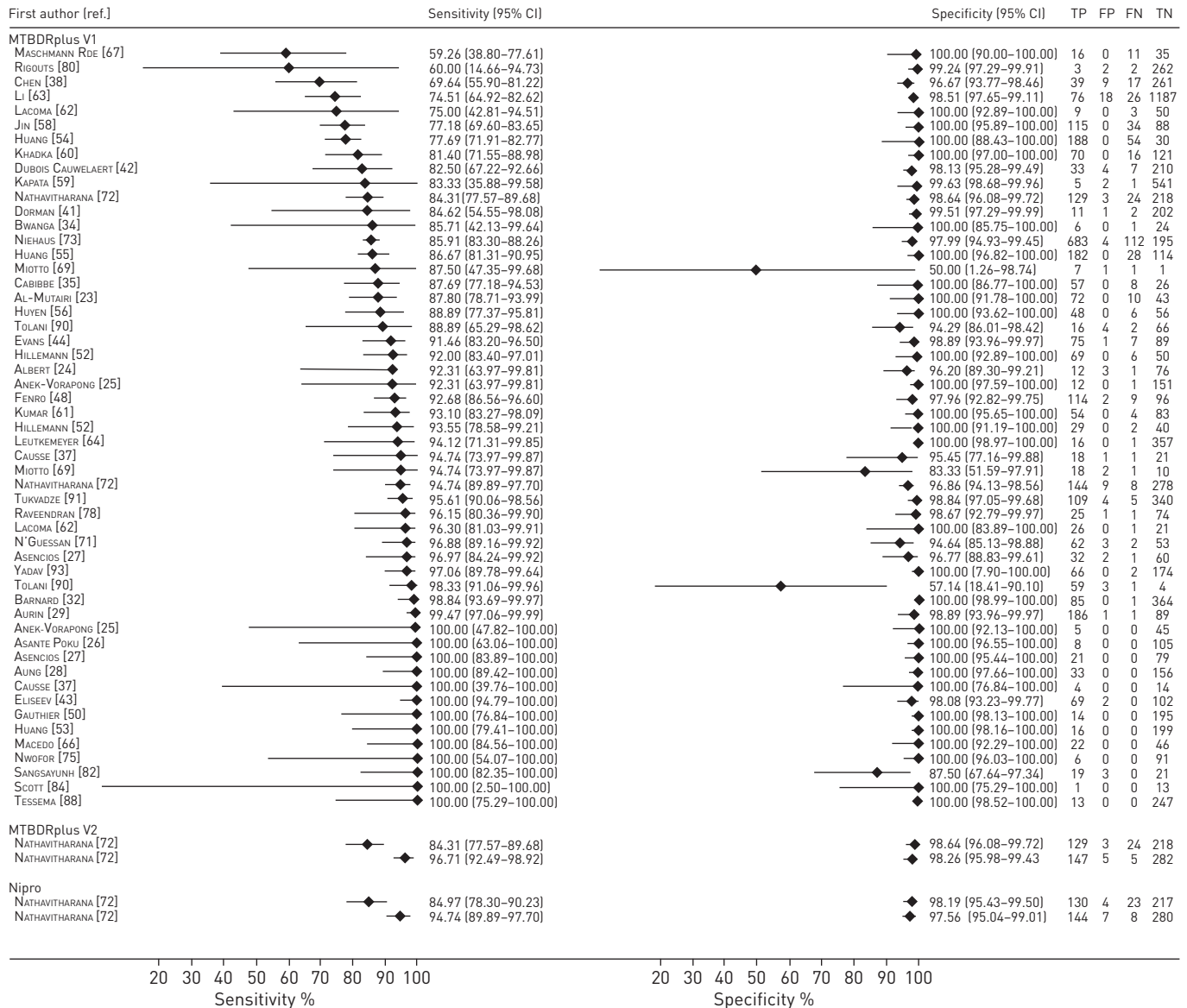


FIGURE 6 Forest plots demonstrating sensitivity and specificity of all line probe assays for multidrug-resistant tuberculosis detection for both specimen types compared with phenotypic drug susceptibility testing. TP: true positive; FP: false positive; FN: false negative; TN: true negative.

96.9% respectively for RIF, 89.2% and 91.0% for INH). Specificity was slightly increased for indirect testing (99.3% compared with 98.2% for RIF, 99.7% compared with 98.4% for INH). The summary point estimates approach the upper left-hand corner of the plots, suggesting good accuracy of LPAs for detection of RIF and INH resistance whether tested directly or indirectly. No studies performed LPA testing on specimens and culture isolates from the same patients precluding direct within-study comparisons.

Diagnosis of pulmonary M. tuberculosis using a culture-based reference standard

Data to answer this question were limited, as the majority of LPA studies identified by our search criteria did not report results for *M. tuberculosis* detection. Of the 21 datasets that did report data on *M. tuberculosis* detection, 15 studies were excluded because they either tested patients who were on treatment or did not specify that patients on treatment were excluded.

Six datasets were included in the bivariate analysis [15, 41, 47, 49, 64, 84], with a total of 3451 samples that included 1277 (37%) confirmed *M. tuberculosis* cases tested directly with LPA. Meta-analysis of datasets that reported both sensitivity and specificity revealed a pooled sensitivity of 85.0% (95% CI 70.0–93.3%) and specificity of 98.0% (95% CI 96.2–99.0%) independent of smear-status (table 4 and figure 3d). Of note, a *post hoc* bivariate analysis of the 21 datasets (including those that did not exclude patients on treatment) revealed a sensitivity of 94.8% (95% CI 87.8–97.9%) and specificity of 95.7% (95% CI 85.0–98.9%).

TABLE 4 Diagnostic accuracy of line probe assays for all three assays for *Mycobacterium tuberculosis* (MTB) detection

Reference standard	Test	Direct or indirect	Smear status	Datasets (samples) n	Sensitivity (95% CI)	Specificity (95% CI)
Culture reference	MTB	Direct	All	6 (3451)	85.0% [70.0–93.3]	98.0% [96.2–99.0]
	MTB	Direct	Positive	5 (802) [¶]	94.4% [89.4–99.4]	#
	MTB	Direct	Negative	5 (961)	44.4% [29.2–71.7]	98.9% [95.4–99.7]
Culture reference	MTB	Direct: fresh	Both	4	83.0% [61.9–93.6]	98.8% [97.2–99.5]
Culture reference	MTB	Direct: frozen	Both	2 ⁺	#	#

+: meta-analysis was not possible based on the number of datasets identified for this subset. #: not estimable. ¶: two of these five studies only reported on sensitivity so bivariate meta-analysis for specificity was not possible. Of the three studies that contributed data on smear positive specificity, estimates were 50% (*n.b.* only had 2 specimens that were MTB negative), 100% and 100%.

Investigation of heterogeneity

Moderate heterogeneity was seen among the sensitivity estimates for *M. tuberculosis* detection, which suggests that this pooled estimate has to be considered with caution (figure 7). Possible explanations include testing of a high proportion of smear negatives by CRUDU *et al.* [15] and DORMAN *et al.* [41] (74% and 50% respectively). SCOTT *et al.* [84] reported that they used frozen samples for LPA in comparison with fresh samples and used a non-standard protocol for phenotypic DST testing, which may have decreased sensitivity. Specificity estimates were homogeneous.

Primary sub-group analysis by smear status

Five out of six included datasets that evaluated LPA on direct testing of clinical specimens reported smear status and were further evaluated. Two studies only reported on sensitivity and so bivariate meta-analysis was not possible for the other three studies.

Univariate analysis of the smear positive data for all five studies, which accounted for 802 samples, of which 781 were confirmed *M. tuberculosis* cases, revealed a sensitivity of 94.4% (95% CI 89.4–99.4%). For the three studies that contributed data to specificity, estimates were 50%, 100% and 100% [15, 49, 64]. The specificity of 50% (95% CI 0.01–98.7%) by CRUDU *et al.* [15] represented one out of two *M. tuberculosis*-negative specimens.

Five studies provided data on *M. tuberculosis* detection in smear negative cases, which accounted for 961 samples, of which 487 were confirmed *M. tuberculosis* cases. Sensitivity estimates across the studies ranged from 0% to 76%. Four studies only contributed data to both sensitivity and specificity and a bivariate meta-analysis revealed a pooled sensitivity of 44.4% (95% CI 20.2–71.7%) and specificity of 98.9% (95% CI 95.4–99.7%). The dataset by FRIEDRICH *et al.* [49] only tested one non-*M. tuberculosis* smear-negative specimen that was misidentified as *M. tuberculosis* by LPA. Given the substantial heterogeneity and the small number of studies, these estimates have to be interpreted with caution.

Secondary sub-group analysis by specimen condition

Given the low numbers of datasets reporting information on specimen condition, only a limited analysis was possible. Two datasets performed LPA testing on frozen specimens and reported a sensitivity of 94.7% and 76.1% respectively [47, 84]. Bivariate meta-analysis of the four datasets that performed LPA testing on

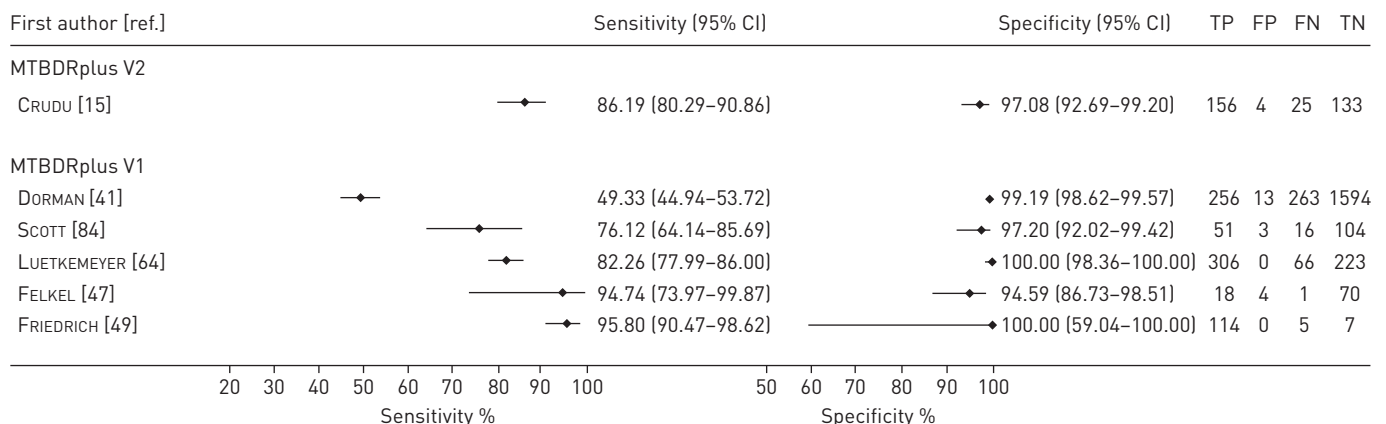


FIGURE 7 Forest plots demonstrating the sensitivity and specificity of all the line probe assays evaluated for the diagnosis of pulmonary *Mycobacterium tuberculosis* compared with culture. TP: true positive; FP: false positive; FN: false negative; TN: true negative.

fresh specimens demonstrated a pooled sensitivity of 83.0% (95% CI 61.9–93.6%). No conclusion could be drawn in regards to the effect of the specimen condition for the sensitivity of the test.

Secondary sub-group analysis by smear grade

It was not possible to perform a sub-group analysis to evaluate smear grade on the accuracy of LPA for *M. tuberculosis* detection due to the lack of reported data on this variable. NATHAVITHARANA *et al.* [72] found that smear grade affected indeterminate rates for RIF and INH detection (see section on indeterminate results) and other studies also mentioned that smear grade affected the number of valid results, often resulting in studies only evaluating smear-positive specimens or selecting specimens with the highest smear grade for analysis.

Sensitivity analyses

We assessed whether excluding studies that: selected for MDR-TB risk, used convenience sampling, used a case–control design or did not blind operators (or if studies were unclear on these criteria). These sensitivity analyses made no difference to any of the findings (appendix F, tables S4 and S5 in the supplementary material).

Discussion

Principal findings

For the detection of RIF resistance, pooled bivariate analyses from 21 225 samples (91 datasets) demonstrated a sensitivity of 96.7% and specificity of 98.8%. For the detection of INH resistance, pooled bivariate analyses from 20 954 samples (87 datasets) demonstrated a sensitivity of 90.2% and specificity of 99.2%. Sensitivity and specificity were similar for direct and indirect testing for both RIF and INH. While INH resistance was only detected with moderate sensitivity (90.2%), INH resistance is also highly correlated with RIF resistance in high-burden settings [95] and a negative result in the context of RIF resistance needs to be treated with caution. LPAs demonstrated good sensitivity (92.9%) and high specificity (99.3%) for MDR-TB detection.

For the detection of *M. tuberculosis*, data were far more limited and results have to be interpreted with caution. Pooled bivariate analyses from 3451 samples (six datasets) demonstrated a sensitivity of 85.0% and specificity of 98.0% on directly tested specimens. Data on smear status were limited. However, our analysis demonstrated that the assay performs well in smear-positive samples (sensitivity 94.4%) but only detects about 44% of cases in smear-negative specimens. This compares with the 67% sensitivity for Xpert MTB/RIF on smear-negative specimens when used as an add-on test [96]. However the smear-negative sensitivity estimate is derived from only four datasets and a univariate analysis that included datasets with patients on treatment provided a higher sensitivity estimate (58%). These estimates must therefore be considered with caution given the substantial remaining heterogeneity observed.

Strengths and limitations of the meta-analysis

Our study includes the largest number of studies (74 studies, 21 225 samples) evaluated as part of a systematic review to assess the diagnostic test accuracy of LPAs for RIF resistance, INH resistance and *M. tuberculosis* detection. The prior WHO-commissioned systematic review on LPAs in 2008 only assessed detection of multidrug resistance without including accuracy for *M. tuberculosis* detection. It also evaluated two assays (INNO-LiPA and Hain MTBDR), which did not detect INH resistance due to *inhA* mutations and are no longer commercially available. Although the majority of datasets included in this study evaluated Hain V1, our analyses included data on Hain V2 and Nipro, which have not previously been reviewed systematically. A recent study demonstrated that the assays are non-inferior in respect to resistance detection, while slightly favouring Hain V2 due to lower indeterminate results on paucibacillary samples [72]. Differences between the index LPAs are therefore more likely to be due to variation in the study populations than due to true differences in accuracy between the tests.

Overall, studies included for the assessment of diagnostic accuracy for RIF and INH resistance were fairly homogeneous. Data from this systematic review reinforces the diagnostic accuracy estimates from the previous systematic review [13] and again demonstrates a greater degree of heterogeneity for INH compared to RIF. This is attributed to INH-resistance mutations being detected in a wider range of genetic loci than for RIF. Other explanations for residual heterogeneity include the predominance of different mutations between datasets due to strain and patient diversity with different mutations being seen in mono-resistant *versus* MDR strains and heteroresistance also being more common in patients that develop resistance on treatment rather than having transmitted resistance [97, 98]. Furthermore predominant mutations have been described both for INH and RIF resistance that differ by geographic locations [10, 17, 99].

Targeted sequencing was only performed in approximately one-third of studies and often only on discrepant results between LPA and phenotypic DST, which limits the validity of the composite reference

standard analysis. Nonetheless, this analysis suggests that decreases in LPA sensitivity are likely due to mutations outside of the probe hotspots or other mechanisms of resistance such as efflux pumps that are also not detected by targeted sequencing. Use of the composite reference standard increased the specificity for RIF and INH resistance detection due to reclassification of some LPA false positives when mutations known to be of clinical significance were detected by LPA but missed by phenotypic DST [100, 101].

Aside from the large number of studies identified, strengths of this study included the use of a standard protocol and predefined analyses using bivariate random-effects models. However, we also recognise some limitations. Despite our comprehensive search strategy, we acknowledge that some relevant studies may have been missed. Publication and selection bias could have resulted in overly optimistic estimates of the diagnostic accuracy of LPAs. Many studies were laboratory based and did not report on the selection criteria of patients. Others included patients that were exposed to MDR-TB patients or already identified as MDR-TB, which may bias the expected pool of mutations detected. If LPAs are utilised more broadly in patients independent of risk factors for drug resistance, then a slightly different pool of mutations and more heteroresistance may impact their sensitivity [97, 98]. Available data for Hain V2 and Nipro were more limited. Further research evaluating the effect of smear status, smear grade and other covariates such as HIV on the diagnostic accuracy of LPAs is needed. A more comprehensive approach to a sequencing reference standard is needed as many studies suffered from bias due to a discrepant analysis. It is also essential for authors of diagnostic accuracy studies to follow the Standards for Reporting Diagnostic accuracy studies (STARD) criteria [102] and QUADAS-2 framework to ensure methodological quality and adequate reporting [103].

Potential clinical and epidemiologic relevance of LPA use in practice

The first pillar of integrated patient-centred care and prevention of the End TB Strategy published by WHO in 2015 [104] states the need for the “early diagnosis of tuberculosis including universal drug susceptibility testing”. This highlights the importance of LPAs for the rapid diagnosis of tuberculosis and multidrug resistance. LPAs are also recommended by the International Standards for Tuberculosis Control (3rd Edition) [105] and represent a widely used assay for the diagnosis of MDR-TB, particularly in settings where there are heightened concerns for INH mono-resistance. This review provides evidence to support the ongoing use of LPAs based on their diagnostic accuracy when used directly on sputum smear-positive specimens or indirectly on culture isolates, as an initial test or in parallel with culture-based DST for the detection of *M. tuberculosis* and multidrug resistance based on data acquired from a range of laboratory settings in different countries. In low- and middle-incidence countries, LPAs may also serve as a critical tool for tuberculosis elimination efforts as part of laboratory surveillance as well as the prompt diagnosis of tuberculosis including MDR-TB in high-risk groups, such as migrants [106–108].

WHO also released recommendations in support of the use of the shorter MDR-TB regimen [109] in May 2016 [110]. This regimen contains kanamycin, high-dose moxifloxacin, prothionamide (or ethionamide), clofazimine, high-dose INH, pyrazinamide and ethambutol, given together in an initial phase of 4–6 months, followed by 5 months of treatment with moxifloxacin, clofazimine, pyrazinamide and ethambutol. Although the guidelines state that INH resistance does not preclude the use of this regimen, the efficacy of high-dose INH in patients with *katG* mutations is unclear [111–114] and is currently the subject of an ongoing clinical trial [115]. Similarly, due to cross-resistance, strains with *inhA* mutations are typically resistant to ethionamide (and also prothionamide) although these patients may benefit from high-dose INH [73]. Therefore patients with *katG* and/or *inhA* mutations may potentially have between one and two fewer active drugs in the regimen. Many patients started on the short MDR-TB regimen will have been diagnosed by Xpert MTB/RIF, which does not detect INH resistance, and therefore knowledge of INH resistance mutations obtained from first line LPAs (as evaluated in this study), while not required, may provide additional valuable information to clinicians, provided it does not delay the start of therapy. Although culture-based DST may also provide these answers, this usually takes several weeks and is not frequently done. This highlights a possible adjunctive role for LPAs in the appropriate early management of MDR-TB [116, 117].

Conclusions

In adults with pulmonary tuberculosis, LPAs demonstrated high accuracy overall for the detection of RIF resistance. LPAs demonstrated high specificity for INH resistance detection with good sensitivity. The accuracy of LPAs for *M. tuberculosis* detection on smear-positive specimens is high, but suboptimal in smear-negative samples. These results were used to inform updated WHO policy recommendations.

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References

- 1 World Health Organization. Global Tuberculosis Report. Geneva, World Health Organization, 2016. Available from: http://www.who.int/tb/publications/global_report/en/
- 2 World Health Organization. Multidrug and Extensively Drug-resistant TB (M/XDR-TB): 2010 Global Report on Surveillance and Response. Geneva, World Health Organization, 2010. Available from: http://apps.who.int/iris/bitstream/10665/44286/1/9789241599191_eng.pdf
- 3 Heifets LB, Cangelosi GA. Drug susceptibility testing of *Mycobacterium tuberculosis*: a neglected problem at the turn of the century. *Int J Tuberc Lung Dis* 1999; 3: 564–581.
- 4 Dinnes J, Deeks J, Kunst H, et al. A systematic review of rapid diagnostic tests for the detection of tuberculosis infection. *Health Technol Assess* 2007; 11: 1–196.
- 5 Small PM, Pai M. Tuberculosis diagnosis: time for a game change. *N Engl J Med* 2010; 363: 1070–1071.
- 6 Ling DI, Zwerling AA, Pai M. Rapid diagnosis of drug-resistant TB using line probe assays: from evidence to policy. *Expert Rev Respir Med* 2008; 2: 583–588.
- 7 Telenti A, Honore N, Bernasconi C, et al. Genotypic assessment of isoniazid and rifampin resistance in *Mycobacterium tuberculosis*: a blind study at reference laboratory level. *J Clin Microbiol* 1997; 35: 719–723.
- 8 Mokrousov I, Narvskaya O, Otten T, et al. High prevalence of KatG Ser315Thr substitution among isoniazid-resistant *Mycobacterium tuberculosis* clinical isolates from northwestern Russia, 1996 to 2001. *Antimicrob Agents Chemother* 2002; 46: 1417–1424.
- 9 Musser JM, Kapur V, Williams DL, et al. Characterization of the catalase-peroxidase gene (katG) and inhA locus in isoniazid-resistant and -susceptible strains of *Mycobacterium tuberculosis* by automated DNA sequencing: restricted array of mutations associated with drug resistance. *J Infect Dis* 1996; 173: 196–202.
- 10 Seifert M, Catanzaro D, Catanzaro A, et al. Genetic mutations associated with isoniazid resistance in *Mycobacterium tuberculosis*: a systematic review. *PLoS One* 2015; 10: e0119628.
- 11 Sreevatsan S, Pan X, Zhang Y, et al. Analysis of the oxyR-ahpC region in isoniazid-resistant and -susceptible *Mycobacterium tuberculosis* complex organisms recovered from diseased humans and animals in diverse localities. *Antimicrob Agents Chemother* 1997; 41: 600–606.
- 12 World Health Organization. WHO policy statement. Molecular line probe assays for rapid screening of patients at risk of multidrug-resistant tuberculosis (English and Russian), 2008. www.who.int/tb/features_archive/policy_statement.pdf?ua=1 Date last accessed: December 22, 2016.
- 13 Ling DI, Zwerling AA, Pai M. GenoType MTBDR assays for the diagnosis of multidrug-resistant tuberculosis: a meta-analysis. *Eur Respir J* 2008; 32: 1165–1174.
- 14 Barnard M, Gey van Pittius NC, van Helden PD, et al. The diagnostic performance of the GenoType MTBDRplus version 2 line probe assay is equivalent to that of the Xpert MTB/RIF assay. *J Clin Microbiol* 2012; 50: 3712–3716.
- 15 Crudu V, Stratan E, Romancenco E, et al. First evaluation of an improved assay for molecular genetic detection of tuberculosis as well as rifampin and isoniazid resistances. *J Clin Microbiol* 2012; 50: 1264–1269.
- 16 Arentz M, Sorensen B, Horne DJ, et al. Systematic review of the performance of rapid rifampicin resistance testing for drug-resistant tuberculosis. *PLoS One* 2013; 8: e76533.
- 17 Mitarai S, Kato S, Ogata H, et al. Comprehensive multicenter evaluation of a new line probe assay kit for identification of *Mycobacterium* species and detection of drug-resistant *Mycobacterium tuberculosis*. *J Clin Microbiol* 2012; 50: 884–890.
- 18 Moher D, Liberati A, Tetzlaff J, et al. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *PLoS Med* 2009; 6: e1000097.
- 19 Cochrane Handbook for Systematic Reviews of Diagnostic Test Accuracy. The Cochrane Collaboration, London, 2008. <http://methods.cochrane.org/sdt/handbook-dta-reviews> Date last accessed: December 22, 2016.
- 20 Whiting PF, Rutjes AW, Westwood ME, et al. QUADAS-2: a revised tool for the quality assessment of diagnostic accuracy studies. *Ann Intern Med* 2011; 155: 529–536.
- 21 Chu H, Cole SR. Bivariate meta-analysis of sensitivity and specificity with sparse data: a generalized linear mixed model approach. *J Clin Epidemiol* 2006; 59: 1331–1332.
- 22 Reitsma JB, Glas AS, Rutjes AW, et al. Bivariate analysis of sensitivity and specificity produces informative summary measures in diagnostic reviews. *J Clin Epidemiol* 2005; 58: 982–990.
- 23 Al-Mutairi NM, Ahmad S, Mokaddas E. Performance comparison of four methods for detecting multidrug-resistant *Mycobacterium tuberculosis* strains. *Int J Tuberc Lung Dis* 2011; 15: 110–115.
- 24 Albert H, Bwanga F, Mukkada S, et al. Rapid screening of MDR-TB using molecular line probe assay is feasible in Uganda. *BMC Infect Dis* 2010; 10: 41.
- 25 Anek-Vorapong R, Sinthuwattanawibool C, Podewils LJ, et al. Validation of the GenoType MTBDRplus assay for detection of MDR-TB in a public health laboratory in Thailand. *BMC Infect Dis* 2010; 10: 123.
- 26 Asante-Poku A, Otchere ID, Danso E, et al. Evaluation of GenoType MTBDRplus for the rapid detection of drug-resistant tuberculosis in Ghana. *Int J Tuberc Lung Dis* 2015; 19: 954–959.
- 27 Asencios L, Galarza M, Quispe N, et al. Molecular test GenoTypeMTBDRplus, an alternative to rapid detection of multidrug resistance tuberculosis. *Revista Peruana de Medicina Experimental y Salud Publica* 2012; 29: 92–98.
- 28 Aung WW, Ei PW, Nyunt WW, et al. Phenotypic and genotypic analysis of anti-tuberculosis drug resistance in *Mycobacterium tuberculosis* isolates in Myanmar. *Ann Lab Med* 2015; 35: 494–499.
- 29 Aurin TH, Munshi SK, Kamal SM, et al. Molecular approaches for detection of the multi-drug resistant tuberculosis (MDR-TB) in Bangladesh. *PLoS One* 2014; 9: e99810.
- 30 Bablishvili N, Tukvadze N, Avaliani Z, et al. A comparison of the Xpert MTB/RIF and GenoType MTBDRplus assays in Georgia. *Int J Tuberc Lung Dis* 2015; 19: 676–678.
- 31 Banu S, Rahman SMM, Khan MSR, et al. Discordance across several methods for drug susceptibility testing of drug-resistant *Mycobacterium tuberculosis* isolates in a single laboratory. *J Clin Microbiol* 2014; 52: 156–163.
- 32 Barnard M, Albert H, Coetzee G, et al. Rapid molecular screening for multidrug-resistant tuberculosis in a high-volume public health laboratory in South Africa. *Am J Respir Crit Care Med* 2008; 177: 787–792.
- 33 Brossier F, Veziris N, Jarlier V, et al. Performance of MTBDR plus for detecting high/low levels of *Mycobacterium tuberculosis* resistance to isoniazid. *Int J Tuberc Lung Dis* 2009; 13: 260–265.

- 34 Bwanga F, Joloba ML, Haile M, *et al.* Evaluation of seven tests for the rapid detection of multidrug-resistant tuberculosis in Uganda. *Int J Tuberc Lung Dis* 2010; 14: 890–895.
- 35 Cabibbe AM, Miotto P, Moure R, *et al.* A lab-on-chip based platform for fast molecular diagnosis of multi-drug resistant tuberculosis. *J Clin Microbiol* 2015.
- 36 Catanzaro A, Rodwell TC, Catanzaro DG, *et al.* Performance comparison of three rapid tests for the diagnosis of drug-resistant tuberculosis. *PLoS One* 2015; 10: e0136861.
- 37 Causse M, Ruiz P, Gutierrez JB, *et al.* Evaluation of new GenoType MTBDRplus for detection of resistance in cultures and direct specimens of *Mycobacterium tuberculosis*. *Int J Tuberc Lung Dis* 2008; 12: 1456–1460.
- 38 Chen C, Kong W, Zhu L, *et al.* Evaluation of the GenoType MTBDRplus line probe assay on sputum-positive samples in routine settings in China. *Int J Tuberc Lung Dis* 2014; 18: 1034–1039.
- 39 Chryssanthou E, Angeby K. The GenoType MTBDRplus assay for detection of drug resistance in *Mycobacterium tuberculosis* in Sweden. *APMIS* 2012; 120: 405–409.
- 40 Daum LT, Rodriguez JD, Worthy SA, *et al.* Next-generation ion torrent sequencing of drug resistance mutations in *Mycobacterium tuberculosis* strains. *J Clin Microbiol* 2012; 50: 3831–3837.
- 41 Dorman SE, Chihota VN, Lewis JJ, *et al.* Genotype MTBDRplus for direct detection of *Mycobacterium tuberculosis* and drug resistance in strains from gold miners in South Africa. *J Clin Microbiol* 2012; 50: 1189–1194.
- 42 Dubois Cauwelaert N, Ramarokoto H, *et al.* DNA extracted from stained sputum smears can be used in the MTBDRplus assay. *J Clin Microbiol* 2011; 49: 3600–3603.
- 43 Eliseev PI, Maryandyshev AO, Nikishova EI, *et al.* Epidemiological analyses of tuberculosis in Archangelsk, Russia and implementation of a rapid assay for detection of resistance in this high burden setting. *Int J Mycobacteriol* 2013; 2: 103–108.
- 44 Evans J, Stead MC, Nicol MP, *et al.* Rapid genotypic assays to identify drug-resistant *Mycobacterium tuberculosis* in South Africa. *J Antimicrob Chemother* 2009; 63: 11–16.
- 45 Fabre M, Hauck Y, Pourcel C, *et al.* Performances of the assay MTBDRplus in the surveillance of rifampicin resistance in *Mycobacterium tuberculosis*. *Pathologie Biologie* 2011; 59: 94–96.
- 46 Farooqi JQ, Khan E, Alam SM, *et al.* Line probe assay for detection of rifampicin and isoniazid resistant tuberculosis in Pakistan. *J Pak Med Assoc* 2012; 62: 767–772.
- 47 Felkel M, Exner R, Schleucher R, *et al.* Evaluation of *Mycobacterium tuberculosis* drug susceptibility in clinical specimens from Nigeria using genotype MTBDRplus and MTBDRsl assays. *Eur J Microbiol Immunol (Bp)* 2013; 3: 252–257.
- 48 Ferro BE, Garcia PK, Nieto LM, *et al.* Predictive value of molecular drug resistance testing of *Mycobacterium tuberculosis* isolates in Valle del Cauca, Colombia. *J Clin Microbiol* 2013; 51: 2220–2224.
- 49 Friedrich SO, Venter A, Kayigire XA, *et al.* Suitability of Xpert MTB/RIF and genotype MTBDRplus for patient selection for a tuberculosis clinical trial. *J Clin Microbiol* 2011; 49: 2827–2831.
- 50 Gauthier M, Somoskovi A, Berland JL, *et al.* Stepwise implementation of a new diagnostic algorithm for multidrug-resistant tuberculosis in Haiti. *Int J Tuberc Lung Dis* 2014; 18: 220–226.
- 51 Gitti Z, Mantadakis E, Maraki S, *et al.* GenoType(R) MTBDRplus compared with conventional drug-susceptibility testing of *Mycobacterium tuberculosis* in a low-resistance locale. *Future Microbiol* 2011; 6: 357–362.
- 52 Hillemann D, Rusch-Gerdes S, Richter E. Evaluation of the GenoType MTBDRplus assay for rifampin and isoniazid susceptibility testing of *Mycobacterium tuberculosis* strains and clinical specimens. *J Clin Microbiol* 2007; 45: 2635–2640.
- 53 Huang FL, Jin JL, Chen S, *et al.* MTBDRplus results correlate with treatment outcome in previously treated tuberculosis patients. *Int J Tuberc Lung Dis* 2015; 19: 319–325.
- 54 Huang WL, Chen HY, Kuo YM, *et al.* Performance assessment of the GenoType MTBDRplus test and DNA sequencing in detection of multidrug-resistant *Mycobacterium tuberculosis*. *J Clin Microbiol* 2009; 47: 2520–2524.
- 55 Huang WL, Hsu ZJ, Chang TC, *et al.* Rapid and accurate detection of rifampin and isoniazid-resistant *Mycobacterium tuberculosis* using an oligonucleotide array. *Clin Microbiol Infect* 2014; 20: O542–O549.
- 56 Huyen MN, Tiemersma EW, Lan NT, *et al.* Validation of the GenoType MTBDRplus assay for diagnosis of multidrug resistant tuberculosis in South Vietnam. *BMC Infect Dis* 2010; 10: 149.
- 57 Imperiale BR, Zumarraga MJ, Weltman G, *et al.* First evaluation in Argentina of the GenoType MTBDRplus assay for multidrug-resistant *Mycobacterium tuberculosis* detection from clinical isolates and specimens. *Rev Argent Microbiol* 2012; 44: 283–289.
- 58 Jin J, Zhang Y, Fan X, *et al.* Evaluation of the GenoType MTBDRplus assay and identification of a rare mutation for improving MDR-TB detection. *Int J Tuberc Lung Dis* 2012; 16: 521–526.
- 59 Kapata N, Mbulo G, Cobelens F, *et al.* The Second Zambian National Tuberculosis Drug Resistance survey: a comparison of conventional and molecular methods. *Trop Med Int Health* 2015; 20: 1492–1500.
- 60 Khadka JB, Rai SK, Shrestha S, *et al.* Study of rifampicin and isoniazid resistance mutation genes of *M. tuberculosis* isolates in Nepal. *Nepal Med Coll J* 2011; 13: 147–151.
- 61 Kumar P, Balooni V, Sharma BK, *et al.* High degree of multi-drug resistance and hetero-resistance in pulmonary TB patients from Punjab state of India. *Tuberculosis (Edinb)* 2014; 94: 73–80.
- 62 Lacombe A, Garcia-Sierra N, Prat C, *et al.* GenoType MTBDRplus assay for molecular detection of rifampin and isoniazid resistance in *Mycobacterium tuberculosis* strains and clinical samples. *J Clin Microbiol* 2008; 46: 3660–3667.
- 63 Li Q, Dong HY, Pang Y, *et al.* Multicenter evaluation of the molecular line probe assay for multidrug resistant *Mycobacterium tuberculosis* detection in China. *Biomed Environ Sci* 2015; 28: 464–467.
- 64 Luetkemeyer AF, Kendall MA, Wu X, *et al.* Evaluation of two line probe assays for rapid detection of *Mycobacterium tuberculosis*, tuberculosis (TB) drug resistance, and non-TB Mycobacteria in HIV-infected individuals with suspected TB. *J Clin Microbiol* 2014; 52: 1052–1059.
- 65 Lyu J, Kim MN, Song JW, *et al.* GenoType MTBDRplus assay detection of drug-resistant tuberculosis in routine practice in Korea. *Int J Tuberc Lung Dis* 2013; 17: 120–124.
- 66 Macedo R, Amorim A, Pereira E. Multidrug-resistant tuberculosis: rapid molecular detection with MTBDRplus assay in clinical samples. *Rev Port Pneumol* 2009; 15: 353–365.

- 67 Maschmann Rde A, Sa Spies F, Nunes, *et al.* Performance of the GenoType MTBDRplus assay directly on sputum specimens from Brazilian patients with tuberculosis treatment failure or relapse. *J Clin Microbiol* 2013; 51: 1606–1608.
- 68 Miotto P, Piana F, Cirillo DM, *et al.* Genotype MTBDRplus: a further step toward rapid identification of drug-resistant *Mycobacterium tuberculosis*. *J Clin Microbiol* 2008; 46: 393–394.
- 69 Miotto P, Saleri N, Dembele M, *et al.* Molecular detection of rifampin and isoniazid resistance to guide chronic TB patient management in Burkina Faso. *BMC Infect Dis* 2009; 9: 142.
- 70 Mironova S, Pimkina E, Kontsevaya I, *et al.* Performance of the GenoType MTBDRplus assay in routine settings: a multicenter study. *Eur J Clin Microbiol Infect Dis* 2012; 31: 1381–1387.
- 71 N'Guessan K, Assi JS, Ouassa T, *et al.* Assessment of the genotype MTBDRplus assay for rifampin and isoniazid resistance detection on sputum samples in Cote d'Ivoire. *Eur J Microbiol Immunol (Bp)* 2014; 4: 166–173.
- 72 Nathavitharana RR, Hillemann D, Schumacher SG, *et al.* Multicenter noninferiority evaluation of Hain GenoType MTBDRplus Version 2 and Nipro NTM+MDRTB line probe assays for detection of rifampin and isoniazid resistance. *J Clin Microbiol* 2016; 54: 1624–1630.
- 73 Niehaus AJ, Mlisana K, Gandhi NR, *et al.* High prevalence of inhA promoter mutations among patients with drug-resistant tuberculosis in KwaZulu-Natal, South Africa. *PLoS One* 2015; 10: e0135003.
- 74 Nikolayevskiy V, Balabanova Y, Simak T, *et al.* Performance of the Genotype MTBDRplus assay in the diagnosis of tuberculosis and drug resistance in Samara, Russian Federation. *BMC Clin Pathol* 2009; 9: 2.
- 75 Nwofor AC, Nyamngee A, Nwabuisi C, *et al.* Performance of Genotype MTBDRplus in the detection of resistance to rifampicin and isoniazid among clinical mycobacteria isolates in Ilorin, Nigeria. *Curr HIV Res* 2015; 13: 308–314.
- 76 Ocheretina O, Escuyer VE, Mabou MM, *et al.* Correlation between genotypic and phenotypic testing for resistance to rifampin in *Mycobacterium tuberculosis* clinical isolates in Haiti. Investigation of cases with discrepant susceptibility results. *Plos One* 2014; 9: e90569.
- 77 Raizada N, Sachdeva KS, Chauhan DS, *et al.* A multi-site validation in India of the line probe assay for the rapid diagnosis of multi-drug resistant tuberculosis directly from sputum specimens. *PLoS One* 2014; 9: e88626.
- 78 Raveendran R, Wattal C, Oberoi JK, *et al.* Utility of GenoType MTBDRplus assay in rapid diagnosis of multidrug resistant tuberculosis at a tertiary care centre in India. *Indian J Med Microbiol* 2012; 30: 58–63.
- 79 Rienthong S, Boonin C, Chaiyasinrote B, *et al.* Evaluation of a novel line-probe assay for genotyping-based diagnosis of *Mycobacterium tuberculosis* in Thailand. *Int J Tuberc Lung Dis* 2015; 19: 817–822.
- 80 Rigouts L, Hoza AS, De Rijk P, *et al.* Evaluation of the Genotype MTBDRplus assay as a tool for drug resistance surveys. *Int J Tuberc Lung Dis* 2011; 15: 959–965.
- 81 Rufai SB, Kumar P, Singh A, *et al.* Comparison of Xpert MTB/RIF with line probe assay for detection of rifampin-monoresistant *Mycobacterium tuberculosis*. *J Clin Microbiol* 2014; 52: 1846–1852.
- 82 Sangsayunh P, Chuchothawon C. Genotype MTBDR plus (Hain) test in suspected MDR-TB patients. *J Med Assoc Thai* 2014; 97: 1028–1032.
- 83 Schon T, Jureen P, Chryssanthou E, *et al.* Rifampicin-resistant and rifabutin-susceptible *Mycobacterium tuberculosis* strains: a breakpoint artefact? *J Antimicrob Chemother* 2013; 68: 2074–2077.
- 84 Scott LE, McCarthy K, Gous N, *et al.* Comparison of Xpert MTB/RIF with other nucleic acid technologies for diagnosing pulmonary tuberculosis in a high HIV prevalence setting: a prospective study. *PLoS Med* 2011; 8: e1001061.
- 85 Shubladeze N, Tadumadze N, Bablshvili N. Molecular patterns of multidrug resistance of in Georgia. *Int J Mycobacteriol* 2013; 2: 73–78.
- 86 Simons SO, van der Laan T, de Zwaan R, *et al.* Molecular drug susceptibility testing in the Netherlands: performance of the MTBDRplus and MTBDRsl assays. *Int J Tuberc Lung Dis* 2015; 19: 828–833.
- 87 Singhal R, Arora J, Lal P, *et al.* Comparison of line probe assay with liquid culture for rapid detection of multi-drug resistance in *Mycobacterium tuberculosis*. *Indian J Med Res* 2012; 136: 1044–1047.
- 88 Tessema B, Beer J, Emmrich F, *et al.* Analysis of gene mutations associated with isoniazid, rifampicin and ethambutol resistance among *Mycobacterium tuberculosis* isolates from Ethiopia. *BMC Infect Dis* 2012; 12: 37.
- 89 Tho DQ, Lan NT, Chau NV, *et al.* Multiplex allele-specific polymerase chain reaction for detection of isoniazid resistance in *Mycobacterium tuberculosis*. *Int J Tuberc Lung Dis* 2011; 15: 799–803.
- 90 Tolani MP, D'Souza DT, Mistry NF. Drug resistance mutations and heteroresistance detected using the GenoType MTBDRplus assay and their implication for treatment outcomes in patients from Mumbai, India. *BMC Infect Dis* 2012; 12: 9.
- 91 Tukvadze N, Kempker RR, Kalandadze I, *et al.* Use of a molecular diagnostic test in AFB smear positive tuberculosis suspects greatly reduces time to detection of multidrug resistant tuberculosis. *PLoS One* 2012; 7: e31563.
- 92 Vijdea R, Stegger M, Sosnovskaja A, *et al.* Multidrug-resistant tuberculosis: rapid detection of resistance to rifampin and high or low levels of isoniazid in clinical specimens and isolates. *Eur J Clin Microbiol Infect Dis* 2008; 27: 1079–1086.
- 93 Yadav RN, Singh BK, Sharma SK, *et al.* Comparative evaluation of GenoType MTBDRplus line probe assay with solid culture method in early diagnosis of multidrug resistant tuberculosis (MDR-TB) at a tertiary care centre in India. *PLoS One* 2013; 8: e72036.
- 94 Yordanova S, Bachiyska E, Atanasova Y, *et al.* Multidrug resistant tuberculosis in Bulgaria—at microbiological aspects. *Probl Infect Parasit Dis* 2013; 41: 5–8.
- 95 Smith SE, Kurbatova EV, Cavanaugh JS, *et al.* Global isoniazid resistance patterns in rifampin-resistant and rifampin-susceptible tuberculosis. *Int J Tuberc Lung Dis* 2012; 16: 203–205.
- 96 Steingart KR, Schiller I, Horne DJ, *et al.* Xpert MTB/RIF assay for pulmonary tuberculosis and rifampicin resistance in adults. *Cochrane Database Syst Rev* 2014; 1: CD009593.
- 97 Gillespie SH. Evolution of drug resistance in *Mycobacterium tuberculosis*: clinical and molecular perspective. *Antimicrob Agents Chemother* 2002; 46: 267–274.
- 98 Muller B, Borrell S, Rose G, *et al.* The heterogeneous evolution of multidrug-resistant *Mycobacterium tuberculosis*. *Trends Genet* 2013; 29: 160–169.
- 99 Sanchez-Padilla E, Merker M, Beckert P, *et al.* Detection of drug-resistant tuberculosis by Xpert MTB/RIF in Swaziland. *N Engl J Med* 2015; 372: 1181–1182.

- 100 Van Deun A, Aung KJ, Bola V, *et al.* Rifampin drug resistance tests for tuberculosis: challenging the gold standard. *J Clin Microbiol* 2013; 51: 2633–2640.
- 101 Van Deun A, Aung KJ, Hossain A, *et al.* Disputed rpoB mutations can frequently cause important rifampicin resistance among new tuberculosis patients. *Int J Tuberc Lung Dis* 2015; 19: 185–190.
- 102 Bossuyt PM, Reitsma JB, Bruns DE, *et al.* STARD 2015: an updated list of essential items for reporting diagnostic accuracy studies. *BMJ* 2015; 351: h5527.
- 103 Fontela PS, Pant Pai N, Schiller I, *et al.* Quality and reporting of diagnostic accuracy studies in TB, HIV and malaria: evaluation using QUADAS and STARD standards. *PLoS One* 2009; 4: e7753–e7700.
- 104 Uplekar M, Weil D, Lonnroth K, *et al.* WHO's new end TB strategy. *Lancet* 2015; 385: 1799–1801.
- 105 TB CARE I. International Standards for Tuberculosis Care. 3rd Edn. The Hague, TB CARE I, 2014.
- 106 Lonnroth K, Migliori GB, Abubakar I, *et al.* Towards tuberculosis elimination: an action framework for low-incidence countries. *Eur Respir J* 2015; 45: 928–952.
- 107 Rendon A, Fuentes Z, Torres-Duque CA, *et al.* Roadmap for tuberculosis elimination in Latin American and Caribbean countries: a strategic alliance. *Eur Respir J* 2016; 48: 1282–1287.
- 108 D'Ambrosio L, Dara M, Tadolini M, *et al.* Tuberculosis elimination: theory and practice in Europe. *Eur Respir J* 2014; 43: 1410–1420.
- 109 Van Deun A, Maug AK, Salim MA, *et al.* Short, highly effective, and inexpensive standardized treatment of multidrug-resistant tuberculosis. *Am J Respir Crit Care Med* 2010; 182: 684–692.
- 110 World Health Organization. WHO treatment guidelines for drug-resistant tuberculosis. 2016 update WHO/HTM/TB/2016.04. World Health Organization, Geneva, 2016. Available from: www.who.int/tb/MDRTBguidelines2016.pdf
- 111 Caminero JA, Sotgiu G, Zumla A, *et al.* Best drug treatment for multidrug-resistant and extensively drug-resistant tuberculosis. *Lancet Infect Dis* 2010; 10: 621–629.
- 112 Jacobson KR, Theron D, Victor TC, *et al.* Treatment outcomes of isoniazid-resistant tuberculosis patients, Western Cape Province, South Africa. *Clin Infect* 2011; 53: 369–372.
- 113 Vilcheze C, Jacobs WR Jr. The mechanism of isoniazid killing: clarity through the scope of genetics. *Annu Rev Microbiol* 2007; 61: 35–50.
- 114 Warren RM, Streicher EM, Gey van Pittius NC, *et al.* The clinical relevance of Mycobacterial pharmacogenetics. *Tuberculosis (Edinb)* 2009; 89: 199–202.
- 115 High-Dose Isoniazid Among Adult Patients With Different Genetic Variants of INH-Resistant Tuberculosis (TB). <https://clinicaltrials.gov/ct2/show/NCT0193683>. Date last accessed August 25, 2016.
- 116 Sotgiu G, Tiberi S, D'Ambrosio L, *et al.* Faster for less, the new 'Shorter' regimen for multidrug-resistant tuberculosis. *Eur Respir J* 2016; 48: 1503–1507.
- 117 Sotgiu G, Tiberi S, D'Ambrosio L, *et al.* WHO recommendations on shorter treatment of multidrug-resistant tuberculosis. *Lancet* 2016; 387: 2486–2487.