Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV (Review)

Bjerrum S, Schiller I, Dendukuri N, Kohli M, Nathavitharana RR, Zwerling AA, Denkinger CM, Steingart KR, Shah M

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Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV (Review)
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Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV

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ABSTRACT

Background
The lateral flow urine lipoarabinomannan (LF-LAM) assay Alere Determine™ TB LAM Ag is recommended by the World Health Organization (WHO) to help detect active tuberculosis in HIV-positive people with severe HIV disease. This review update asks the question, "does new evidence justify the use of LF-LAM in a broader group of people?", and is part of the WHO process for updating guidance on the use of LF-LAM.

Objectives
To assess the accuracy of LF-LAM for the diagnosis of active tuberculosis among HIV-positive adults with signs and symptoms of tuberculosis (symptomatic participants) and among HIV-positive adults irrespective of signs and symptoms of tuberculosis (unselected participants not assessed for tuberculosis signs and symptoms).

The proposed role for LF-LAM is as an add on to clinical judgement and with other tests to assist in diagnosing tuberculosis.

Search methods
We searched the Cochrane Infectious Diseases Group Specialized Register; MEDLINE, Embase, Science Citation Index, Web of Science, Latin American Caribbean Health Sciences Literature, Scopus, the WHO International Clinical Trials Registry Platform, the International Standard Randomized Controlled Trial Number Registry, and ProQuest, without language restriction to 11 May 2018.
Selection criteria
Randomized trials, cross-sectional, and observational cohort studies that evaluated LF-LAM for active tuberculosis (pulmonary and extrapulmonary) in HIV-positive adults. We included studies that used the manufacturer’s recommended threshold for test positivity, either the updated reference card with four bands (grade 1 of 4) or the corresponding prior reference card grade with five bands (grade 2 of 5). The reference standard was culture or nucleic acid amplification test from any body site (microbiological). We considered a higher quality reference standard to be one in which two or more specimen types were evaluated for tuberculosis diagnosis and a lower quality reference standard to be one in which only one specimen type was evaluated.

Data collection and analysis
Two review authors independently extracted data using a standardized form and REDCap electronic data capture tools. We appraised the quality of studies using the Quality Assessment of Diagnostic Accuracy Studies-2 (QUADAS-2) tool and performed meta-analyses to estimate pooled sensitivity and specificity using a bivariate random-effects model and a Bayesian approach. We analyzed studies enrolling strictly symptomatic participants separately from those enrolling unselected participants. We investigated pre-defined sources of heterogeneity including the influence of CD4 count and clinical setting on the accuracy estimates. We assessed the certainty of the evidence using the GRADE approach.

Main results
We included 15 unique studies (nine new studies and six studies from the original review that met the inclusion criteria): eight studies among symptomatic adults and seven studies among unselected adults. All studies were conducted in low- or middle-income countries. Risk of bias was high in the participant selection and reference standard domains, mainly because studies excluded participants unable to produce sputum and used a lower quality reference standard.

Participants with tuberculosis symptoms
LF-LAM pooled sensitivity (95% credible interval (CrI) ) was 42% (31% to 55%) (moderate-certainty evidence) and pooled specificity was 91% (85% to 95%) (very low-certainty evidence), (8 studies, 3,449 participants, 37% with tuberculosis).

For a population of 1000 people where 300 have microbiologically-confirmed tuberculosis, the utilization of LF-LAM would result in: 189 to be LF-LAM positive: of these, 63 (33%) would not have tuberculosis (false-positives); and 811 to be LF-LAM negative: of these, 174 (21%) would have tuberculosis (false-negatives).

By clinical setting, pooled sensitivity was 52% (40% to 64%) among inpatients versus 29% (17% to 47%) among outpatients; and pooled specificity was 87% (78% to 93%) among inpatients versus 96% (91% to 99%) among outpatients. Stratified by CD4 cell count, pooled sensitivity increased, and specificity decreased with lower CD4 cell count.

Unselected participants not assessed for signs and symptoms of tuberculosis
LF-LAM pooled sensitivity was 35% (22% to 50%), (moderate-certainty evidence) and pooled specificity was 95% (89% to 96%), (low-certainty evidence), (7 studies, 3365 participants, 13% with tuberculosis).

For a population of 1000 people where 100 have microbiologically-confirmed tuberculosis, the utilization of LF-LAM would result in: 80 to be LF-LAM positive: of these, 45 (56%) would not have tuberculosis (false-positives); and 920 to be LF-LAM negative: of these, 65 (7%) would have tuberculosis (false-negatives).

By clinical setting, pooled sensitivity was 62% (41% to 83%) among inpatients versus 31% (18% to 47%) among outpatients; pooled specificity was 84% (48% to 96%) among inpatients versus 95% (87% to 99%) among outpatients. Stratified by CD4 cell count, pooled sensitivity increased, and specificity decreased with lower CD4 cell count.

Authors’ conclusions
We found that LF-LAM has a sensitivity of 42% to diagnose tuberculosis in HIV-positive individuals with tuberculosis symptoms and 35% in HIV-positive individuals not assessed for tuberculosis symptoms, consistent with findings reported previously. Regardless of how people are enrolled, sensitivity is higher in inpatients and those with lower CD4 cell, but a concomitant lower specificity. As a simple point-of-care test that does not depend upon sputum evaluation, LF-LAM may assist with the diagnosis of tuberculosis, particularly when a sputum specimen cannot be produced.

17 October 2019
Up to date
All studies incorporated from most recent search
All studies identified during the most recent search (11 May, 2018) have been incorporated in the review, and no ongoing studies identified.
**Plain Language Summary**

**Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV**

Why is improving the diagnosis of tuberculosis important?

Tuberculosis causes more deaths in people living with HIV than any other disease. The lateral flow urine lipoarabinomannan assay (LF-LAM, Alere Determine™ TB LAM Ag assay) is a World Health Organization-recommended rapid test to assist in detection of active tuberculosis in HIV-positive people with severe HIV disease. Rapid and early tuberculosis diagnosis may allow for prompt treatment and alleviate severe illness and death. An incorrect tuberculosis diagnosis may result in anxiety and unnecessary treatment.

What is the aim of this review?

To find out how accurate LF-LAM is for diagnosing tuberculosis in HIV-positive people with tuberculosis symptoms (symptomatic participants) and those not assessed for tuberculosis symptoms (unselected participants). This is an update of the 2016 Cochrane Review.

What was studied in this review?

LF-LAM is a commercially available point-of-care test that detects lipoarabinomannan (LAM), a component of the bacterial cell walls, present in some people with active tuberculosis. The test is simple and shows results in 25 minutes. LF-LAM results were measured against culture or molecular tests (benchmark).

What are the main results of this review?

Fifteen studies: eight studies evaluated LF-LAM for tuberculosis among symptomatic participants and seven studies among unselected participants. All studies were conducted in low- or middle-income countries.

Tuberculosis diagnosis among symptomatic participants: LF-LAM registered positive in 42% (sensitivity) of people who actually had tuberculosis and did not register positive in 91% of people who were actually negative (specificity).

Tuberculosis diagnosis among unselected participants: LF-LAM sensitivity was 35% and specificity 95%.

How confident are we in the review’s results?

Several studies excluded participants who could not produce sputum and most studies relied on a lower quality benchmark. Few studies and participants were included in some analyses and only one study was conducted outside of sub-Saharan Africa. Results should be interpreted with caution.

What do the results mean?

Among symptomatic participants, in theory, for a population of 1000 people where 300 have microbiologically-confirmed tuberculosis, the utilization of LF-LAM would result in: 189 to be LF-LAM positive: of these, 63 (33%) would not have tuberculosis (false-positives); and 811 to be LF-LAM negative: of these, 174 (21%) would have tuberculosis (false-negatives).

Among unselected participants, in theory, for a population of 1000 people where 100 have microbiologically-confirmed tuberculosis, the utilization of LF-LAM would result in: 80 to be LF-LAM positive: of these, 45 (56%) would not have tuberculosis (false-positives); and 920 to be LF-LAM negative: of these, 65 (7%) would have tuberculosis (false-negatives).

Who do the review’s results apply to?

HIV-positive people with tuberculosis symptoms and those not assessed for tuberculosis symptoms.

What are the implications of this review?

LF-LAM has sensitivity around 40% to detect tuberculosis. As the test does not require sputum collection, LF-LAM may be the only way to diagnose tuberculosis when sputum cannot be produced.

How up-to-date is this review?

To 11 May 2018.
**SUMMARY OF FINDINGS**

**Summary of findings 1. LF-LAM for symptomatic participants**

Review question: what is the diagnostic accuracy of LF-LAM for the diagnosis of active tuberculosis in HIV-positive adults with signs and symptoms of tuberculosis?

Studies: cross-sectional studies and randomized controlled trials

Participants: HIV-positive adults with tuberculosis signs and symptoms

Setting: all settings (inpatient and outpatient)

Index test: LF-LAM

Threshold for index tests: manufacturer’s recommended threshold for positivity i.e. grade 1 of 4 (revised reference card) or the corresponding grade 2 of 5 (prior reference card)

Reference standard: microbiological (mycobacterial culture and/or nucleic acid amplification test)

Role: an add on to clinical judgement and with other tests to assist in tuberculosis diagnosis

Pooled sensitivity (95% CrI): 42% (31% to 55%); pooled specificity (95% CrI): 91% (85% to 95%)

| Test result            | Number of results per 1000 participants tested (95% CrI) | Number of participants (studies) | Certainty of the evidence (GRADE) |
|------------------------|-----------------------------------------------------------|----------------------------------|-----------------------------------|
| True positives         |                                                           |                                  |                                   |
| Prevalence 1%          | 4 (3 to 6)                                                 |                                  |                                   |
| False negatives        | 6 (4 to 7)                                                 |                                  |                                   |
| True negatives         | 901 (842 to 941)                                           |                                  |                                   |
| False positives        | 89 (49 to 148)                                             |                                  |                                   |

| Prevalence 10%         | 42 (31 to 55)                                              |                                  |                                   |
| False negatives        | 58 (45 to 69)                                              |                                  |                                   |
| True negatives         | 819 (765 to 855)                                           |                                  |                                   |
| False positives        | 81 (45 to 135)                                             |                                  |                                   |

| Prevalence 30%         | 126 (93 to 165)                                            |                                  |                                   |
| False negatives        | 174 (135 to 207)                                           |                                  |                                   |
| True negatives         | 637 (595 to 665)                                           |                                  |                                   |
| False positives        | 63 (35 to 105)                                             |                                  |                                   |

| Prevalence 30%         | 1277 (8)                                                   |                                  | MODERATE a,b,c,d                   |
| False negatives        | 58 (45 to 69)                                              |                                  |                                   |
| True negatives         | 2172 (8)                                                   |                                  | VERY LOW c,e,f                     |

| Prevalence 30%         | 1277 (8)                                                   |                                  |                                    |
| False negatives        | 174 (135 to 207)                                           |                                  |                                    |
| True negatives         | 2172 (8)                                                   |                                  |                                    |

| Prevalence 30%         | 1277 (8)                                                   |                                  |                                    |
| False negatives        | 174 (135 to 207)                                           |                                  |                                    |
| True negatives         | 2172 (8)                                                   |                                  |                                    |

Abbreviations: CrI: credible interval.

Explanations

\(^{a}\)As assessed by QUADAS-2, in the patient selection domain, we judged six studies (75%) at high risk of bias because they did not avoid inappropriate exclusions. We downgraded one level for risk of bias.

\(^{b}\)For individual studies, sensitivity estimates ranged from 23% to 68%. We thought that differences in enrolment criteria (different populations targeted) or CD4 count could explain in part the heterogeneity. We did not downgrade for inconsistency.
The median tuberculosis prevalence in the studies was 42% and thus the results tend to be more applicable to settings with a higher tuberculosis prevalence. For tuberculosis prevalence of 1% and 10%, whether or not to downgrade remains unclear. It is possible the test will perform differently at lower prevalences. We did not downgrade for indirectness.

We thought the 95% CrIs around true positives and false negatives would likely not lead to different decisions depending on which credible limits are assumed. We did not downgrade for imprecision.

As assessed by QUADAS-2, in the reference standard domain, we judged seven studies (88%) at high risk of bias because we thought the reference standard used was unlikely to correctly classify the target condition. We downgraded two levels for risk of bias.

We thought the 95% CrIs around true negatives and false positives would likely lead to different decisions depending on which credible limits are assumed. We downgraded one level for imprecision.

**GRADE certainty of evidence** (GRADEpro 2015; Balsch 2011)

*High:* we are very confident that the true effect lies close to that of the estimate of the effect.

*Moderate:* we are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

*Low:* our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect.

*Very low:* we have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect.

The table displays normalized frequencies within a hypothetical cohort of 1000 patients at three different prevalences of tuberculosis (pre-test probabilities): 1%, 10% and 30%. Credible limits (CrIs) were estimated based on those around the point estimates for pooled sensitivity and specificity.

**Note:** the results on this table should not be interpreted in isolation from the results of the individual included studies contributing to each summary test accuracy measure. These are reported in the main body of the text of the review.

### Summary of findings 2. LF-LAM for unselected participants

**Review question:** what is the diagnostic accuracy of LF-LAM for the diagnosis of active tuberculosis in HIV-positive adults irrespective of signs and symptoms of tuberculosis?

**Studies:** cross-sectional studies and randomized controlled trials

**Participants:** HIV-positive unselected adults not assessed for signs and symptoms of tuberculosis i.e. irrespective of symptoms

**Setting:** all settings (inpatient and outpatient)

**Index test:** LF-LAM

**Threshold for index tests:** manufacturer’s recommended threshold for positivity i.e. grade 1 of 4 (revised reference card) or the corresponding grade 2 of 5 (prior reference card)

**Reference standard:** microbiological (mycobacterial culture and/or nucleic acid amplification test)

**Role:** an add on to clinical judgement and with other tests to assist in tuberculosis diagnosis

**Pooled sensitivity (95% CrI):** 35% (22% to 50%); **pooled specificity (95% CrI):** (89% to 98%)
| Test result            | Number of results per 1000 participants tested (95% CrI) | Number of participants (studies) | Certainty of the evidence (GRADE) |
|-----------------------|-------------------------------------------------------|---------------------------------|----------------------------------|
|                       | Prevalence 1%                                         |                                 |                                  |
|                       | Typically seen in asymptomatic persons in outpatient  |                                 |                                  |
|                       | settings                                             |                                 |                                  |
|                       | True positives                                       | 3 (2 to 5)                       | MODERATEa,b,c                     |
|                       | False negatives                                       | 7 (5 to 8)                       |                                  |
|                       | True negatives                                        | 941 (881 to 970)                 | LOWd,e,f                         |
|                       | False positives                                       | 49 (20 to 109)                   |                                  |
|                       | Prevalence 10%                                        | 35 (22 to 50)                    |                                  |
|                       | Typically seen in symptomatic persons in all settings |                                 |                                  |
|                       | True positives                                       | 105 (66 to 150)                  |                                  |
|                       | False negatives                                       | 65 (50 to 78)                    |                                  |
|                       | True negatives                                        | 855 (801 to 882)                 |                                  |
|                       | False positives                                       | 45 (18 to 99)                    |                                  |
|                       | Prevalence 30%                                        | 35 (14 to 77)                    |                                  |
|                       | Typically seen in seriously ill persons in inpatient  |                                 |                                  |
|                       | settings                                              |                                 |                                  |
|                       | True positives                                       |                                  |                                  |
|                       | False negatives                                       |                                  |                                  |
|                       | True negatives                                        |                                  |                                  |
|                       | False positives                                       |                                  |                                  |

**Abbreviations:** CrI: Credible interval.

**Explanations**

a As assessed by QUADAS-2, in the patient selection domain, we judged four studies (57%) at high risk of bias because they did not avoid inappropriate exclusions. We downgraded one level for risk of bias.

b For individual studies, sensitivity ranged from 0% to 67%. We thought that the percentage of patients with tuberculosis symptoms or differences in CD4 count could explain in part the heterogeneity. One study, LaCourse 2016, with sensitivity of 0% differed from the other studies by including a) a population of exclusively pregnant women attending an antenatal care setting, b) a low proportion of symptomatic participants (19%), c) a low tuberculosis prevalence (1%), and d) a high median CD4 cell count (437 cells per μL). One study, Thit 2017, with sensitivity 67% differed from the other studies by being conducted in Myanmar, and was the only study included in this review that evaluated LF-LAM in a setting outside sub-Saharan Africa. We did not downgrade for inconsistency.

c We thought the 95% CrI around true positives and false negatives would likely not lead to different decisions depending on which credible limits are assumed. We did not downgrade for imprecision.

d As assessed by QUADAS-2, in the reference standard domain, we judged five studies (71%) at high risk of bias because we thought the reference standard used was unlikely to correctly classify the target condition. We downgraded one level for risk of bias.

e For individual studies, specificity ranged from 67% to 99%. Eighty-six per cent (6/7) of the included studies had specificity of 94% or higher. One study, Thit 2017, with specificity 67% differed from the other studies by being conducted in Myanmar, and is the only study included in this review that evaluated LF-LAM in a setting outside sub-Saharan Africa. We did not downgrade for inconsistency.

f We thought the wide 95% Crls around true negatives and false positives would lead to different decisions depending on which credible limits are assumed. We downgraded one level for imprecision.

**GRADE certainty of evidence (GRADEpro 2015; Balshem 2011)**

High: we are very confident that the true effect lies close to that of the estimate of the effect.

Moderate: we are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

Low: our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect.

Very low: we have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of the effect.

The table displays normalized frequencies within a hypothetical cohort of 1000 patients at three different prevalences of tuberculosis (pre-test probabilities): 1%, 10% and 30%. Credible limits (CrIs) were estimated based on those around the point estimates for pooled sensitivity and specificity.
Note: the results on this table should not be interpreted in isolation from the results of the individual included studies contributing to each summary test accuracy measure. These are reported in the main body of the text of the review.
**BACKGROUND**

Tuberculosis is an infectious airborne disease caused by the bacillus *Mycobacterium tuberculosis* (*M. tuberculosis*). In 2017, an estimated 10.0 million people developed tuberculosis disease of which 920,000 (9%) occurred among people living with HIV (WHO Global Tuberculosis Report 2018). Although the number of tuberculosis deaths has fallen since 2000, tuberculosis was responsible for 1.6 million deaths in 2017 (WHO Global Tuberculosis Report 2018), and tuberculosis has surpassed HIV as the world’s leading cause of death from an infectious disease. Among people living with HIV, tuberculosis is the major cause of hospitalisation and in-hospital death despite increased access to antiretroviral treatment (ART) (Ford 2016). A systematic review of the prevalence of tuberculosis identified at autopsy in resource-limited settings, suggests that tuberculosis is responsible for up to 40% of all HIV-related deaths and that tuberculosis often is disseminated and undiagnosed at the time of death (Gupta 2015). Globally in 2017, only 51% of the incident tuberculosis cases were reported among people living with HIV (WHO Global Tuberculosis Report 2018). However, most deaths from tuberculosis are preventable if the disease is detected early and effectively treated. Overall, it is estimated that 45 million lives were saved between 2000 and 2017 through effective diagnosis and treatment (WHO Global Tuberculosis Report 2018).

Geographically, HIV and tuberculosis are often concentrated in areas of poverty with limited resources for diagnosis, treatment, and prevention of tuberculosis. Most of the 30 high tuberculosis/HIV burden countries are situated in sub-Saharan Africa where HIV-infection represents a major driver of the tuberculosis epidemic (WHO Global Tuberculosis Report 2018). HIV-positive individuals have a 20- to 37-fold increased risk of developing tuberculosis compared to HIV-negative individuals (Getahun 2010). The risk of tuberculosis increases with decreasing CD4 cell count, and remains elevated throughout the course of HIV (Gupta 2012; Lawn 2009).

Tuberculosis predominantly affects the lungs (pulmonary tuberculosis), but can affect most parts of the body, such as the lymph nodes, pleura, brain, or spine (extrapulmonary tuberculosis). Extrapulmonary tuberculosis represented 14% of the incident tuberculosis cases notified in 2017 (WHO Global Tuberculosis Report 2018). In comparison with HIV-negative people, HIV-positive people have higher rates of extrapulmonary tuberculosis or mycobacteraemia (tuberculosis bloodstream infection) (Pai 2016; Shrivakoti 2017). Signs and symptoms of tuberculosis in people living with HIV vary depending on the progression of immunodeficiency and involved site(s). Complaints are often non-specific such as fever, weight loss, and fatigue. A cough for longer than two weeks is a common distinguishing feature of tuberculosis that prompts diagnostic testing for tuberculosis in people who are HIV negative, but is present in less than a third of people with tuberculosis who are HIV positive (Cain 2010). Similarly, radiographic features of tuberculosis in people living with HIV may be misleading or atypical. Whereas upper lobe cavitory lesions are often seen in HIV-negative people with tuberculosis, such lesions are less common in HIV-positive people with tuberculosis (Cain 2010). Identifying those who warrant further testing for tuberculosis can therefore be challenging. Tuberculosis diagnosis further relies heavily on a sputum-based test strategy that may have limited accuracy as well as applicability among a population that may be sputum scarce and frequently has extrapulmonary and disseminated tuberculosis disease. Moreover, people living with HIV often have paucibacillary tuberculosis disease that makes diagnosis of tuberculosis by smear microscopy, nucleic acid amplification tests (NAATs), and culture less sensitive.

Non-sputum-based point-of-care tuberculosis diagnostic tests are highly desired to improve tuberculosis case detection in people living with HIV and ensure timely treatment (WHO TTP 2014). Desired characteristics of such a test would include minimal or non-invasive sample collection, short time to result (under one hour), and ability to implement the test without need for special instruments, electricity, or specimen preparation (WHO TTP 2014).

Detection of mycobacterial antigen in urine has attracted great attention over time. Urine-based antigen testing would allow for a tuberculosis diagnosis that is non-site specific. In addition, urine is easy to collect and store, and lacks the infection control risks associated with sputum collection. Multiple platforms have been developed to detect lipoarabinomannan (LAM), a component of mycobacterial cell walls, initially as enzyme-linked immunosorbent (ELISA) assays that were evaluated in several clinical settings (Minion 2011). Later, the lateral flow urine lipoarabinomannan (LF-LAM, Alere Determine TB-LAM assay) was developed as a simple point-of-care test for diagnosis of active tuberculosis in people living with HIV. The test is commercially available, does not require access to special laboratory equipment, and produces a result after 25 minutes (Alere 2017), meeting many of the desired target product-profile requirements (WHO TTP 2014).

The original Cochrane Review of LF-LAM for tuberculosis included 12 studies (Shah 2016). In that review, six studies evaluated accuracy among symptomatic individuals and found a low pooled sensitivity of 45% (Credible Interval (Crl) 29% to 63%) and specificity of 92% (Crl 80% to 97%) against a microbiological reference standard (Shah 2016). In participants with CD4 ≤ 100 cells/µL, pooled sensitivity increased to 56% (Crl 41% to 70%) and specificity decreased to 90% (81% to 95%). In the original review, the accuracy of LF-LAM was also evaluated in participants not assessed of symptoms (i.e. termed ‘TB screening’ and now described as studies among ‘unselected participants’) with sensitivity ranging from 0% to 44% (Shah 2016).

In 2015, informed by the original Cochrane Review, the WHO made a conditional recommendation for using LF-LAM to assist with the diagnosis of tuberculosis in HIV-positive people with advanced disease (described below) and a strong recommendation against using the test “as a screening test for tuberculosis” based on the data among unselected participants (WHO Lipoarabinomannan Policy Guidance 2015).

Since 2015, additional evidence for the use and clinical impact of LF-LAM has emerged. This Cochrane Review update includes published studies evaluating the accuracy of the commercially available LF-LAM, Alere Determine TB LAM Ag assay for diagnosis of active tuberculosis (pulmonary and extrapulmonary tuberculosis) in people living with HIV and informed the updated WHO guidelines on the use of the test. Of note, in 2018, preliminary performance characteristics of a second commercially developed lateral flow assay to detect LAM for the diagnosis of tuberculosis was announced based on data from frozen biobank specimens (Fujifilm SILVAMP TB LAM, Japan; FujifLAM) (Broger 2019). The test is projected to become commercially available in 2020; no studies of FujilAM accuracy are included in this updated review.
Target condition being diagnosed

The target condition is active tuberculosis, which includes pulmonary and extrapulmonary tuberculosis.

Index test(s)

The urine-based lateral flow lipoarabinomannan immunocapture assay (LF-LAM) is a commercially available point-of-care test for active tuberculosis (Alera Determine™ TB LAM Ag, Abbott, Palatine, IL, USA, previous Alera Inc., Waltham, MA, USA). Lipoarabinomannan (LAM) is a lipopolysaccharide present in mycobacterial cell walls (Brennan 2003), which is released from metabolically active or degenerating bacterial cells during tuberculosis disease (Briken 2004). LAM is detectable in urine of people with active tuberculosis disease and evaluated for both LAM ELISA and the LF-LAM testing platforms (Lawn 2012; Minion 2011; Peter 2010; Shah 2016). The original Cochrane Review of LF-LAM (Shah 2016) and a meta-analysis of an earlier generation LAM ELISA test (Minion 2011) both demonstrated that the accuracy of urinary LAM detection was improved among people living with HIV with advanced immunosuppression. Several hypotheses may explain the higher sensitivity of urine LAM detection in people living with HIV including higher bacillary burden and antigen load (Shah 2010), greater likelihood of genitourinary tract tuberculosis involvement, and greater glomerular permeability to allow increased antigen levels in urine (Lawn 2016; Minion 2011).

LF-LAM testing is performed manually by applying 60 µl of unprocessed urine to the sample pad of the Determine™ TB LAM Ag test and leave the strip test to incubate at room temperature for 25 minutes (Alera 2017; Appendix 1). The strip is then inspected by eye. The intensity of any visible band is graded by comparing it with the intensities of the bands on a manufacturer-supplied reference scale card. Of note, the reference scale was revised in January 2014. Prior to January 2014, the reference scale card included five bands (grade 1 representing a very low intensity band to grade 5 representing a high/dark intensity band). After January 2014, the manufacturer revised the reference scale card to have four reference bands, such that the band intensity for the new grade 1 corresponded to the band intensity for the previous grade 2 (Appendix 2). Under the current manufacturer recommendations (using the revised four bands reference card), only bands that are grade 1 or higher are considered positive (Alera 2017; Appendix 2).

Clinical pathway

Based on current WHO guidelines (WHO Lipoarabinomannan Policy Guidance 2015), the proposed role for the LF-LAM test is as an ‘add on’ to clinical judgement and with other tests to assist in tuberculosis diagnosis. The test does not have a role as a replacement or triage test. The reason for specifying ‘clinical judgement’ is that, in our view, clinical judgement for HIV-associated tuberculosis carries more importance than in many other cases (i.e. tuberculosis treatment may be provided regardless of the test results). The reason for specifying ‘add on’ is that for settings where the test is not available, the same clinical judgement is used; hence for settings where the test is available, it is used ‘in addition’. Importantly, some HIV-positive people may have difficulties in producing a good quality sputum specimen or any sputum at all.

In the WHO policy guidance on the use of LAM, it is recommended that LF-LAM “may be used to assist in the diagnosis of tuberculosis in HIV-positive adult inpatients with signs and symptoms of tuberculosis (pulmonary/and/or extrapulmonary) and a CD4 cell count less than or equal to 100 cells/µL, or in people living with HIV who are ‘seriously ill’ regardless of CD4 count or if the CD4 count is unknown” (WHO Lipoarabinomannan Policy Guidance 2015). The recommendations also apply to HIV-positive outpatients and children with signs and symptoms of tuberculosis (pulmonary and/or extrapulmonary) based on the generalization of data from adult inpatients while acknowledging the limitation of available data (WHO Lipoarabinomannan Policy Guidance 2015). The WHO recommends that LF-LAM should not be used for general tuberculosis screening “owing to suboptimal sensitivity” (WHO Lipoarabinomannan Policy Guidance 2015).

In 2016, WHO developed an algorithm for managing people living with HIV presumed to have tuberculosis that was published as part of the guidelines on use of ART for treating and preventing HIV infection (WHO ART Guidelines 2016). The algorithm recommends the Xpert® MTB/RIF assay (Cepheid, Sunnyvale, USA) as the initial diagnostic test in adults and children with presumed HIV-associated tuberculosis (WHO Xpert® Policy Update 2013). It is recommended that Xpert® MTB/RIF should be used rather than conventional microscopy and culture for testing sputum and specific extra-pulmonary specimens like cerebrospinal fluid and lymph nodes (WHO Xpert® Policy Update 2013). The algorithm includes recommendations on use of LF-LAM as a test that may assist in diagnosing active tuberculosis among seriously ill adults and children living with HIV, regardless of CD4 count (WHO ART Guidelines 2016). Use of LF-LAM is also included as one of the components in the package for diagnosis of tuberculosis in the WHO guidelines for managing people presenting with advanced HIV (WHO Managing Advanced HIV Disease 2017).

When extrapulmonary tuberculosis is suspected, it is recommended to obtain “appropriate specimens from the suspected sites of involvement for microscopy, culture, and histopathological examination” (TB CARE 1 2014). A newly published Cochrane Review found that Xpert MTB/RIF may be helpful in confirming the diagnosis of extrapulmonary tuberculosis (Kohl 2018). However, evaluation for extrapulmonary tuberculosis often requires invasive diagnostic procedures that may have low yield even in people with advanced disease.

To identify those who need referral for tuberculosis diagnostic testing, the WHO guidelines on intensified case-finding in people living with HIV recommend that, in resource-constrained settings, people living with HIV who report “any one of the symptoms of current cough, fever, weight loss, or night sweats should be evaluated for tuberculosis and other diseases” referred to as the WHO symptom screening rule (Getahun 2011; WHO ICF 2011). People living with HIV with tuberculosis may however not exhibit common symptoms of tuberculosis disease. In some settings with high prevalence of tuberculosis, clinicians have therefore considered evaluating all patients presenting for care for tuberculosis without assessment or consideration of specific signs or symptoms of tuberculosis.

In the context of LF-LAM being recommended in combination with existing tuberculosis tests to increase early tuberculosis diagnosis and treatment, the downstream consequences of LF-LAM testing include the following.
• True-positive (TP): people would benefit from rapid non-site specific diagnosis and early initiation of tuberculosis treatment.

• True-negative (TN): people would be spared unnecessary treatment and would benefit from reassurance and pursuit of an alternative diagnosis.

• False-positive (FP): people would likely experience anxiety and morbidity caused by additional testing, unnecessary treatment, and possible adverse effects; possible stigma associated with a tuberculosis diagnosis; and a FP result may halt further diagnostic evaluation. However, as FP results increase (specificity decrease) as CD4 count decrease, the observed FP results may be due to an inability of the sickest patients to produce a sputum specimen and are TP results being misclassified as FP. Given the high mortality in persons living with HIV, acting on all positive LF-LAM results may balance the possibly adverse effects associated with unnecessary diagnosis and treatment.

• False-negative (FN): people would experience increased risk of morbidity and mortality and delayed treatment initiation; there would be continuous risk of tuberculosis transmission. These concerns may, however, be limited when using LF-LAM in combination with existing tuberculosis tests.

Importantly, it should be noted that LF-LAM does not provide information about drug resistance and some individuals with unidentified drug-resistant tuberculosis may be inappropriately treated with a regimen appropriate for drug-susceptible tuberculosis.

Alternative test(s)

In this section, we briefly describe selected alternative tests for detection of tuberculosis. For a comprehensive review of these tests, we refer the reader to several excellent resources (Lewinsohn 2017; Unitaid 2017).

Smear microscopy (light microscopy (Ziehl-Neelsen), fluorescence microscopy, or light-emitting diode (LED) fluorescence microscopy) is the examination of smears for acid-fast bacilli (tuberculosis bacteria) under a microscope. Advantages of smear microscopy include its simplicity, low cost, speed, and high specificity in high tuberculosis burden areas. In addition, smear microscopy identifies the most infectious tuberculosis patients. Smear microscopy can be performed in basic laboratories. Drawbacks of smear microscopy include the need for specialized training and its relatively low sensitivity, 50% to 60% on average for a direct smear. Among people living with HIV sensitivity is reported as low as 22% to 43% (Getahun 2007). Around 5000 to 10,000 organisms per mL must be present in the specimen for tuberculosis bacteria to be visible by microscopy (Lewinsohn 2017). Although, the sensitivity of microscopy can be improved by approximately 10% with fluorescence (Steingart 2006), a large number of tuberculosis cases still go undiagnosed if tuberculosis diagnosis solely relies on sputum smear microscopy. Smear-negative tuberculosis is disproportionately more common in HIV-positive than HIV-negative individuals, accounting for 24% to 61% of all pulmonary cases in people living with HIV, the yield decreasing with lower CD4 cell counts (Getahun 2007; Perkins 2007).

Mycobacterial culture is a method used to grow bacteria on nutrient-rich media. In comparison with microscopy, a positive culture requires only around 100 organisms per mL and therefore can detect lower numbers of tuberculosis bacteria (Lewinsohn 2017). Additionally, culture is essential for species identification and drug susceptibility testing. However, culture may take up to six to eight weeks and requires a highly equipped laboratory.

Nucleic acid amplification tests (NAATs) are molecular systems that can detect small quantities of genetic material (DNA or RNA) from microorganisms, such as M. tuberculosis. The key advantage of NAATs is that they are rapid diagnostic tests, potentially providing results in a few hours. A variety of molecular amplification methods are available, of which polymerase chain reaction (PCR) is the most common. NAATs are available as commercial kits and in-house tests (based on a protocol developed in a laboratory) and are used routinely in high-income countries for tuberculosis detection. In-house PCR is widely used in low-income countries because these tests are less expensive than commercial kits. However, in-house PCR is known to produce highly inconsistent results (Flores 2005).

Xpert MTB/RIF and Xpert Ultra, the newest version (Cepheid, Sunnyvale, USA), are fully automated NAATs, that simultaneously and rapidly detect M. tuberculosis complex and rifampicin resistance (WHO Xpert® Policy Update 2013; WHO Xpert Ultra 2017). A Cochrane Review found that Xpert MTB/RIF was sensitive and specific for both pulmonary tuberculosis detection and rifampicin resistance detection (Horne 2019). Compared with Xpert MTB/ RIF, Xpert Ultra had higher sensitivity and lower specificity for tuberculosis and similar sensitivity and specificity for rifampicin resistance (one study) (Horne 2019). Although sputum testing with Xpert MTB/RIF has high sensitivity for smear-positive pulmonary tuberculosis (98%), sensitivity is lower for smear-negative pulmonary tuberculosis (67%) (Horne 2019). In another Cochrane Review including 66 studies evaluating Xpert MTB/RIF for detection of extrapulmonary tuberculosis, pooled sensitivity was found to vary across different types of specimens from 31% in pleural tissue to 97% in bone or joint fluid, whereas specificity varied less from 82% in bone or joint tissue to 99% in pleural fluid and urine (Kohli 2018). These findings have bearing for HIV-associated tuberculosis where smear-negative tuberculosis and extrapulmonary tuberculosis are disproportionately higher. In 2017, based on a non-inferiority analysis of Xpert Ultra compared with Xpert MTB/RIF, the WHO stated that recommendations on the use of Xpert MTB/RIF also apply to the use of Xpert Ultra as the initial diagnostic test for all adults and children with signs and symptoms of tuberculosis (WHO Xpert Ultra 2017).

The loop-mediated isothermal amplification test, TB-LAMP (Eiken Chemical Co., Tokyo, Japan), has been recommended by the WHO since 2016, for diagnosing pulmonary tuberculosis in adults (WHO TB-LAMP 2016). It is a manual and simple assay that can be performed directly on sputum samples with a result provided in less than one hour (Yuan 2014). It is suitable for use at peripheral health centres and is promoted as a test more sensitive than microscopy, but inferior to Xpert MTB/RIF (WHO TB-LAMP 2016).

FujiLAM is a novel, lateral flow urine-based, point-of-care test for tuberculosi diagnosis in people living with HIV in low-resource settings. This assay was developed to detect LAM with results available in less than one hour and was announced on 26 September 2018, the day of the first United Nations General Assembly high-level meeting on tuberculosis (FIND 2018). Fuji LAM has been evaluated on frozen biobank samples originating from various studies of hospitalized HIV-positive patients with an estimated sensitivity of 70% (Broger 2019). Studies are ongoing
to evaluate FujiLAM performance on bio-samples stored from outpatient HIV-cohorts and prospective studies to evaluate the diagnostic accuracy of FujiLAM are now called for as the next step forward. These trials are being supported by the Global Health Innovative Technology fund and the German Federal Ministry of Education and Research. The Foundation for Innovative New Diagnostics (FIND) participated in development of the test, that was supported by the governments of the Netherlands and Australia, UK aid from the UK government, and the Bill & Melinda Gates Foundation.

Rationale
To address tuberculosis as the leading cause of morbidity and mortality among people living with HIV, new tests and strategies for detection of tuberculosis are urgently needed. Among the key priorities identified by the WHO, healthcare providers, patients, and advocacy groups is development of point-of-care, non-site specific tests for tuberculosis (Batz 2011; Pai 2012; Weyer 2011; WHO TTP 2014). To date, LF-LAM is the only commercially available point-of-care test for tuberculosis. LF-LAM, if sufficiently accurate, would satisfy many of the established minimum specifications for a point-of-care test for tuberculosis (Appendix 3; Batz 2011). LF-LAM testing could provide obvious benefits for HIV-positive people by earlier detection of pulmonary tuberculosis that may be missed by sputum smear microscopy and sputum Xpert MTB/RIF and extrapulmonary tuberculosis that may be missed by sputum-based testing. Studies that evaluate the impact of the use of LF-LAM on mortality and other patient outcomes are becoming available.

WHO guidelines on the use of urine LF-LAM were published in 2015 (WHO Lipoarabinomannan Policy Guidance 2015). Since 2015, additional evidence on the diagnostic accuracy of LF-LAM has emerged and is summarized in this Cochrane Review update. A draft of this review was used to informed the 2019 updated WHO guidelines on the use of LF-LAM.

OBJECTIVES

Primary objectives
We had two primary objectives:

1. To assess the accuracy of the lateral flow urine lipoarabinomannan assay (LF-LAM) for the diagnosis of active tuberculosis among HIV-positive adults with signs and symptoms of tuberculosis.
2. To assess the accuracy of LF-LAM for the diagnosis of active tuberculosis among HIV-positive adults irrespective of signs and symptoms of tuberculosis (i.e. unselected participants without consideration or assessment of tuberculosis signs and symptoms).

To estimate accuracy in HIV-positive individuals with signs and symptoms of tuberculosis (Objective 1), we combined studies in which presentation with signs and symptoms suggestive of tuberculosis was an inclusion criterion and refer to these as ‘Studies with symptomatic participants’. To estimate accuracy in HIV-positive adults irrespective of signs and symptoms of tuberculosis (Objective 2), we combined studies that considered all HIV-positive individuals eligible to participate, including both individuals with and individuals without symptoms of tuberculosis and refer to these as ‘Studies with unselected participants’. These studies enrolled participants without assessment or consideration of specific signs or symptoms of tuberculosis.

Secondary objectives
Our secondary objective was to investigate potential sources of heterogeneity in test accuracy, including clinical setting, CD4 cell count, and prevalence of tuberculosis in the studies.

METHODS

Criteria for considering studies for this review

Types of studies
We included primary studies that evaluated the diagnostic accuracy of urine LF-LAM assay for the detection of active tuberculosis in people living with HIV and compared the index test results with a defined microbiological reference standard. We included studies from which we could extract true positives (TP), false positives (FP), true negatives (TN), and false negatives (FN) values.

Diagnostic studies for tuberculosis are largely cross-sectional in design, but may include some clinical follow-up as part of patient classification. We included cross-sectional studies and observational cohort studies. In addition, we included randomized controlled trials that evaluated the use of the index test on patient health outcomes, but that also reported sensitivity and specificity. Although the study design was a randomized trial for the purpose of determining the impact of the test on participant outcomes, the study design was a cross-sectional study for the purpose of determining the diagnostic accuracy of the index test in this review. We excluded case-control studies and other study designs. We excluded data reported only in abstracts, reviews, comments, and editorial notes. We did not include unpublished studies.

Participants
People living with HIV are at increased risk of tuberculosis and may present with symptoms of tuberculosis but may also be asymptomatic or have symptoms not routinely associated with tuberculosis. We included participants who were adults (15 years and older is considered ‘adult’ for purpose of tuberculosis surveillance) and HIV positive. We included studies in which there was a suspicion of tuberculosis among study participants based on the presence of signs and symptoms compatible with tuberculosis (studies with symptomatic participants), as well as studies that included participants who presented for medical care irrespective of signs and symptoms of tuberculosis (studies with unselected participants). Signs and symptoms of tuberculosis include cough, fever, weight loss, and night sweats. Participants who were known to have active tuberculosis and were taking anti-tuberculosis drugs were not included.

Index tests
We included studies that evaluated the lateral flow lipoarabinomannan (LF-LAM) assay Alere Determine™ TB LAM Ag test (Abbott, Palatine, IL, USA, previous Alere Inc., Waltham, MA, USA) on urine samples. As of May 2019, the Alere Determine™ TB LAM Ag test was the only commercially available LF-LAM assay that had been evaluated in published studies.
We included studies that evaluated the test at the manufacturer’s recommended threshold for positivity, i.e. grade 1 and above on the updated reference scale card with four band intensities graded on a scale of 1 to 4. For studies that used the prior reference scale card with band intensities graded on a scale of 1 to 5, we included those that evaluated the test at grade 2 and above corresponding to the current recommended positivity threshold. We excluded studies that did not use a positivity threshold corresponding to the manufacturer’s recommendations. Results summarizing diagnostic accuracy at older thresholds (grade 1 on a scale of 1 to 5) can be found in the original review (Shah 2016).

Target conditions
The target condition was active tuberculosis, which includes pulmonary and extrapulmonary tuberculosis.

Reference standards
We required studies to diagnose tuberculosis using the following microbiological reference standard.

- ‘Tuberculosis’ is defined as a positive M tuberculosis culture or NAAT.
- ‘Not tuberculosis’ is defined as a negative M tuberculosis culture and NAAT (if performed).

NAAT tests included: Enhanced Amplified Mycobacterium Tuberculosis Direct Test (E-MTD, Gen-Probe, San Diego, USA); Amplicor Mycobacterium tuberculosis Test (Amplicor, Roche Diagnostics, Basel, Switzerland); COBAS® TaqMan® MTB Test (Roche Diagnostics); GenoType MTBDRplus (HAIN Lifesciences, Nehren, Germany); Xpert® MTB/RIF assay (Cepheid, Sunnyvale, USA); and Xpert® MTB/RIF Ultra.

We considered a higher quality reference standard to be one in which two or more specimen types were evaluated for tuberculosis diagnosis in all participants as part of a standardized study algorithm. We considered a lower quality reference standard to be one in which only one specimen type was evaluated for tuberculosis diagnosis, or if there was no algorithm defined to ensure a standardized approach for specimen collection and testing.

A microbiological reference standard, primarily culture, is considered the best reference standard. We expected all studies to obtain sputum specimens and some studies to obtain additional specimens for culture. However, the primary concern with relying on sputum culture alone is that tuberculosis diagnosis may be missed for the following reasons: people living with HIV may not be able to provide sputum specimens of sufficient quality; sputum bacillary load is typically low in people living with HIV; and a substantial proportion of people with HIV-associated tuberculosis cannot produce sputum at all (Lawn 2013), or have extrapulmonary tuberculosis without pulmonary tuberculosis. This means that index test TPs may be misclassified as FPs by sputum culture. Therefore, when evaluating LF-LAM with respect to sputum culture, the number of FPs (classified as positive by the index test and negative by the reference test) may be increased and LF-LAM specificity may be underestimated (Lawn 2015). This misclassification may also lead to underestimation of sensitivity. Increasing the sensitivity of the reference standard by evaluating multiple specimens, including evaluating specimens from sites of disease for extrapulmonary tuberculosis, may reduce the number of cases of tuberculosis incorrectly classified as ‘not tuberculosis’ by culture or NAAT if performed.

In the original Cochrane Review, we additionally considered a ‘composite microbiological and clinical reference standard’ recognizing that microbiological reference standards alone may fail to detect tuberculosis in patients with tuberculosis. However, our original review found relatively little data using a composite reference standard; found heterogeneity in defining and applying composite reference standards; and found a relatively modest impact on pooled estimates of sensitivity and specificity comparing microbiological and composite reference standards. Results assessing diagnostic accuracy against a composite reference standard can be found in the original review (Shah 2016).

Given the limitations of the reference standard, we might have considered pursuing latent class analysis (Chu 2009; Kohli 2018). However, we lacked patient-level data on the type of specimen and LAM is not site-specific, meaning a positive LAM alone does not tell us whether the patient has pulmonary or extrapulmonary tuberculosis.

Search methods for identification of studies
We attempted to identify all relevant studies regardless of language or publication status (published, unpublished, in press, and ongoing). As mentioned, we only included published studies in this review.

Electronic searches
We performed literature searches up to 11 May 2018 in the following databases using the search terms reported in Appendix 4: the Cochrane Infectious Diseases Group Specialized Register; MEDLINE (PubMed, from 1966); Embase (OVID, from 1947); Science Citation Index Expanded (SCI-EXPANDED, from 1900), Conference Proceedings Citation Index-Science (CPCI-S, from 1900), and BIOSIS Previews (from 1926), all three using the Web of Science platform; LILACS (BIREME, from 1982); and SCOPUS (from 1995). We also searched Clinicaltrials.gov and the search portal of the WHO International Clinical Trials Registry Platform (WHO ICTRP, www.who.int/trialsearch) to identify ongoing trials, and ProQuest Dissertations & Theses A&A (from 1861) to identify relevant dissertations. We included search results from the original review and re-evaluated previously included studies to determine if the studies met the refined inclusion criteria.

Searching other resources
We further examined reference lists of relevant reviews and studies and searched the WHO websites.

Data collection and analysis
Selection of studies
We used Covidence systematic review software to manage the selection of studies (Covidence 2017). Two review authors (MS and SB) independently examined all titles and abstracts identified from the electronic search to determine potentially eligible studies. We obtained the full-text articles of these potentially eligible studies and the same two review authors independently assessed inclusion based on predefined inclusion and exclusion criteria. We resolved disagreements through discussion and, if necessary, consulted a third review author (KRS). We included studies from the original
review if still eligible according to the predefined eligibility criteria. We maintained a list of excluded studies and the reasons for exclusion, and recorded these details in the Characteristics of excluded studies table and we prepared a PRISMA diagram.

Data extraction and management

We developed a standardized data extraction form and piloted the form on two of the included studies. Based on the pilot, we finalized the form (Appendix 5). Then two review authors (MS and SB) independently extracted data from each included study on the following characteristics.

- Author, publication year, study design, country/countries, clinical setting (outpatient or inpatient).
- Participants: age, gender, HIV-status, CD4 count, tuberculosis history, clinical status (asymptomatic, symptomatic).
- Quality Assessment of Diagnostic Accuracy Studies-2 (QUADAS-2) items.
- Cut-off used for determining a positive index test result and the reference card used.
- Samples collected (spum and/or extrapulmonary samples).
- Reference standard(s).
- The number of tuberculosis cases in the study.
- Number of TP, FN, FP, and TN values.
- Missing or unavailable test results.

We assigned country income status (high income, upper- and lower-middle income, and low income) as classified by the World Bank (World Bank 2018). In addition, we classified a country as being high burden or not high burden for tuberculosis/HIV according to the post-2015 era classification by the WHO (WHO Global Tuberculosis Report 2018). For studies that included both participants with HIV and without HIV infection, we extracted data only for participants with HIV. We contacted study authors for clarifications on the LF-LAM positivity threshold used if data were missing.

We used REDCap electronic data capture tools (Harris 2009) hosted at OPEN, Odense Patient data Explorative Network, Odense University Hospital, Odense, Denmark (SDU Open) to collect and manage study data. REDCap (Research Electronic Data Capture) is a secure, web-based application designed to support data capture for research studies, providing: 1) an intuitive interface for validated data entry; 2) audit trails for tracking data manipulation and export procedures; 3) automated export procedures for seamless downloads to common statistical packages; and 4) procedures for importing data from external sources. With regard to the use of REDCap, the content in this review is solely the responsibility of the authors.

Assessment of methodological quality

We used the QUADAS-2 tool tailored to this review to assess the quality of the included studies (Whiting 2011; Appendix 6). QUADAS-2 consists of four domains: patient selection, index test, reference standard, and flow and timing (flow and timing domain includes differential verification of tuberculosis status for study participants). We assessed all domains for risk of bias and the first three domains for concerns regarding applicability. As recommended, we first developed the guidance on how to appraise the questions in each domain. Then, one review author (SB) piloted the tool with two of the included studies and finalized the QUADAS-2 tool. Two review authors (MS and SB) independently completed the QUADAS-2 assessment. We resolved disagreements through discussion or consulted a third review author (KRS). We present the results of this quality assessment in the text, tables, and graphs.

Statistical analysis and data synthesis

We performed descriptive analyses of the characteristics of the included studies using Stata 15 (StataCorp 2017), and presented key study characteristics in the 'Characteristics of included studies' table. We used the number of TP5, FP5, FN5, and TN5 to calculate the individual study estimates of sensitivity and specificity and their 95% confidence intervals (CIs). We presented individual study results graphically by plotting the estimates of sensitivity and specificity (and their 95% CIs) in forest plots using Review Manager 5 (RevMan 5) (Review Manager 2014).

We presented results at the current manufacturer reference scale card for test interpretation, with band intensities graded 1 to 4, and considered all test results at grade 1 and above as positive. The prior reference scale card with five band intensities was used in the original Cochrane Review with grade 2 considered as positivity threshold that corresponds to the current grade 1 band intensity (Appendix 2). The original review also included several analyses at grade 1 which is no longer recommended for determining test positivity. To allow consistent comparisons, we converted results from older studies that used the ‘grade 2’ threshold and treated these as ‘grade 1’ in the updated review. As such, analyses labelled at ‘grade 2’ in the original Cochrane Review are in this review considered according to the new manufacturer reference card as ‘grade 1’. Studies in the original review that used the ‘grade 1’ threshold on the prior reference card were not included as this threshold is no longer recommended for determining test positivity.

We grouped the studies evaluating LF-LAM for: (I) diagnosis of tuberculosis in HIV-positive people with signs and symptoms of tuberculosis i.e. ‘Studies with symptomatic participants’ and (II) diagnosis of tuberculosis in HIV-positive people, irrespective of signs and symptoms of tuberculosis i.e. ‘Studies with unsellected participants’.

When data were sufficient, we carried out meta-analyses to estimate LF-LAM pooled sensitivity and specificity with a bivariate random-effects model (Chu 2006; Reitsma 2005). This approach allowed us to calculate pooled sensitivity and specificity while dealing with potential sources of variation caused by: (1) imprecision of sensitivity and specificity estimates within individual studies; (2) correlation between sensitivity and specificity across studies; and (3) variation in sensitivity and specificity between studies.

We estimated all models using a Bayesian approach implemented using OpenBUGS (Lunn 2009). Under the Bayesian approach, all unknown parameters must be provided a prior distribution that defines the range of possible values of the parameter and the weight of each of those values, based on information external to the data. Because most meta-analyses involved few studies (eight or less), which could lead the model to be just identified, we chose to use low-information prior distributions for most parameters and a more informative prior on the between-study standard deviations.
which are particularly sensible in meta-analyses with few studies (Spiegelhalter 2004).

We defined prior distributions on the log-odds scale over the pooled sensitivity and specificity parameters, their corresponding between-study standard deviations (SDs) and the correlation between the sensitivities and specificities across studies. For the pooled log odds of the sensitivity or log odds of the specificity, we used a normal prior distribution with mean 0 and a variance of 4 (or a precision of 0.25). This corresponds to a roughly uniform distribution over the pooled sensitivity and pooled specificity on the probability scale. The 2.5% and 97.5% prior distribution quantiles for the pooled sensitivity or pooled specificity are 2.0% and 98.0%, slightly wider than for a standard normal distribution. For the between-study precision we used a gamma distribution with a shape parameter of two and rate parameter of 0.5. This corresponds to a 95% prior credible interval (CrI) for the between-study SD in the log odds of sensitivity or log odds of specificity ranging from roughly 0.29 to 1.44, corresponding to moderate to high values of between-study heterogeneity. The resulting median 2.5% and 97.5% prior distribution quantiles for the predicted sensitivity or predicted specificity are 0.1% and 99.9%. Covariance terms followed a uniform prior distribution whose upper and lower limits were determined by the sensitivity of the two tests. We have summarized the models we used (including the prior distributions) and the OpenBUGS programs we used to estimate them in Appendix 7.

To study the sensitivity of our results to the choice of prior distributions given above, we considered alternative prior distributions that were less informative, which allowed a wider range of possible values. We increased the variance of the normal distributions over the pooled log odds of the sensitivity or specificity to 100. We used a uniform prior distribution ranging from zero to three over the between-study SD on the log odds scale. We found that the pooled estimates remained roughly the same with these alternative priors, though the posterior CrIs were wider, as expected. We combined information from the prior distribution with the likelihood of the observed data, in accordance with Bayes’ theorem in the OpenBUGS program, which resulted in a sample from the posterior distribution of each unknown parameter. Using this sample, we calculated various descriptive statistics of interest. We estimated the median pooled sensitivity and specificity and their 95% CrI. The median or the 50% quantile is the value below which 50% of the posterior sample lies. We reported the median because the posterior distributions of some parameters may be skewed, and the median would be considered a better point estimate of the unknown parameter than the mean in such cases. The 95% CrI is the Bayesian equivalent of the classical (frequentist) 95% confidence interval (CI) (we indicated 95% CI for individual study estimates and 95% CrI for pooled study estimates as appropriate). The 95% CrI may be interpreted as an interval that has a 95% probability of capturing the true value of the unknown parameter given the observed data and the prior information.

We also estimated the predicted sensitivity and specificity in a future study together with their 95% CrIs. The predicted estimate is our best guess for the estimate in a future study and is the same as the pooled estimate. The CrIs, however, may be different. These values were derived from the predicted region typically reported in a bivariate meta-analysis plot. If there is no heterogeneity between the included studies, the CrI around the predicted estimate will be the same as the CrI around the pooled estimate. On the other hand, if there is considerable heterogeneity between studies, the CrI around the predicted estimate will be much wider than the CrI around the pooled estimate. We generated summary plots displaying the individual study estimates for sensitivity and specificity, the pooled estimate for sensitivity and specificity with the 95% credible region and the 95% prediction region using R (R Statistical Computing 2016).

In our original review we evaluated the incremental change in sensitivity and specificity when combining LF-LAM with smear microscopy or Xpert MTB/RIF (Shah 2016). We did not undertake analysis of incremental benefit in the current review as it was beyond the scope of this review, and data within published manuscripts were limited.

**Approach to uninterpretable LF-LAM results**

We excluded uninterpretable test results from the meta-analyses of sensitivity and specificity, but we reported the number and proportion of uninterpretable test results from each study when such data were available.

**Investigations of heterogeneity**

Initially, we investigated heterogeneity through visual examination of forest plots of sensitivities and specificities and through visual examination of the ROC plot of the raw data. When data were sufficient, we performed subgroup analyses with the following categorial covariates: clinical setting (inpatient versus outpatient); CD4 count (CD4 ≤ 200; CD4 ≤ 100; CD4 101-200; CD4 > 200 and; CD4 > 100 cells per μL). To further investigate heterogeneity, we performed a subgroup analysis by the prevalence of tuberculosis in the studies and classified prevalence as greater than the median value versus less than or equal to the median value. We investigated heterogeneity separately for studies with symptomatic participants and studies with unselected participants. We generated the plots depicting the pooled results within CD4 count categories using R (R Statistical Computing 2016).

**Sensitivity analyses**

We performed sensitivity analyses by limiting inclusion in the meta-analysis to the following:

- Studies that avoided inappropriate exclusions, for example, studies that included participants who could not produce sputum. For this analysis we included studies that we scored as ‘yes’ for the QUADAS-2 question, “Did the study avoid inappropriate exclusions?” (low risk of bias for participant selection).

- Studies with a higher quality reference standard, for example studies that included two or more specimen types. For this analysis, we included studies that we scored as ‘yes’ for the QUADAS-2 question, “Is the reference standard likely to correctly classify the target condition?” (low risk of bias for the reference standard).

- Studies that used only fresh urine specimens for LAM testing.

- Studies initially categorized as ‘studies among unselected participants’ that included more than 80% of symptomatic participants were re-categorized as ‘studies with symptomatic participants’. We conducted this analysis to explore the possibility that these studies represented a comparable population to the studies of symptomatic participants even...
though participants were not explicitly enrolled in the study on
the basis of specific tuberculosis symptoms.

Assessment of reporting bias
We did not carry out a formal assessment of publication bias
using methods such as funnel plots or regression tests because
such techniques have not been helpful for diagnostic test accuracy
studies (Macaskill 2010).

Assessment of the certainty of the evidence
We assessed the certainty of evidence as recommended using the
GRADE approach (Balshem 2011; Schünemann 2008; Schünemann
2016). As recommended, we rated the certainty of evidence as
either high (not downgraded), moderate (downgraded by one
level), low (downgraded by two levels), or very low (downgraded
by more than two levels) based on five domains: risk of bias,
indirectness, inconsistency, imprecision, and publication bias. For
each outcome, the certainty of evidence started as high when there
were high-quality observational studies (cross-sectional or cohort
studies) that enrolled participants with diagnostic uncertainty. If
we found a reason for downgrading, we used our judgement to
classify the reason as either serious (downgraded by one level) or
very serious (downgraded by two levels).

Four review authors (SB, MS, ND, and KRS) discussed judgments
and applied GRADE in the following way.

- Risk of bias: we used QUADAS-2 to assess risk of bias.
- Indirectness: we used QUADAS-2 for concerns of applicability
  and looked for important differences between the populations
  studied (for example, in the spectrum of disease), the setting,
  and index test and asked are differences sufficient to lower
  certainty in results?
- Inconsistency: GRADE recommends downgrading for
  unexplained inconsistency in sensitivity and specificity
  estimates. We carried out pre-specified analyses to investigate
  potential sources of heterogeneity and did not downgrade when
  we felt we could explain inconsistency in the accuracy estimates.
- Imprecision: we considered a precise estimate to be one that
  would allow a clinically meaningful decision. We considered
  the width of the CrI, and asked ourselves, “Would we make
  a different decision if the lower or upper boundary of the CrI
  represented the truth?” In addition, we worked out projected
  ranges for TP, FN, TN, and FP for a given prevalence of
  tuberculosis and made judgements on imprecision from these
calculations.
- Publication bias: we rated publication bias as undetected (not
  serious) for several reasons including the comprehensiveness
  of the literature search and extensive outreach to tuberculosis
  researchers to identify studies.

RESULTS

Results of the search
We identified 15 unique studies that met the inclusion criteria
of this review. We included data from six published manuscripts
from the original WHO Guidelines (WHO Lipoarabinomannan Policy
Guidance 2015), and Cochrane Review (Shah 2016) that met the
refined inclusion criteria (Bjerrum 2015; Drain 2015a; LaCourse
2016; Nakiyangi 2014; Peter 2012a; Peter 2015), and nine new
studies identified in the updated search (Drain 2016; Floridia 2017;
Hanifa 2016; Huerga 2017; Juma 2017; Lawn 2017; Pandie 2016;
Peter 2016; Thit 2017). Of six previously included studies, three were
excluded because they did not use the currently recommended
threshold for test positivity (Balcha 2014; Drain 2014a; Lawn 2012a);
one previously included abstract (Lawn 2014) was added as an
updated published manuscript (Lawn 2017); one abstract remained
unpublished (Andrews 2014); and one abstract was published but
did not provide diagnostic accuracy data (Drain 2014b). Eight
studies evaluated the accuracy of LF-LAM for tuberculosis diagnosis
in participants with signs and symptoms suggestive of tuberculosis.
Seven studies evaluated the accuracy of LF-LAM for diagnosis
of unselected participants without assessment of symptoms (i.e.
patients may or may not have had tuberculosis signs and symptoms
at enrolment).

Figure 1 shows the flow of studies in the review. We listed
the excluded studies and the reasons for their exclusion in the
‘Characteristics of excluded studies’ section. All studies were
written in English.
Methodological quality of included studies

Risk of bias and applicability concerns for each of the 15 included studies is shown in Figure 2; Figure 3.
Figure 2. Studies with symptomatic participants - Risk of bias and applicability concerns summary: review authors' judgements about each domain for each included study.
Studies with symptomatic participants

Eight studies were included that evaluated LF-LAM for tuberculosis diagnosis among symptomatic participants suspected of tuberculosis. Risk of bias and applicability concerns for the studies with symptomatic participants are shown in Figure 2.

In the patient selection domain, we considered six studies (75%) to be at high risk of bias because: (1) the study excluded all smear-positive participants (Drain 2016); (2) the studies excluded participants who could not expectorate or produce sputum despite sputum induction (Drain 2016; Nakiyingi 2014; Peter 2015); (3) the study excluded participants from analysis if, in the absence of a positive reference standard result, there was one sample with no Xpert MTB/RIF result or contaminated result (culture) (Huerga 2017); (4) the study only included patients suspected of extrapulmonary tuberculosis and excluded patients suspected of pulmonary tuberculosis (Juma 2017); (5) the study only included participants with pericardial effusion and suspected tuberculosis and excluded participants suspected of other forms of tuberculosis (Pandie 2016). All studies were cross-sectional, cohort or randomized controlled studies. Regarding applicability, seven studies (88%) had low concern in the patient selection domain because the studies included the appropriate participants and settings. We judged one study (12%) to have high concern for applicability as the participants did not resemble people with presumed HIV/tuberculosis co-infection i.e. participants were smear-negative HIV-positive and HIV-negative patients with a Karnofsky Performance score < 50 (Drain 2016).

In the index test domain, we judged one study (12%) at high risk of bias as the study used grade 2 (on the updated reference scale card) as the test positivity threshold, as opposed to the

Figure 3. Studies with unselected participants - Risk of bias and applicability concerns summary: review authors' judgements about each domain for each included study.
current manufacturer recommendation to use grade 1 to define test positivity (Juma 2017). We also considered this study to have high concern of applicability because the test procedure was inconsistent with the manufacturers’ recommendations (Juma 2017). The remaining studies all used the recommended threshold for positivity and interpreted the test without knowledge of the results of the reference standard, and we considered them to have low concern for applicability.

In the reference standard domain, we considered seven studies (88%) to be at high risk of bias because: (1) the studies did not include testing of any extrapulmonary specimens (Drain 2016, Peter 2015); (2) the study did not include testing of any respiratory samples (Juma 2017); (3) the study only tested respiratory samples for some of the participants (Pandie 2016); (4) the study only tested extrapulmonary specimens in addition to respiratory samples for some of the participants (Huerga 2017); (5) health providers selected the sites for testing based on their own clinical suspicion (Peter 2012a; Peter 2016). We deemed three studies at high concern for applicability as they lacked a study or protocol directed testing (Pandie 2016; Peter 2012a; Peter 2016). In these studies, health providers selected the sites for testing based on their own clinical suspicion, and it was unclear if their choice of reference standard would correctly classify tuberculosis.

In the flow and timing domain, we considered four studies (50%) to be at high risk of bias because not all participants received the same reference standard (Huerga 2017; Peter 2012a; Peter 2016), or because not all participants were included in the two-by-two tables (Huerga 2017; Pandie 2016). We judged the remaining studies to be at low risk of bias because all participants received the index test, the same reference standard and no participants were excluded from the two-by-two table.

Studies with unselected participants

Seven studies contributed data for the purpose of evaluating LF-LAM for tuberculosis diagnosis among unselected participants who may or may not have tuberculosis signs or symptoms. Risk of bias and applicability concerns for the studies with symptomatic participants are shown in Figure 3.

In the patient selection domain, we considered four studies (57%) to be at high risk of bias because these studies excluded participants who could not expectorate or produce sputum samples (Bjerrum 2015; Drain 2015a; Floridia 2017; LaCourse 2016). All studies were cross-sectional or cohort studies. Regarding applicability, we judged that all studies (100%) included the appropriate participants and settings.

In the index test domain, we considered all studies at low risk of bias as all studies used LF-LAM, pre-specified the grade used for positivity, and interpreted the test at the recommended positivity threshold without knowledge of the results of the reference standard. We considered the test conduct and interpretation in all studies to be applicable.

In the reference standard domain, we considered five studies (71%) to be at high risk of bias because these studies did not include microbiological testing on extrapulmonary specimens (Bjerrum 2015; Drain 2015a; Floridia 2017; LaCourse 2016; Thit 2017). Thit 2017 also did not report if the reference standard results were interpreted without knowledge of the index test result. One study (Thit 2017) did not report if they speciated mycobacteria isolates and was judged to have unclear concern for applicability. We judged the remaining six studies to be of low concern in terms of applicability.

In the flow and timing domain, we considered two studies (29%) to be at high risk of bias because the studies collected specimens for index and reference standard tests up to six months apart (Hanifa 2016; Thit 2017). We also noted that five studies (71%) to be at low risk of bias because all participants received the index test and the same reference standard, and no participants enrolled were excluded from the two-by-two table.

Findings

The 15 included studies involved 6814 participants, 1761 (26%) with tuberculosis. Eight studies evaluated the accuracy of LF-LAM for tuberculosis diagnosis in participants with signs and symptoms suggestive of tuberculosis involving 3449 participants, 1277 (37%) with tuberculosis. Seven studies evaluated the accuracy of LF-LAM for diagnosis of unselected participants that may or may not have had tuberculosis signs and symptoms at enrollment involving 3365 participants, 432 (13%) with tuberculosis.

All studies were performed in countries with a high tuberculosis/ HIV burden (WHO Global Tuberculosis Report 2018), and classified as low-income or middle-income countries (World Bank 2018). We noted substantial differences in the studies for the following characteristics: type of population (‘studies with symptomatic participants’ and ‘studies with unselected participants’); setting (inpatients versus outpatients); median CD4 cell count; tuberculosis prevalence; inclusion and exclusion of participants based on whether or not they could produce sputum; and whether patients were evaluated for pulmonary tuberculosis, extrapulmonary tuberculosis, or both. The key study characteristics are summarized in Table 1 (Summary characteristics of included studies) and in Characteristics of included studies.

Table 2 presents pooled sensitivity and specificity results for LF-LAM grouped by the type of population, ‘studies among symptomatic participants’ and ‘studies among unselected participants’.

I. Diagnostic accuracy of LF-LAM for tuberculosis diagnosis in HIV-positive adults with signs and symptoms of tuberculosis

Of the 15 included studies, eight evaluated the accuracy of LF-LAM for tuberculosis diagnosis in participants with signs and symptoms suggestive of tuberculosis (Drain 2016; Huerga 2017; Juma 2017; Nakiyongi 2014; Pandie 2016; Peter 2012a; Peter 2015; Peter 2016). The suggestive signs and symptoms of tuberculosis varied from study to study, as described in Table 1 and in Characteristics of included studies, but were often based on any of cough, fever, weight loss, or night sweats as per the WHO symptom screening rule WHO ICF 2011. The tuberculosis prevalence in the study population ranged from 29% to 63%. Two studies were conducted exclusively among patients with presumed extrapulmonary tuberculosis (Juma 2017; Pandie 2016). Four studies were conducted exclusively in an inpatient setting (Juma 2017; Pandie 2016; Peter 2012a; Peter 2016); two studies exclusively in an outpatient setting (Drain 2016; Peter 2015); and two studies in both inpatient and outpatient settings (Huerga 2017; Nakiyongi 2014). The median CD4 cell count ranged from 81 to 210

Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV (Review)

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cells per µL across the eight studies, lower in studies evaluating inpatients (median CD4 between 81 to 139 cells per µL) compared to studies evaluating outpatients (median CD4 was 168 to 210 cells per µL). See Table 1 and Characteristics of included studies.

Primary analysis, LF-LAM for tuberculosis in HIV-positive adults with signs and symptoms of tuberculosis

For determining the overall accuracy of LF-LAM in HIV-positive adults with signs and symptoms of tuberculosis, eight studies provided data for 3449 participants, including 1277 (37%) with tuberculosis. Sensitivity estimates ranged from 23% to 68%, and specificity estimates from 75% to 100% (Figure 4).

Figure 4. Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies among symptomatic participants. The individual studies are ordered by decreasing sensitivity. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).

Juma 2017 evaluated diagnostic accuracy for extrapulmonary tuberculosis (all forms) exclusively and had the highest sensitivity of 68%. Pandie 2016 evaluated accuracy for pericardial tuberculosis and found sensitivity of 33%. Sensitivity was lowest in the studies by Peter 2015 and Drain 2016 that both reported a sensitivity of 23%. Differences between these studies and the other studies in this analysis were the setting (outpatient only), focus on pulmonary tuberculosis (no extrapulmonary samples were taken), and exclusion of participants unable to produce sputum. In particular, Drain 2016 included smear-negative participants with presumed tuberculosis and a small number of HIV-negative participants. Drain 2016 further excluded participants with a low Karnofsky score in order to target relatively well outpatients, where smear-negative tuberculosis is often seen. Specificity was lowest for Peter 2012a a study that included only inpatients and differed from other studies in that in both sputum and non-sputum-based sampling was performed at the discretion of the attending clinical team and not study directed. Pandie 2016 reported the highest specificity of 100%. This study, as mentioned, differed from others by evaluating accuracy only for pericardial tuberculosis. The authors further excluded a number of participants in the analysis for unknown reasons and reported two true negatives and zero false positives that may have inflated specificity. The pooled sensitivity and specificity (95% credible interval [CRI]) were 42% (21% to 55%) and 91% (85% to 95%) (Table 2). Appendix 8 presents the pooled and predicted sensitivity and specificity estimates together with the credible and prediction regions for LF-LAM for tuberculosis detection. Prediction intervals were very wide in all meta-analysis models indicating between-study heterogeneity (data not shown).

Uninterpretable index test results

Studies reported few uninterpretable test results. Peter 2012a reported that 1% to 2% of 423 tests remained ‘indeterminate’ (defined as a broken band in the patient window) after repeat testing. Peter 2016 reported that 0.3% (3/1172) test results remained ‘invalid’ after a repeat test (defined as no control band identified in the patient window or a broken/ incomplete band in the patient window). Peter 2015 reported that fewer than 1% of LF-LAM strip tests failed on the first attempt and required a second strip to produce valid results. Four studies reported that a valid LF-LAM result was obtained on the first attempt for all tests (Drain 2016; Huerga 2017; Juma 2017; Nakiyingi 2014), and one study did not provide information on uninterpretable tests (Pandie 2016).

Investigations of heterogeneity

LF-LAM accuracy among symptomatic participants, stratified by setting

LF-LAM accuracy in inpatient settings

Six studies were conducted among inpatients involving 2253 participants, 868 (39%) with tuberculosis (Huerga 2017; Juma 2017; Nakiyingi 2014; Pandie 2016; Peter 2012a; Peter 2016). Sensitivity estimates ranged from 33% to 69% and specificity estimates ranged from 75% to 100% (Figure 5). The highest sensitivity (69%) was reported by Huerga 2017 with a relatively low specificity (78%). This study enrolled inpatients with tuberculosis symptoms who had at least one of the following features: who were severely ill or who had CD4 < 200 cell per µL (median CD4 109) or who had a low body mass index (BMI). The study did not include microbiological or histological evaluation of extrapulmonary specimens for tuberculosis in their reference standard, which may have led to LAM-positive participants with extrapulmonary tuberculosis being misclassified as ‘false-positive’ and lowered specificity. Pandie 2016 reported the lowest sensitivity (33%) and a specificity of 100%. As mentioned, this study evaluated accuracy for pericardial tuberculosis and excluded participants from specificity analysis that may have affected the specificity.
estimate. The pooled sensitivity and specificity (95% CrI) among inpatients were 52% (40% to 64%) and 87% (78% to 93%) (Table 2).

**Figure 5.** Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies among symptomatic participants, by health care setting. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. The individual studies are ordered by decreasing sensitivity. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).

| Symptomatic adults, inpatients |
|-------------------------------|
| Study            | TP  | FP  | FN  | TN  | Sensitivity (95% CI) | Specificity (95% CI) |
| Huerga 2017       | 73  | 17  | 33  | 60  | 0.69 [0.50, 0.79]     | 0.78 [0.67, 0.87]    |
| Juma 2017         | 15  | 3   | 7   | 42  | 0.68 [0.46, 0.88]     | 0.93 [0.82, 0.96]    |
| Peter 2012a       | 58  | 31  | 68  | 94  | 0.60 [0.41, 0.53]     | 0.75 [0.67, 0.82]    |
| Nakinyingi 2014   | 114 | 19  | 132 | 287 | 0.46 [0.30, 0.53]     | 0.94 [0.80, 0.98]    |
| Peter 2016        | 166 | 04  | 186 | 726 | 0.46 [0.30, 0.51]     | 0.89 [0.86, 0.91]    |
| Pandie 2016       | 12  | 0   | 24  | 2   | 0.33 [0.19, 0.51]     | 1.00 [1.00, 1.00]    |

| Symptomatic adults, outpatients |
|-------------------------------|
| Study            | TP  | FP  | FN  | TN  | Sensitivity (95% CI) | Specificity (95% CI) |
| Huerga 2017       | 29  | 2   | 21  | 46  | 0.56 [0.43, 0.72]     | 0.95 [0.84, 0.98]    |
| Drain 2016        | 13  | 1   | 44  | 32  | 0.23 [0.05, 0.41]     | 0.97 [0.84, 1.00]    |
| Peter 2015        | 41  | 27  | 140 | 351 | 0.25 [0.17, 0.33]     | 0.93 [0.90, 0.95]    |
| Nakinyingi 2014   | 22  | 1   | 59  | 322 | 0.16 [0.12, 0.26]     | 0.99 [0.96, 1.00]    |

**LF-LAM accuracy in outpatient settings**

Four studies were conducted among outpatients involving 1196 participants, 409 (34%) with tuberculosis (Drain 2016; Huerga 2017; Nakinyingi 2014; Peter 2015). Sensitivity estimates ranged from 18% to 58% and specificity estimates ranged from 93% to 99% (Figure 5). The highest sensitivity (58%) was reported by Huerga 2017 where outpatients included were severely ill, or had a CD4 < 200 cell per μL, or had a low BMI below 17 kg/m². Among outpatients, pooled sensitivity and specificity were 29% (17% to 47%) and 96% (91% to 99%) (Table 2).

**LF-LAM accuracy among symptomatic participants, stratified by CD4 count**

We included five studies in the analyses by CD4 count (Nakinyingi 2014; Peter 2012a; Peter 2015; Peter 2016). See Figure 6 and Figure 7.
Figure 6. Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies with symptomatic participants, stratified by CD4 (CD4 > 200 and CD4 ≤ 200; CD4 > 100 and CD4 ≤ 100). The individual studies are ordered by decreasing sensitivity. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).
Figure 7. Plot by CD4 of diagnostic accuracy in adults with signs and symptoms of tuberculosis. (A) Sensitivity by CD4 strata; (B) Specificity by CD4 strata. The circle represents the pooled estimates (median), with bars representing 95% credible intervals.

LF-LAM accuracy stratified by CD4 > 200 cells per µL and ≤ 200 cells per µL

Three studies evaluated LF-LAM in participants with CD4 > 200 cells per µL (Nakiyingi 2014; Peter 2012a; Peter 2016). Sensitivity estimates ranged from 9% to 27% and specificity estimates ranged from 83% to 99% (Figure 6). In the four studies that evaluated participants with CD4 ≤ 200 cells per µL, sensitivity estimates ranged from 24% to 58% and specificity estimates ranged from 72% to 95% (Nakiyingi 2014; Peter 2012a; Peter 2015; Peter 2016, Figure 6). The pooled sensitivity (95% CI) was higher among participants with CD4 ≤ 200 cells per µL at 45% (31% to 61%) (4 studies, 1825 participants, 40% with tuberculosis) versus 16% (8% to 31%) among those with CD4 > 200 cells per µL (3 studies, 738 participants, 22% with tuberculosis) (Table 2); a significant difference in pooled sensitivity (95% CI) of -29% (-47% to -9%). The pooled specificity was 89% (77% to 94%) for participants with CD4 ≤ 200 cells per µL and 94% (81% to 97%) for those with CD4 > 200 cells per µL (Table 2), a difference in pooled specificity of 5% (-8% to 17%).

When we limited the analysis to studies involving inpatients with CD4 ≤ 200 cells per µL, the pooled sensitivity and specificity were 54% (34% to 73%) and 80% (58% to 91%) (2 studies, 1009 participants, 34% with tuberculosis) (Peter 2012a; Peter 2016). Only one study reported data for outpatients with CD4 ≤ 200 cells per µL (Peter 2015).

LF-LAM accuracy stratified by CD4 > 100 cells per µL and ≤ 100 cells per µL

Four studies evaluated LF-LAM in participants with CD4 > 100 cells per µL (Nakiyingi 2014; Pandie 2016; Peter 2015; Peter 2016). Sensitivity estimates ranged from 12% to 19% and specificity estimates ranged from 92% to 100% (Figure 6). In the five studies that evaluated participants with CD4 ≤ 100 cells per µL, sensitivity estimates ranged from 30% to 65% and specificity estimates ranged from 75% to 94% (Nakiyingi 2014; Pandie 2016; Peter 2012a; Peter 2015; Peter 2016). One study had no estimable specificity, as the authors reported zero true negatives (Pandie 2016). The pooled sensitivity (95% CI) was higher among participants with CD4 ≤ 100 cells per µL at 54% (38% to 69%) (4 studies, 1239 participants, 41% with tuberculosis) versus 17% (10% to 27%), (4 studies, 1519 participants, 28% with tuberculosis) among those with CD4 > 100 cells per µL, a significant difference in pooled sensitivity of -37% (-53% to -19%). The pooled specificity was 88% (77% to 94%) for participants with CD4 ≤ 100 cells per µL and 95% (89% to 98%) for those with CD4 > 100 cells per µL (Table 2), a difference in pooled specificity of 7% (-1% to 18%).

When we limited the analysis to studies involving inpatients with CD4 ≤ 100 cells per µL (Peter 2012a; Peter 2016), the pooled sensitivity and specificity were 61% (40% to 78%) and 81% (61% to 91%). Only one study reported data for outpatients with CD4 ≤ 100 cells per µL (Peter 2015).

LF-LAM accuracy stratified by CD4 strata

We observed that LF-LAM pooled sensitivity increased as the degree of immunodeficiency increased, from 16% (8% to 31%) in patients with CD4 cell count >200 cells per µL; to 24% (14% to 38%) in patients with CD4 count between 101 to 199; to 54% (38% to 69%) in patients with CD4 ≤ 100 (Figure 7). Also, we observed that most participants contributing data for the CD4 ≤ 200 cells per µL stratum (1825 participants including 722 (40%) with tuberculosis) were participants with CD4 ≤ 100 cells per µL (1239 participants including 512 (41%) with tuberculosis) (Table 2).

LF-LAM accuracy among symptomatic participants, stratified by tuberculosis prevalence

The median prevalence of tuberculosis in studies with symptomatic participants was 43% (interquartile range (IQR) 32% to 60%). In an analysis by tuberculosis prevalence, we found that pooled sensitivity and specificity for symptomatic participants in settings with tuberculosis prevalence of greater than 43% were 44% (27% to 59%) and 94% (60% to 97%), respectively.
Sensitivity analyses

In evaluation of the accuracy of LF-LAM for detection of tuberculosis, we undertook sensitivity analyses by limiting inclusion in the meta-analysis to the following:

- Studies with low risk of bias for patient selection (two studies)
- Studies with low risk of bias in the reference standard domain (one study)
- Studies that used fresh urine samples (four studies) rather than stored urine sample

Two studies with greater than 80% symptomatic participants (Bjerrum 2015; Lawn 2017) were re-assigned from ‘studies of unselected participants’ to ‘studies of symptomatic participants’.

These sensitivity analyses only included few studies and participants and made little difference to any of the findings (Table 3).

II. Diagnostic accuracy of LF-LAM for tuberculosis diagnosis in HIV-positive unselected adults irrespective of signs and symptoms for tuberculosis

Of the 15 studies included, seven studies evaluated the accuracy of LF-LAM for diagnosis in participants, irrespective of sign and symptoms (‘unselected participants’) (Bjerrum 2015; Drain 2015a; Floridia 2017; Hanifa 2016; LaCourse 2016; Lawn 2017; Thit 2017).

The tuberculosis prevalence in the study population varied from 1% in LaCourse 2016 to 33% in Lawn 2017. Six of the studies reported the proportion of symptomatic participants that were included (e.g. having a positive WHO symptoms screen) which varied from 19% (LaCourse 2016) to more than 90% of participants in two studies (Bjerrum 2015; Lawn 2017). No studies reported results specifically for individuals who were asymptomatic. Four studies were carried out in an outpatient setting (Drain 2015a; Floridia 2017; Hanifa 2016; LaCourse 2016), one study exclusively in an inpatient setting (Lawn 2017), and two studies in both inpatient and outpatient settings (Bjerrum 2015; Thit 2017). The median CD4 cell count across studies of unselected adults ranged from 111 to 437 cells per μL across studies. See Table 1 and Characteristics of included studies.

Primary analysis, LF-LAM for tuberculosis in HIV-positive adults irrespective of signs and symptoms of tuberculosis

For determining the overall accuracy of LF-LAM in HIV-positive adults irrespective of signs and symptoms of tuberculosis, seven studies provided data for 3365 participants, including 432 (13%) with tuberculosis. Sensitivity estimates ranged from 0% to 67%, and specificity estimates from 64% to 99% (Figure 8).

Figure 8. Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies among unselected participants not assessed for signs and symptoms of tuberculosis. The individual studies are ordered by decreasing sensitivity. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).

| Study     | TP   | FP   | FN   | TN   | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|-----------|------|------|------|------|----------------------|----------------------|----------------------|----------------------|
| Thit 2017 | 36   | 165  | 10   | 293  | 0.67 (0.53, 0.76)    | 0.54 (0.63, 0.68)    |                      |                      |
| Bjerrum 2015 | 24   | 21   | 31   | 393  | 0.44 (0.30, 0.56)    | 0.95 (0.92, 0.97)    |                      |                      |
| Lawn 2017  | 53   | 3    | 83   | 274  | 0.93 (0.81, 0.96)    | 0.93 (0.92, 0.97)    |                      |                      |
| Florida 2017 | 26   | 8    | 64   | 674  | 0.29 (0.20, 0.38)    | 0.98 (0.98, 1.00)    |                      |                      |
| Drain 2015a | 15   | 16   | 36   | 250  | 0.20 (0.16, 0.24)    | 0.94 (0.90, 0.97)    |                      |                      |
| Hanifa 2016 | 3    | 5    | 37   | 363  | 0.07 (0.02, 0.26)    | 0.98 (0.97, 1.00)    |                      |                      |
| LaCourse 2016 | 0    | 13   | 3    | 250  | 0.00 (0.00, 0.41)    | 0.95 (0.92, 0.97)    |                      |                      |

Sensitivity was lowest (0%) in LaCourse 2016, that differed from the other studies by including a) a population of exclusively pregnant women attending an antenatal care setting, b) a low proportion of symptomatic participants (19%), c) a low tuberculosis prevalence (1%), and d) a high median CD4 cell count (437 cells per μL). Specificity was lowest (64%) in the study by Thit 2017 that also reported the highest sensitivity (67%). This study reported that more than 90% of the FP results were grade 1 positive results (classified as positive according to current manufacturer recommendations). Participants included had a median CD4 at 270 cells per μL and 33% were symptomatic at enrolment. The study evaluated sputum samples only and allowed a follow-up for six months from LF-LAM testing at enrolment to final classification of participants as ‘Tuberculosis’ or ‘Not tuberculosis’ cases. Thit 2017 further differed from the other studies by being conducted in Myanmar, and is the only study included in this review that evaluated LF-LAM in a setting outside sub-Saharan Africa. The pooled sensitivity and specificity (95% CI) were 35% (22% to 50%) and 95% (89% to 98%) (Table 2). Appendix 9 presents the pooled and predicted sensitivity and specificity estimates together with the credible and prediction regions for LF-LAM for tuberculosis detection. Prediction intervals were very wide in all meta-analysis models indicating between-study heterogeneity (data not shown).

Uninterpretable index test results

The included studies reported no uninterpretable results.
Investigations of heterogeneity

LF-LAM accuracy among unselected participants, stratified by setting

LF-LAM accuracy in inpatient settings

We identified three studies that evaluated accuracy among inpatients involving 537 participants, 159 (30%) with tuberculosis.

Figure 9. Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies among unselected participants, by health care setting. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. The individual studies are ordered by decreasing sensitivity. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).

Unselected adults, inpatients

| Study     | TP  | FP  | FN  | IN  | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|-----------|-----|-----|-----|-----|----------------------|----------------------|----------------------|----------------------|
| Thit 2017 | 7   | 20  | 1   | 18  | 0.89 [0.47, 1.00]    | 0.29 [0.06, 0.55]    |                      |                      |
| Bjerrum 2015 | 10  | 6   | 5   | 47  | 0.67 [0.36, 0.88]    | 0.65 [0.72, 0.94]    |                      |                      |
| Lawn 2017 | 53  | 3   | 63  | 274 | 0.39 [0.31, 0.49]    | 0.99 [0.87, 1.03]    |                      |                      |

Unselected adults, outpatients

| Study     | TP  | FP  | FN  | IN  | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|-----------|-----|-----|-----|-----|----------------------|----------------------|----------------------|----------------------|
| Thit 2017 | 29  | 13  | 17  | 200 | 0.63 [0.48, 0.77]    | 0.67 [0.60, 0.75]    |                      |                      |
| Bjerrum 2015 | 14  | 13  | 26  | 346 | 0.35 [0.21, 0.52]    | 0.96 [0.94, 0.98]    |                      |                      |
| Florida 2017 | 26  | 8   | 64  | 674 | 0.25 [0.20, 0.38]    | 0.98 [0.97, 1.00]    |                      |                      |
| Drain 2015a | 15  | 16  | 38  | 250 | 0.26 [0.16, 0.42]    | 0.94 [0.90, 0.97]    |                      |                      |
| Hanifa 2016 | 3   | 5   | 37  | 363 | 0.07 [0.02, 0.20]    | 0.99 [0.97, 1.00]    |                      |                      |
| LaCourse 2016 | 0  | 13  | 3   | 250 | 0.00 [0.00, 0.12]    | 0.95 [0.92, 0.97]    |                      |                      |

Thit 2017 reported a very low specificity (39%) and a high sensitivity (88%), based on a relatively small sample size of 54 inpatients; eight (15%) with tuberculosis. The authors reported that 41 of the inpatients (76%) were symptomatic at enrolment with a median CD4 of 96 (IQR 37 to 277) cells per µL, which is comparable to evaluation of LF-LAM in a population with advanced HIV disease. The study further differed from the other studies by allowing a follow-up for six months for classification of participants and by being conducted in Myanmar as mentioned above. Most of the participants (91%) included in the studies by Bjerrum 2015 and Lawn 2017 were symptomatic and thus comparable to studies with symptomatic participants. See Table 1 and Characteristics of included studies. The pooled sensitivity and specificity (95% CrI) among unselected inpatients were 62% (41% to 83%) and 84% (48% to 96%) (Table 2).

LF-LAM accuracy in outpatient settings

Six studies were conducted among unselected outpatients, involving 2828 participants; 273 (10%) with tuberculosis (Bjerrum 2015; Drain 2015a; Florida 2017; Hanifa 2016; LaCourse 2016; Thit 2017). Sensitivity estimates ranged from 0% to 63% and specificity estimates ranged from 67% to 99% (Figure 9). Pooled sensitivity and specificity (95% CrI) were 31% (18% to 47%) and 95% (87% to 99%) (Table 2).

LF-LAM accuracy among unselected participants, stratified by CD4 count

There were limited data to evaluate LF-LAM by CD4 threshold for unselected participants. Five studies contributed data to the analyses by CD4 count (Bjerrum 2015; Drain 2015a; Hanifa 2016; LaCourse 2016; Lawn 2017). See Appendix 10; Appendix 11 and Table 2.

LF-LAM accuracy stratified by CD4 ≤ 200 cells per µL and > 200 cells per µL

Two studies evaluated LF-LAM in unselected participants with CD4 ≤ 200 cells per µL, all settings. Sensitivity and specificity were 45% and 93% for Bjerrum 2015, and 7% and 99% for Hanifa 2016, (Appendix 10). Pooled sensitivity and specificity were 26% (9% to 56%) and 96% (87% to 98%) (706 participants, 12% with tuberculosis) (Table 2).

For unselected outpatients with a CD4 ≤ 200 cells per µL, two studies contributed data (652 participants; 10% with tuberculosis). Sensitivity and specificity were 36% and 94% for Bjerrum 2015, and 7% and 99% for Hanifa 2016 (Appendix 10). Pooled sensitivity and specificity were 21% (8% to 48%) and 96% (89% to 99%) (Table 2).

When we limited the analysis to unselected inpatients with CD4 ≤ 200 cells per µL, only one study (Bjerrum 2015) contributed data and found a sensitivity of 64% (95% CI: 35% to 87%) and specificity of 82% (67% to 93%) (54 participants, 26% with tuberculosis).
For comparison, we assessed diagnostic accuracy among participants with CD4 > 200 cells per μL, all settings. Only one study reported data for participants with CD4 > 200 cells per μL and reported a sensitivity of 27% (95% CI: 6% to 61%) and specificity of 99% (95% CI: 96% to 100%) (Bjerrum 2015).

**LF-LAM accuracy stratified by CD4 ≤ 100 cells per μL and > 100 cells per μL**

Three studies evaluated patients with CD4 ≤ 100 cells per μL, all settings, and sensitivity and specificity ranged from 37% to 55% and specificity from 85% to 98% (Appendix 11). The pooled sensitivity and specificity (95% CI) among participants with CD4 ≤ 100 cells per μL were 47% (30% to 64%) and 90% (77% to 96%) (3 studies, 417 participants, 31% with tuberculosis) (Table 2).

When we limited the analysis to studies involving unselected inpatients with CD4 ≤ 100 cells per μL, two studies contributed data, (200 participants; 42% with tuberculosis). Sensitivity and specificity were 60% and 80% for Bjerrum 2015, and 55% and 98% for Lawn 2017 (Appendix 11). The pooled sensitivity and specificity were 57% (33% to 79%) and 90% (69% to 97%) (Table 2).

Two studies contributed data for unselected outpatients with CD4 ≤ 100 cells per μL (Bjerrum 2015; Hanifa 2016) (Appendix 11). The pooled sensitivity and specificity were 40% (20% to 64%) and 87% (68% to 94%) (217 participants, 21% with tuberculosis) (Table 2).

Four studies evaluated LF-LAM in participants with CD4 > 100 cells per μL (Bjerrum 2015; Drain 2015a; LaCourse 2016; Lawn 2017). Sensitivity estimates ranged from 0% to 33% and sensitivity estimates ranged from 95% to 99% (Appendix 11). Pooled sensitivity and specificity (95% CI) were 20% (10% to 35%) and 98% (95% to 99%), (952 participants, 12% with tuberculosis) (Table 2).

**LF-LAM accuracy among unselected participants, stratified by tuberculosis prevalence**

The median prevalence of tuberculosis in studies with unselected participants was 10% (IQR 9% to 17%). In the analysis by tuberculosis prevalence, we found that pooled sensitivity and specificity for unselected participants in settings with tuberculosis prevalence of 10% or more were 45% (31% to 61%) and 52% (79% to 97%) (4 studies) compared to 16% (5% to 36%) and 98% (94% to 99%) in settings with tuberculosis prevalence less than 10% (3 studies). In general, tuberculosis prevalence increased in studies with a higher proportion of symptomatic participants.

**Sensitivity analyses**

As for studies among symptomatic participants, we undertook sensitivity analyses by limiting inclusion in the meta-analysis to the following.

- Studies with low risk of bias for patient selection (three studies)
- Studies with low risk of bias in the reference standard domain (two studies)
- Studies that used fresh urine samples (five studies) rather than stored urine sample

We further excluded the two studies with more than 80% of participants being symptomatic at inclusion (Bjerrum 2015; Lawn 2017). These sensitivity analyses made little difference to any of the findings (Table 3).

**DISCUSSION**

This systematic review updates the current literature and includes 15 unique studies on the accuracy of lateral flow urine lipoolarabinomannan assay (LF-LAM, Alere Determine™ TB LAM Ag) for tuberculosis in adults living with HIV and integrates nine new studies identified since the original WHO guidelines (WHO Lipoolarabinomannan Policy Guidance 2015) and Cochrane Review (Shah 2016). A priori, we chose to evaluate performance of LF-LAM separately for studies that enrolled strictly symptomatic participants and studies that enrolled unselected participants irrespective of signs and/or symptoms of tuberculosis, and we conducted investigations of heterogeneity based on CD4 counts and study setting (inpatient/outpatient). Eight studies used LF-LAM for tuberculosis diagnosis in symptomatic adult participants with signs and symptoms suggestive of tuberculosis. These studies largely focused on inpatient settings and had high tuberculosis prevalence. Seven studies evaluated LF-LAM for diagnosing tuberculosis in unselected participants who may or may not have had signs or symptoms suggestive of tuberculosis when enrolled in the study. The studies with unselected participants were conducted predominantly in outpatient settings and, compared to studies with exclusively symptomatic participants, had lower tuberculosis prevalence and involved patients with higher CD4 counts; the proportion of symptomatic participants in these studies ranged from 19% to 90%. All studies were conducted in low- and middle-income countries with a high tuberculosis/HIV burden, and only one study outside sub-Saharan Africa.

**Summary of main results**

We have summarized the main findings in **Summary of findings 1; Summary of findings 2**. The main findings of this updated review include the following.

For tuberculosis diagnosis in HIV-positive adults presenting with signs and symptoms of tuberculosis, the diagnostic accuracy of LF-LAM is (95% credible interval (CrI)):

- In all settings, sensitivity 42% (31% to 55%) and specificity 91% (85% to 95%);
- In inpatient settings, sensitivity 52% (40% to 64%) and specificity 87% (78% to 93%);
- In outpatient settings, sensitivity 29% (17% to 47%) and specificity 96% (91% to 99%).

For tuberculosis diagnosis in HIV-positive adults irrespective of signs and symptoms of tuberculosis, the diagnostic accuracy of LF-LAM is:

- In all settings, sensitivity 35% (22% to 50%) and specificity 95% (89% to 98%);
- In inpatient settings, sensitivity 62% (41% to 83%) and specificity 84% (48% to 96%);
- In outpatient settings, sensitivity 31% (18% to 47%) and specificity 95% (87% to 99%).

We note that the band intensity of grade 1 in this review corresponds to the current manufacturer threshold for positivity (equivalent to that of grade 2 on the old manufacturer reference
Collaboration.

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card) and all results were evaluated against a microbiological reference standard.

**LF-LAM for tuberculosis diagnosis in studies with symptomatic participants**

**Summary of findings 1.**

In planned investigations of heterogeneity, we found an inverse correlation between LF-LAM sensitivity and CD4 count, with increasing sensitivity as patient CD4 count decreased (increased from 16% (CI 8% to 21%) in patients with CD4 cell count > 200 cells per μL to 24% (CI 14% to 38%) in patients with CD4 cell count between 101 to 199 cells per μL, to 54% (CI 38% to 69%) in patients with CD4 < 100 cells per μL). Similarly, we a priori planned to investigate and expected to find higher sensitivity in patients who were hospitalized while specificity decreased.

Results of these studies indicate that in theory, for a population of 1000 people where 300 have microbiologically-confirmed tuberculosis, the utilization of LF-LAM would result in: 189 to be LF-LAM positive: of these, 63 (33%) would not have tuberculosis (false-positives); and 811 to be LF-LAM negative: of these, 174 (21%) would have tuberculosis (false-negatives).

**LF-LAM for tuberculosis diagnosis in studies with unselected participants (signs and symptoms not assessed as a condition of study inclusion)**

**Summary of findings 2.**

In the investigations of heterogeneity, we expected and found a higher sensitivity in patients with CD4 ≤ 100 cells per μL (47%) and among inpatients (62%) compared to patients with CD4 > 100 cells per μL (20%) among outpatients (31%) respectively, though data to inform subgroup analyses were limited. We noted that participants included in the studies with unselected participants often presented with signs and symptoms suggestive of tuberculosis (a positive WHO tuberculosis screen), and in the studies evaluating inpatients most participants (> 80%) were in fact presenting with signs and symptoms suggestive of tuberculosis. These studies may be considered more similar to studies with exclusively symptomatic participants. In additional analysis of heterogeneity, we examined diagnostic accuracy based on tuberculosis prevalence within the studied cohort, as an alternative surrogate to presence of symptoms or CD4 count as an assessment of pre-test probability. We found that pooled sensitivity was 45% when tuberculosis prevalence within the study population was ≥ 10%, compared to only 16% when tuberculosis prevalence in the study population was < 10%.

Results of these studies indicate that in theory, for a population of 1000 people where 100 have microbiologically-confirmed tuberculosis, the utilization of LF-LAM would result in: 80 to be LF-LAM positive: of these, 45 (56%) would not have tuberculosis (false-positives); and 920 to be LF-LAM negative: of these, 65 (7%) would have tuberculosis (false-negatives).

**LF-LAM for tuberculosis diagnosis, overall**

The findings of this updated review are consistent with those of the original review (Shah 2016; WHO Lipopolysaccharide Policy Guidance 2015). Inclusion of additional studies in this updated review provided the basis for a more precise estimate of the LF-LAM overall sensitivity and specificity. It further allowed us to address key questions regarding test accuracy and sources of heterogeneity including clinical setting and CD4 cell count in studies with symptomatic individuals and in studies with unselected participants.

Overall, we found lower sensitivity for diagnosis of tuberculosis among people living with HIV than the internationally suggested target of minimum 65% overall for rapid non-sputum tuberculosis tests (WHO TTP 2014). We found that sensitivity increased when considering inpatients and individuals with lower CD4 counts, whether considering studies with exclusively symptomatic participants or those with unselected participants. The reasons for this heterogeneity remain unclear, and several hypotheses have been postulated including differences in bacterial burden, sites of tuberculosis disease, or degree of glomerular dysfunction that may be different across different settings or CD4 counts.

When restricting analysis to studies that included participants unable to produce a sputum sample, the estimates of sensitivity increased. Sputum-scarce patients may be the potential target population to benefit the most from urine-based testing as they cannot have other sputum-based diagnostic testing and are likely to have a high yield of urine lipopolysaccharide (LAM) test positivity (Sabur 2017). However, only a few studies included participants who could not provide sputum samples for diagnostic testing. To the extent that inability to produce sputum is correlated with severity of tuberculosis disease and/or LAM positivity, this approach to participant selection could have lowered sensitivity estimates within these studies. Sensitivity analysis further revealed a higher sensitivity among studies evaluating LF-LAM on fresh non-stored urine samples without it affecting specificity. However, no study has made a direct comparison of performance on fresh versus frozen/stored urine samples and the significance of this is unclear.

Overall, we found that the estimated specificity approached the recommended targets for non-site-specific, non-sputum based test (WHO TTP 2014), although lower specificity was found among inpatients and those with advanced immunosuppression compared to outpatients and those with higher CD4 counts. We expected that, if restricting the analysis to studies using a higher quality reference standard (e.g. inclusion of more than one specimen type), that estimates of specificity would increase, but had limited data to conduct such a sensitivity analysis. When evaluating an assay that is not site-specific, future studies may consider systematic sampling of multiple sites to improve the quality of the reference standard. Future studies should include systematic sampling of sputum, blood, and urine for mycobacterial culture and/or nucleic acid amplification test (NAAT) testing from all participants as common sites of tuberculosis that may be identified by urinary LAM antigen detection; additional site-specific samples (e.g. cerebrospinal fluid (CSF), tissue biopsies) should also be collected for culture and/or NAAT based on clinical presentation. We decided a priori to evaluate performance of LF-LAM in HIV-positive individuals with signs and symptoms of tuberculosis (symptomatic) separately from HIV-positive individuals irrespective of signs and symptoms of tuberculosis (unselected participants). We considered evaluating LF-LAM performance among specifically asymptomatic (i.e. exclusively those without symptoms) participants to assess the use of LF-LAM for the purpose of tuberculosis screening or community-based active case-finding, but such data were lacking among included
studies. We did find that several studies among unselected participants reported that a high proportion of study participants had signs and symptoms of tuberculosis, suggesting relative similarities to studies that enrolled exclusively symptomatic participants. Consequently, the overall performance of LF-LAM among asymptomatic patients remains largely unknown.

The overall differences in pooled estimates of sensitivity and specificity between studies of symptomatic versus unselected participants may have been attributable to differences in study setting and relative degree of immunosuppression of included participants, rather than type of population (i.e. unselected versus symptomatic participants). When examining inpatients, the pooled estimates for sensitivity were 52% (40% to 64%) and 62% (41% to 83%), when comparing studies of symptomatic participants and those including unselected participants. Among outpatients, the pooled sensitivity was 29% (17% to 47%) compared to 31% (18% to 47%) among studies of symptomatic participants and unselected participants, respectively. This indicates that other characteristics than lower CD4 may explain the higher sensitivity among inpatients like higher tuberculosis prevalence, higher mycobacterial burden, renal or genitourinary tract tuberculosis with LAM secretion in urine.

Overall, our findings suggest that the diagnostic accuracy of LF-LAM may vary by study setting, CD4 count, and tuberculosis prevalence among the target population. The authors hypothesize that these attributes (inpatients, low CD4 counts, or high tuberculosis prevalence) may collectively be surrogate indicators of participants with advanced tuberculosis disease or higher bacillary burden and LAM antigenemia in whom LF-LAM may aid in the diagnosis of tuberculosis, including both pulmonary and extrapulmonary tuberculosis. Although subgroup comparisons in diagnostic accuracy reviews are observational and suffer from the same limitations as all observational findings (for example, confounding between characteristics), there is a scientific rationale for these findings in that inpatients, those with low CD4, or cohorts with higher tuberculosis prevalence are likely to have higher disease severity or higher bacillary burden. While the test does not identify all tuberculosis cases, our findings suggest that it may be of particular value in diagnosing tuberculosis among patients with increased disease severity. Other factors that may be considered in evaluating LF-LAM may include ability to perform the test on individuals unable to produce sputum who cannot be diagnosed with other tuberculosis diagnostic tests, and ability to implement the test at the point-of-care with non-invasive specimen collection (WHO TTP 2014).

We identified three published studies of LF-LAM in children as the result of a broader search for studies in adults and children using the same inclusion criteria (Kroidl 2015; LaCourse 2018a; Nicol 2014). All three studies took place in countries in Africa with a high tuberculosis/HIV burden. Prevalence of microbiologically-confirmed tuberculosis in the studies was 7% in LaCourse 2018a, 22% in Nicol 2014; and 40% in Kroidl 2015. Kroidl 2015 enrolled children six weeks to 14 years, median age (interquartile range (IQR)) 6.8 years (3.9 to 9.5) for all participants, including HIV-positive and HIV-negative children. LaCourse 2018a enrolled children aged 12 years or less, median age (IQR) 24 months (13 to 58). Nicol 2014 enrolled children aged 15 years or less, median age (IQR) 42.5 months (19.1 to 66.3) for all participants, including HIV-positive and HIV-negative children. A table summarizing the characteristics of studies in children can be found in Appendix 12. Kroidl 2015 and Nicol 2014 involved HIV-positive children with tuberculosis symptoms. LaCourse 2018a involved HIV-positive children hospitalized for acute illness irrespective of tuberculosis signs and symptoms. Kroidl 2015 was conducted in an outpatient setting. LaCourse 2018a in an inpatient setting, and Nicol 2014 in both an inpatient and an outpatient setting (Appendix 12). Given the differences in population and setting, we did not perform meta-analyses and report sensitivity and specificity estimates for the individual studies, all measured against a microbiological reference standard. Sensitivity and specificity (95% CI) were 42% (15% to 72%) and 94% (73% to 100%), (30 participants) (Kroidl 2015); 56% (21% to 86%) and 95% (90% to 98%), (130 participants) (LaCourse 2018a); and 43% (23% to 66%) and 80% (69% to 88%), (106 participants) (Nicol 2014). We found limited evidence on the accuracy of LF-LAM in children living with HIV. There were too few studies and participants to draw conclusions.

We acknowledge that patient outcomes are clearly important to patients, decision makers, and the wider tuberculosis community. While, the primary focus of this Cochrane Review update was to evaluate the diagnostic accuracy of LF-LAM, in the process of performing the review, we also assessed available data on LF-LAM and patient-important outcomes including mortality. We identified two randomized controlled trials (Gupta-Wright, 2018a; Peter 2016) that included data on the impact of LF-LAM implementation on mortality and other patient outcomes. Peter 2016 enrolled participants with tuberculosis symptoms, and Gupta-Wright, 2018a enrolled unselected participants irrespective of symptoms. Both trials were conducted in multiple countries in sub-Saharan Africa. Both trials involved hospitalized, HIV-positive patients, used the results of LF-LAM to guide therapy, and assessed all-cause mortality at eight weeks. Peter 2016 found that mortality was significantly reduced in participants receiving LF-LAM compared to those receiving routine diagnostic tests alone (no LF-LAM); adjusted relative risk reduction of 0.83 (95% CI: 0.73 to 0.96, P = 0.012). In Gupta-Wright, 2018a, the use of rapid urine screening with LF-LAM and urine Xpert MTB/RIF (intervention group) was not associated with a statistically significant reduction in mortality compared with routine diagnostic tests alone (standard-of-care group, no LF-LAM); adjusted risk reduction (aRD) of -2.8% (95% CI -5.8 to 0.3, P = 0.074). However, of importance, Gupta-Wright, 2018a, demonstrated that mortality was significantly lower in the intervention group than in the standard-of-care group for three of 12 prespecified subgroups: CD4 counts less than 100 cells per μL (aRD -7.1% (95% CI -13.7 to -0.4), P = 0.036); severe anaemia (aRD -9.0% (95% CI -16.6 to -1.4), P = 0.021); and participants with clinically suspected tuberculosis (aRD -5.7% (95% CI -10.9 to -0.5), P = 0.033). We will address the impact of testing with LF-LAM on patient outcomes in greater depth in another Cochrane Review (a Cochrane Protocol is underway).

Concerning economic evaluations, we are aware of six studies, all from settings in sub-Saharan Africa (Boyles 2018; Mukora 2018; Orlando 2018; Reddy 2019; Shah 2013; Sun 2013). These study methods and populations were heterogeneous, assessing a range of diagnostic algorithms, and only four of the total studies assessed cost-effectiveness (Orlando 2018; Reddy 2019; Shah 2013; Sun 2013), while Mukora 2018 performed a detailed costing study, including both at clinical and above clinic level costs, and Boyles 2018 assessed whether the addition of LF-LAM in diagnostic algorithms could reduce total program costs. studies employed a range of different willingness to pay thresholds, all four studies
were consistent in concluding LF-LAM could be cost-effective in a population of African adults living with HIV (particularly amongst hospitalized patients). Models found cost-effectiveness of LF-LAM to be robust across a variety of sensitivity analyses, and across different country settings and scenarios investigated. Key parameters that are likely most influential on cost-effectiveness include: tuberculosis prevalence/target population, and LF-LAM specificity, cost of treating tuberculosis and HIV and life expectancy post tuberculosis survival. Detailed empirical costing data are limited for LF-LAM implementation, however one such study published in 2018 (Mukora 2018) estimates unit test costs for LF-LAM implementation several fold higher (~$23-24) than most current models ($2-4), where costs of implementation and staff time are not considered. Underestimation of LF-LAM unit costs could result in overly optimistic cost-effectiveness profiles. However, tuberculosis diagnostics costs likely represent just a small proportion of total costs, indeed inclusion of costs associated with antiretroviral treatment (ART) and HIV care resulted in higher incremental cost-effectiveness ratios. While current evidence is consistent in suggesting LF-LAM is likely cost-effective among HIV-positive patients in sub-Saharan Africa, caution should be used when extrapolating from a small number of studies, and additional evidence from a wider range of populations, settings and diagnostic approaches will be necessary. A systematic review of economic evaluations is underway.

**Differences from original Cochrane Review**

In comparison to the original Cochrane Review (Shah 2016), this updated review includes 15 published studies (nine new studies since the original review and six studies from the original review). The original review included data from 12 studies, of which six were excluded as they did not meet the eligibility criteria for this updated review. Three used an older threshold for determining test positivity that is no longer recommended and three were abstracts, of which one (Lawn 2014) was added in this review as an updated published manuscript. For comparison, the main findings from the original review (Shah 2016) are listed in Table 4.

In the evaluation of diagnostic accuracy among symptomatic participants, the pooled estimates for sensitivity (42% [Crl 31% to 55%] versus 45% [Crl 29% to 63%]) and specificity (91% [Crl 85% to 95%] versus 92% [Crl 80% to 97%]) remained similar, comparing the current review and prior review, respectively. When stratified by setting, the pooled estimates among inpatients for sensitivity (52% [Crl 40% to 64%] versus 53% [Crl 38% to 70%]) and specificity (87% [Crl 78% to 93%] versus 90% [Crl 73% to 96%]) did not change substantially when comparing the current review and the prior review, respectively. Pooled estimates among outpatients at the current manufacturer threshold for positivity were not previously available.

In Shah 2016, some studies were classified as ‘tuberculosis (TB) screening’ if they included participants irrespective of symptoms (i.e. with or without symptoms). Recognizing that these studies may have included a large proportion of symptomatic participants, these studies have been more clearly labelled as studies among ‘unselected participants’ in the current review. In the prior review, there were insufficient data to perform meta-analysis among unselected participants at the currently recommended manufacturer threshold for test positivity. There were previously insufficient data to investigate heterogeneity due to study setting or CD4 count. In the current review, we report on diagnostic accuracy among inpatients and outpatients and by CD4. In both the current and prior review, data to assess diagnostic accuracy among asymptomatic participants (without signs or symptoms of tuberculosis) were unavailable.

This updated review did not assess diagnostic accuracy at an older threshold for determining test positivity (grade 1 out of 5, on an older reference card); data on diagnostic accuracy for this threshold can be found in the prior review (Shah 2016). Similarly, in this updated review we did not evaluate accuracy against a composite reference standard, results of which were included in the prior review (Shah 2016). Finally, given the relative lack of data on evaluating the incremental yield of LF-LAM in combination with sputum smear microscopy and Xpert MTB/RIF, we did not include these analyses in the updated review, but a summary of available data is found in the prior review (Shah 2016).

We note that the band intensity of grade 1 in this review corresponds to the current manufacturer threshold for positivity (equivalent to that of grade 2 on the old manufacturer reference card) and all results were evaluated against a microbiological reference standard.

**Strengths and weaknesses of the review**

The findings in this review are based on comprehensive searching, strict inclusion criteria, and standardized data extraction. The strength of our review is that it enabled an assessment of the accuracy of LF-LAM in people living with HIV when applied to different enrolment criteria - accuracy among individuals with signs and symptoms of tuberculosis, and accuracy when used without assessment of signs and symptoms of tuberculosis (i.e. patients may or may not have had symptoms). This updated review included new studies published since the original review. However, we found considerable heterogeneity across studies with respect to clinical setting, CD4 count and tuberculosis prevalence. For some analyses, few studies and participants contributed data and results should, therefore, be interpreted with caution. The review was further limited by the number of studies that used a lower quality reference standard and the high risk of selection bias in several studies owing to exclusion of patients unable to produce sputum.

**Completeness of evidence**

This data set involved comprehensive searching and correspondence with experts in the field and the test manufacturer to identify additional studies, as well as correspondence with study authors to obtain additional and unpublished data on study characteristics. The search strategy included studies published in all languages. However, as diagnostic accuracy studies are poorly indexed, we acknowledge that we may have missed some studies despite the comprehensive search.

**Accuracy of the reference standards used**

In a diagnostic test accuracy systematic review, the reference standard is the best available test to determine the presence or absence of the target condition. We only included studies with a microbiological reference standard, which is considered the best currently available reference standard for tuberculosis. We included studies that evaluated LF-LAM for diagnosis of pulmonary tuberculosis, extrapulmonary tuberculosis, or both pulmonary and extrapulmonary tuberculosis. However, we recognize that a substantial number of tuberculosis cases may not be verified by...
microbiological testing if only sputum is tested and when patients with advanced HIV are assessed. We acknowledge difficulties in diagnosing HIV-associated tuberculosis with extrapulmonary and disseminated forms of disease and considered a standardized reference standard using two or more specimen types consistently for all participants to be of higher quality than a reference standard using one specimen type. The higher quality reference standard is better at classifying which patients have and do not have tuberculosis. A lower quality reference standard may miss some tuberculosis cases and classify some tuberculosis patients as not having tuberculosis. This may make a truly positive LF-LAM result seem like a false-positive result leading to an underestimation of specificity. In this review, only one study (12%) for tuberculosis diagnosis among symptomatic participants and two studies (29%) for tuberculosis diagnosis among unselected participants used a higher quality reference standard for all participants.

We did not assess performance against a composite reference standard that uses microbiological or clinical information to classify tuberculosis. This was done in the original WHO guideline (WHO Liparabinomannan Policy Guidance 2015), and Cochrane Review (Shah 2016), but found little impact on pooled estimates of sensitivity and specificity relative to performance measured against a microbiological reference standard.

We could not determine whether heterogeneity in specificity estimates was fully attributable to misclassification bias. Some studies (Nel 2017; Qvist 2014) have postulated that infection with (disseminated) nontuberculous mycobacteria (NTM) may also result in false-positive results, although this hypothesis is still questioned (Gupta-Wright 2018). Only one study (Thit 2017) was conducted outside of sub-Saharan Africa, and was noted to report the lowest specificity estimates of all included studies. Reasons for potential false-positive results remain unclear and it is unknown if differences in the epidemiology of disseminated NTM and other opportunistic infections across settings could contribute to variation in specificity.

Quality and quality of reporting of the included studies

There were few studies that included participants unable to expectorate sputum and few studies that included extrapulmonary specimens in addition to sputum. We had limited data to address these quality items in sensitivity analyses and acknowledge that these features may have contributed to risk of bias in the accuracy estimates. For studies among symptomatic participants, in the patient selection domain, we considered six studies (75%) to be at high risk of bias, and in the reference standard domain, we considered seven studies (88%) to be at high risk of bias. For studies with unselected participants, in the patient selection domain, we considered four studies (57%) to be at high risk of bias and in the reference standard domain, we considered five studies (71%) to be at high risk of bias. Risk of bias was considered low for the index test and flow and timing domains.

Using the GRADE approach, we judged the evidence for LF-LAM sensitivity to be moderate. This means that we are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different. We judged the evidence for LF-LAM specificity to be low or very low. This means that we have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of effect.

In general, studies were fairly well reported, though we corresponded with some of the primary study authors to ask for clarification mainly in relation to interpretation of test positivity.

Applicability of findings to the review question

Overall, we had low concern about the applicability of the included studies to our review question as assessed by Quality Assessment of Diagnostic Accuracy Studies-2 (QUADAS-2). However, studies of HIV-positive adults irrespective of signs and symptoms had low representation of asymptomatic individuals, as most participants in fact presented with signs and symptoms of tuberculosis. To date, no study has evaluated LF-LAM in an asymptomatic population. This review evaluated LF-LAM at the current manufacturer-recommended cut-off for positivity. Therefore, the findings should be considered applicable to the test. However, it is important to note that this review assessed sensitivity and specificity in applied research settings. Although the participant characteristics and settings matched our review question in most cases, as studies were carried out under research conditions, it is possible that the accuracy of LF-LAM may be lower in routine practice settings. All studies were performed in low- or middle-income countries, but only a single study was conducted outside sub-Saharan Africa. It is therefore possible that the LF-LAM would perform differently in settings outside Africa. For the reference test domain, most studies had low concern for applicability.

Authors’ conclusions

Implications for practice

For people living with HIV, this review found overall lower sensitivity of the lateral flow urine lipoarabinomannan assay (LF-LAM) than the internationally suggested target of minimum 65% overall for non-sputum based tuberculosis diagnostic tests (WHO 2014). This was consistent whether the test is used for diagnosis of HIV-associated tuberculosis (without consideration of sites of disease involvement) among symptomatic (sensitivity of 42%) or unselected (i.e. not assessed for symptoms) participants (sensitivity of 35%). The estimated sensitivity suggests that if LF-LAM were to be used alone, more than half of all tuberculosis cases would be missed. Whether diagnostic accuracy differs for specific sites of tuberculosis is unknown (e.g. pulmonary tuberculosis, tuberculosis meningitis, or lymph node tuberculosis). There was a lack of data to evaluate diagnostic accuracy for active tuberculosis in asymptomatic participants, but studies among unselected outpatients that included a combination of symptomatic and asymptomatic participants (i.e. unselected participants) showed lower sensitivity (31%).

Despite the estimated sensitivity, two randomized controlled trials implementing LF-LAM in high-prevalence settings in sub-Saharan Africa have demonstrated reduced mortality and impact on other clinical outcomes when used to guide tuberculosis treatment in hospitalized HIV-positive adults.

LF-LAM is being considered as a diagnostic test that may be used as an add on to clinical judgement and with other tests for the diagnosis of HIV-associated tuberculosis. The test does not have a role as a replacement or triage test. The test may improve important health outcomes among HIV-positive hospitalized patients because it is feasible (i.e. can be done on urine which is easy to obtain in an inpatient) and may yield a positive result in those that are
most likely to die from tuberculosis and are most likely to present atypically (i.e. patients with advanced HIV). Whether these results may be extrapolated to outpatients with advanced HIV disease or those that are seriously ill is unknown.

Findings suggest that sensitivity increases with lower CD4 counts and in inpatient settings compared to outpatient settings. For unselected participants irrespective of symptoms, we found differences in LF-LAM performance based on tuberculosis prevalence of the target population (when stratified at greater or less than 10%), with higher sensitivity when the study population had higher tuberculosis prevalence (≥ 10%). The difference in diagnostic accuracy of LF-LAM by study setting and by degree of immunosuppression was consistent regardless of the approach to patient selection. Taking all findings into account, we think LF-LAM has a role along with other tests among individuals with tuberculosis symptoms or with low CD4 cell count. However, we had limited data and these findings should be interpreted with caution.

The test does not require sputum collection and is not site-specific. Other favourable test characteristics include low-cost, rapidity (< one hour), ease of use (does not require extensive sample preparation), and the fact that the test does not require electricity or special instruments and equipment (WHO TTP 2014). As a simple point-of-care test that does not depend upon sputum evaluation, LF-LAM testing may be the only possible way to confirm a diagnosis when a sputum sample cannot be produced.

Clinicians must consider the need for additional testing when interpreting negative LF-LAM results. The consequences of false-negative results are increased risk of morbidity and mortality, delayed treatment initiation, and the continued risk of tuberculosis transmission. The consequences of false-positive results are delayed alternative diagnosis, likelihood of anxiety and morbidity caused by additional testing, unnecessary treatment, and possible adverse events; the possible stigma associated with a diagnosis of tuberculosis. As LF-LAM does not offer information about drug resistance, a culture- or molecular-based diagnosis should be attempted to enable drug susceptibility testing to avoid that patients with unidentified drug-resistant tuberculosis may be inappropriately treated with a regimen appropriate only for drug-susceptible disease.

Ultimately, the use of LF-LAM in a clinical algorithm will not be based on accuracy alone, but also on considerations of cost-effectiveness, user perspectives and preferences, and the overall benefits and harms of the test. This updated review contributed accuracy data for the WHO Guideline Development Group Meeting in May 2019. The updated WHO guidelines on the use of Alere LAM are expected later in 2019.

Implications for research

Future studies that evaluate the diagnostic accuracy of non-sputum-based tests for tuberculosis, such as LF-LAM, in people living with HIV should use a reference standard that includes at least two different specimens (e.g. sputum, and urine), and in addition, for presumed extrapulmonary tuberculosis, appropriate specimens from the suspected sites of involvement. Moreover, future studies should include patients unable to expectorate sputum in the analysis as these are most likely to benefit from the test. Characteristics that possibly drive performance of the test in addition to CD4 status, such as burden of disease, renal co-morbidities or other should be evaluated. More studies should evaluate LF-LAM accuracy in HIV-positive children, as currently there are very limited data available (Kroidl 2015; LaCourse 2018a; Nicol 2014). While some studies enrolled unselected participants, our review suggests that a large proportion were symptomatic, particularly in the inpatient setting. Understanding whether there is value in using LF-LAM for diagnosis of tuberculosis among HIV-positive people who do not report any tuberculosis symptoms (particularly for those with CD4 < 100) remains unknown. Further studies among asymptomatic individuals are needed to better define the scope of recommendations on the use of LF-LAM. The indication of increased sensitivity with use of fresh urine and also morning urine needs further investigations, and studies in settings outside sub-Saharan Africa are lacking. Further research on effective implementation of LF-LAM within routine clinical practice is needed because the test can only influence clinical practice if the results are believed and acted upon.

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Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV (Review)

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**CHARACTERISTICS OF STUDIES**

**Characteristics of included studies [ordered by study ID]**

**Bjerrum 2015**

**Study characteristics**

| Patient sampling | Cross-sectional, prospective and consecutive enrolment |
|------------------|--------------------------------------------------------|
| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected adults included irrespective of symptoms. A positive WHO symptom screen was reported for 91% of participants. |
| Age | mean 39 (SD10) |
| Sex, female | 64% |
| HIV infection | 100% |
| Median CD4 cell count per μL | 127 (IQR 35 to 256) |
| History of tuberculosis | 6% |
| Sample size | 469 |
| Clinical setting | outpatient and inpatients |
| Country | Ghana |
| Tuberculosis incidence rate | 152 per 100,000 |
| Number (proportion) of tuberculosis cases in the study | 55 (12%) |

**Index tests**

| Target condition and reference standard(s) | LF-LAM |
|-------------------------------------------|--------|
| Target condition: pulmonary tuberculosis | |
| Reference standard: any of sputum mycobacterial culture (solid and liquid) or nucleic acid amplification test; no extrapulmonary specimens tested | |

**Flow and timing**

| Comparative | All patients were included in the analysis |

**References to other published versions of this review**

**Shah 2014**
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Shah M, Hanrahan C, Wang ZY, Dendukuri N, Lawn SD, Denkinger CM, et al. Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in HIV-positive adults. *Cochrane Database of Systematic Reviews* 2016, Issue 5. [DOI: 10.1002/14651858.CD011420.pub2]
The study authors did not attempt sputum induction and excluded individuals without some respiratory specimen available. There was no attempt to identify extrapulmonary tuberculosis.

### Methodological quality

| Item                                      | Authors' judgement | Risk of bias | Applicability concerns |
|-------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**           |                    |              |                        |
| Was a consecutive or random sample of patients enrolled? | Yes               |              |                        |
| Was a case-control design avoided?        | Yes               |              |                        |
| Did the study avoid inappropriate exclusions? | No                | High         | Low                    |
| **DOMAIN 2: Index Test All tests**        |                    |              |                        |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes               |              |                        |
| If a threshold was used, was it pre-specified? | Yes               | Low          | Low                    |
| **DOMAIN 3: Reference Standard**          |                    |              |                        |
| Is the reference standards likely to correctly classify the target condition? | No                |              |                        |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes               | High         | Low                    |
| **DOMAIN 4: Flow and Timing**             |                    |              |                        |
| Was there an appropriate interval between index test and reference standard? | Yes               |              |                        |
| Did all patients receive the same reference standard? | Yes               |              |                        |
| Were all patients included in the analysis? | Yes               | Low          |                        |

### Drain 2015a

**Study characteristics**
Patient sampling
Cross-sectional, prospective and consecutive enrolment

Patient characteristics and setting
Presenting signs and symptoms: HIV-infected adults included irrespective of symptoms. Proportion of participants that reported with tuberculosis signs and symptoms symptomatic not stated.
Age: mean 33 (SD 9)
Sex, female: 49%
HIV infection: 100%
Median CD4 cell count per μL: 248 (IQR 107 to 379)
History of tuberculosis: 9%
Sample size: 320
Clinical setting: outpatient
Country: South Africa
Tuberculosis incidence rate: 567 per 100,000
Number (proportion) of tuberculosis cases in the study: 54 (17%)

Index tests
LF-LAM

Target condition and reference standard(s)
Target condition: pulmonary tuberculosis
Reference standard: mycobacterial culture of sputum (solid); no extrapulmonary specimens tested

Flow and timing
All patients were included in the analysis

Comparative

Notes
The study authors performed sputum induction for individuals who could not expectorate, excluded individuals without some respiratory specimen available. They did not attempt to identify extrapulmonary tuberculosis.

Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection** | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | No | | |
| **DOMAIN 2: Index Test All tests** | | | |

| | | |
| | | |
| | | |
### Drain 2015a (Continued)

| Question                                                                 | Answer | Risk of Bias |
|--------------------------------------------------------------------------|--------|--------------|
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes    | Low          |
| If a threshold was used, was it pre-specified?                           | Yes    | Low          |

#### DOMAIN 3: Reference Standard

| Question                                                                 | Answer | Risk of Bias |
|--------------------------------------------------------------------------|--------|--------------|
| Is the reference standards likely to correctly classify the target condition? | No     | Low          |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes    | High         |

#### DOMAIN 4: Flow and Timing

| Question                                                                 | Answer | Risk of Bias |
|--------------------------------------------------------------------------|--------|--------------|
| Was there an appropriate interval between index test and reference standard? | Yes    | Low          |
| Did all patients receive the same reference standard?                     | Yes    | Low          |
| Were all patients included in the analysis?                               | Yes    | Low          |

### Drain 2016

#### Study characteristics

| Patient sampling | Cross-sectional, prospective and consecutive enrolment |
|------------------|--------------------------------------------------------|
| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected and HIV-uninfected adults suspected of tuberculosis included if smear microscopy negative x 2; Presenting with at least two of four tuberculosis-related symptoms: cough, fever, weight loss, night sweats for at least two weeks and a Karnofsky performance score >50 Age: mean 37 (SD 9) Sex, female: 49% HIV infection: 93% Median CD4 cell count per μL: 168 (IQR 89 to 256) History of tuberculosis: not stated Sample size: 90 Clinical setting: outpatient Country: South Africa Tuberculosis incidence rate: 567 per 100,000 Number (proportion) of tuberculosis cases in the study: 57 (63%) |
| Index tests | LF-LAM |
| Target condition and reference standard(s) | Target: pulmonary tuberculosis Reference standard: any of sputum mycobacterial culture (solid and liquid) or nucleic acid amplification test; no extrapulmonary specimens tested |
Flow and timing

All patients were included in the analysis

Comparative

Notes

The study authors performed sputum induction for individuals who could not expectorate. They excluded individuals without some respiratory specimen available and did not attempt to identify extrapulmonary tuberculosis

**Methodological quality**

| Item                                                                 | Authors' judgement | Risk of bias | Applicability concerns |
|----------------------------------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**                                      |                    |              |                        |
| Was a consecutive or random sample of patients enrolled?             | Yes                | High         | High                   |
| Was a case-control design avoided?                                   | Yes                |              |                        |
| Did the study avoid inappropriate exclusions?                        | No                 | High         | High                   |
| **DOMAIN 2: Index Test All tests**                                   |                    | Low          | Low                    |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes                |              |                        |
| If a threshold was used, was it pre-specified?                       | Yes                | Low          | Low                    |
| **DOMAIN 3: Reference Standard**                                     |                    |              |                        |
| Is the reference standards likely to correctly classify the target condition? | No                 | High         | Low                    |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes                |              |                        |
| **DOMAIN 4: Flow and Timing**                                        |                    | Low          |                        |
| Was there an appropriate interval between index test and reference standard? | Yes                |              |                        |
| Did all patients receive the same reference standard?                | Yes                |              |                        |
| Were all patients included in the analysis?                          | Yes                |              |                        |
### Study characteristics

**Patient sampling**
Cross-sectional, prospective and consecutive enrolment

**Patient characteristics and setting**
Presenting signs and symptoms: HIV-infected adults included irrespective of symptoms. A positive WHO symptom screen was reported for 34% of participants. 
- Age: median 35 (IQR 30-43)
- Sex, female: 59%
- HIV infection: 100%
- Median CD4 cell count per μL: 278 (IQR 142 to 395)

History of tuberculosis: not stated
- Sample size: 972
- Clinical setting: outpatients
- Country: Mozambique
- Tuberculosis incidence rate: 551 per 100,000
- Number (proportion) of tuberculosis cases in the study: 90 (9%)

### Index tests
LF-LAM

**Target condition and reference standard(s)**
- Target: pulmonary tuberculosis
- Reference standard: nucleic acid amplification test; no extrapulmonary specimens tested

### Flow and timing
All patients were included in the analysis

### Comparative

### Notes
The study authors did not attempt sputum induction and excluded individuals without some respiratory specimen available. They did not attempt to identify extrapulmonary tuberculosis.

### Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection** | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | No | | |
| **DOMAIN 2: Index Test All tests** | | | |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes | | |
| If a threshold was used, was it pre-specified? | Yes | | |
### Floridia 2017 (Continued)

| DOMAIN 3: Reference Standard | Low | Low |
|------------------------------|-----|-----|
| Is the reference standards likely to correctly classify the target condition? | No |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear |

| DOMAIN 4: Flow and Timing | High | Low |
|----------------------------|------|-----|
| Was there an appropriate interval between index test and reference standard? | Yes |
| Did all patients receive the same reference standard? | Yes |
| Were all patients included in the analysis? | Yes |

### Hanifa 2016

| Study characteristics |
|-----------------------|
| **Patient sampling** | Cohort, prospective and consecutive enrolment with six months follow-up |
| **Patient characteristics and setting** | Presenting signs and symptoms: HIV-infected adults included irrespective of symptoms. A positive WHO symptom screen was reported for 53% of participants. |
| | Age: median 39 (IQR 32-45) |
| | Sex, female: 61% |
| | HIV infection: 100% |
| | Median CD4 cell count per µL: 111 (IQR 56 to 161) |
| | History of tuberculosis: 30% |
| | Sample size: 424 |
| | Clinical setting: outpatients |
| | Country: South Africa |
| | Tuberculosis incidence rate: 567 per 100,000 |
| | Number (proportion) of tuberculosis cases in the study: 40(9%) |
| **Index tests** | LF-LAM |
| **Target condition and reference standard(s)** | Target condition: pulmonary tuberculosis, extrapulmonary tuberculosis, mycobacteraemia |
| | Reference standard: any of mycobacterial culture (liquid) or nucleic acid amplification test; extrapulmonary specimens tested for all |
| **Flow and timing** | The study excluded 16 cases defined as 'Clinical tuberculosis' from analysis. |
| **Comparative** |  |
Sputum and non-sputum samples were tested for mycobacteria for all participants; The study authors performed sputum induction for individuals who could not expectorate; The study had specimens collected for index test and reference standard tests greater than seven days apart.

### Methodological quality

| Item                                           | Authors' judgement | Risk of bias | Applicability concerns |
|------------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**                |                    |              |                        |
| Was a consecutive or random sample of patients enrolled? | Yes                | Low          | Low                    |
| Was a case-control design avoided?             | Yes                | Low          | Low                    |
| Did the study avoid inappropriate exclusions?  | Unclear            | Low          | Low                    |
| **DOMAIN 2: Index Test All tests**             |                    |              |                        |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes                | Low          | Low                    |
| If a threshold was used, was it pre-specified? | Yes                | Low          | Low                    |
| **DOMAIN 3: Reference Standard**               |                    |              |                        |
| Is the reference standards likely to correctly classify the target condition? | Yes                | Low          | Low                    |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes                | Low          | Low                    |
| **DOMAIN 4: Flow and Timing**                  |                    |              |                        |
| Was there an appropriate interval between index test and reference standard? | No                 | High         |                        |
| Did all patients receive the same reference standard? | Yes                |              |                        |
| Were all patients included in the analysis?    | No                 |              |                        |
### Study characteristics

**Patient sampling**  
Cross-sectional, prospective and consecutive enrolment

**Patient characteristics and setting**  
Presenting signs and symptoms: HIV-infected adults suspected of tuberculosis; Presenting with cough for more than 2 weeks or any cough and at least any of loss of weight, night sweats, or fever  
Age: median 35 (IQR:29-44)  
Sex, female: 52%  
HIV infection: 100%  
Median CD4 cell count per µL: 109 (IQR 43 to 214)

History of tuberculosis: 25%  
Sample size: 474 (275 included in analysis)  
Clinical setting: outpatient and inpatients  
Country: Kenya  
Tuberculosis incidence rate: 319 per 100,000  
Number (proportion) of tuberculosis cases in the study: 156 (57%)

**Index tests**  
LF-LAM

**Target condition and reference standard(s)**  
Target: pulmonary and extrapulmonary tuberculosis.  
Reference standard: any of mycobacterial culture (solid and liquid) or nucleic acid amplification test. Extrapulmonary specimens collected for participants without respiratory samples available

**Flow and timing**  
The study excluded participants with at least one sample with no result or contaminated result (culture) from analysis in the absence of a positive result.

### Comparative

**Notes**  
The study authors performed sputum induction for individuals who could not expectorate. They included individuals without respiratory specimen available and attempted to identify extrapulmonary tuberculosis; Not all participants received the same reference standard and not all participants were included in the analysis.

### Methodological quality

| Item | Authors’ judgement | Risk of bias | Applicability concerns |
|------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection** | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | | |
| Was a case-control design avoided? | Yes | | |
| Did the study avoid inappropriate exclusions? | No | | |
| | | High | Low |
| **DOMAIN 2: Index Test All tests** | | | |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes | | |
### Huerga 2017 (Continued)

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

| Low | Low |
|-----|-----|

#### DOMAIN 3: Reference Standard

**Is the reference standards likely to correctly classify the target condition?**  
No

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

| High | Low |
|------|-----|

#### DOMAIN 4: Flow and Timing

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

**Did all patients receive the same reference standard?**  
No

**Were all patients included in the analysis?**  
No

| High |
|------|

---

### Juma 2017

#### Study characteristics

| Patient sampling | Cross-sectional, prospective and consecutive enrolment |
|------------------|-------------------------------------------------------|

| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected and HIV-uninfected adults (aged above 14 years) suspected of extrapulmonary tuberculosis.  
Age: 43 (SD: 14.8)  
Sex, female: 46%  
HIV infection: 68%  
Median CD4 cell count per μL: not stated  
History of tuberculosis: 6%  
Sample size: 99 (67 with HIV included)  
Clinical setting: inpatients  
Country: Kenya  
Tuberculosis incidence rate: 319 per 100,000  
Number (proportion) of tuberculosis cases in the study: 29 (33%) |

| Index tests | LF-LAM |
|-------------|--------|

| Target condition and reference standard(s) | Target: extrapulmonary tuberculosis, any site  
Reference standard: any of mycobacterial culture (solid and liquid) or nucleic acid amplification test. No sputum specimens tested |
|---------------------------------------------|-----------------------------------------------------|

| Flow and timing | All patients were included in the analysis |
|-----------------|-----------------------------------------|
The study excluded participants with suspected pulmonary tuberculosis and the reference standard was restricted to extrapulmonary specimens.

### Methodological quality

| Item                                                                 | Authors' judgement | Risk of bias | Applicability concerns |
|----------------------------------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**                                       |                    |              |                        |
| Was a consecutive or random sample of patients enrolled?              | Yes                |              |                        |
| Was a case-control design avoided?                                    | Yes                |              |                        |
| Did the study avoid inappropriate exclusions?                         | No                 | High         | Low                    |
| **DOMAIN 2: Index Test All tests**                                    |                    |              |                        |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes                |              |                        |
| If a threshold was used, was it pre-specified?                        | Yes                | High         | High                   |
| **DOMAIN 3: Reference Standard**                                      |                    |              |                        |
| Is the reference standards likely to correctly classify the target condition? | No                 |              |                        |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes                | High         | Low                    |
| **DOMAIN 4: Flow and Timing**                                         |                    |              |                        |
| Was there an appropriate interval between index test and reference standard? | Yes                |              |                        |
| Did all patients receive the same reference standard?                 | No                 |              |                        |
| Were all patients included in the analysis?                           | Yes                |              |                        |
| **Low**                                                              |                    |              |                        |
**LaCourse 2016 (Continued)**

**Patient sampling**

Cross-sectional, prospective and consecutive enrolment

**Patient characteristics and setting**

Presenting signs and symptoms: HIV-infected pregnant women attending antenatal care was included irrespective of symptoms. A positive WHO symptom screen was reported for 19% of participants.

Other characteristics: pregnant women

- Age: median 25 (IQR 22 to 30)
- Sex, female: 100%
- HIV infection: 100%
- Median CD4 cell count per μL: 437 (IQR 342 to 565)
- History of tuberculosis: 9%
- Sample size: 266
- Clinical setting: outpatient
- Country: Kenya
- Tuberculosis incidence rate: 319 per 100,000
- Number (proportion) of tuberculosis cases in the study: 3 (1%)

**Index tests**

LF-LAM

**Target condition and reference standard(s)**

- Target condition: pulmonary tuberculosis
- Reference standard: expectorated sputum mycobacterial culture (liquid)

**Flow and timing**

All patients were included in the analysis

**Comparative**

**Notes**

The study authors did not attempt sputum induction and excluded individuals without some respiratory specimen available. There was no attempt to identify extrapulmonary tuberculosis.

**Methodological quality**

| Item                                                       | Authors' judgement | Risk of bias | Applicability concerns |
|------------------------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**                            |                    |              |                        |
| Was a consecutive or random sample of patients enrolled?   | Yes                |              |                        |
| Was a case-control design avoided?                         | Yes                |              |                        |
| Did the study avoid inappropriate exclusions?              | No                 |              |                        |
| **DOMAIN 2: Index Test All tests**                         |                    |              |                        |
### LaCourse 2016 (Continued)

| Question                                                                 | Score |
|--------------------------------------------------------------------------|-------|
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes   |
| If a threshold was used, was it pre-specified?                           | Yes   |
| **DOMAIN 3: Reference Standard**                                         |       |
| Is the reference standards likely to correctly classify the target condition? | No    |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes   |
| **DOMAIN 4: Flow and Timing**                                            |       |
| Was there an appropriate interval between index test and reference standard? | Yes   |
| Did all patients receive the same reference standard?                    | Yes   |
| Were all patients included in the analysis?                              | Yes   |

### Lawn 2017

#### Study characteristics

| Patient sampling | Cross-sectional, prospective and consecutive enrolment |
|------------------|--------------------------------------------------------|
| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected adults included irrespective of symptoms. A positive WHO symptom screen was reported for 91% of participants Age: median 36 (IQR 29-42) Sex, female: 61% HIV infection: 100% Median CD4 cell count per μL: 149 (IQR 55 to 312) History of tuberculosis: 6% Sample size: 413 Clinical setting: inpatients Country: South Africa Tuberculosis incidence rate: 567 per 100,000 Number (proportion) of tuberculosis cases in the study: 139 (32.6%) |

| Index tests | LF-LAM |
|-------------|--------|

| Target condition and reference standard(s) | Target: pulmonary tuberculosis, extrapulmonary tuberculosis, mycobacteraemia Reference standard: any of mycobacterial culture (liquid) or nucleic acid amplification test; extrapulmonary specimens tested for all |

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Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV (Review)  
Copyright © 2019 The Authors. Cochrane Database of Systematic Reviews published by John Wiley & Sons, Ltd. on behalf of The Cochrane Collaboration.
Flow and timing

All patients were included in the analysis

Comparative

Notes

The study authors performed sputum induction for individuals who could not expectorate. They included individuals without respiratory specimen and attempted to identify extrapulmonary tuberculosis

Methodological quality

| Item                                           | Authors’ judgement | Risk of bias | Applicability concerns |
|------------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**                |                    |              |                        |
| Was a consecutive or random sample of patients enrolled? | Yes                | Low          | Low                    |
| Was a case-control design avoided?             | Yes                | Low          | Low                    |
| Did the study avoid inappropriate exclusions?  | Yes                | Low          | Low                    |
| **DOMAIN 2: Index Test All tests**             |                    |              |                        |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes                | Low          | Low                    |
| If a threshold was used, was it pre-specified? | Yes                | Low          | Low                    |
| **DOMAIN 3: Reference Standard**               |                    |              |                        |
| Is the reference standards likely to correctly classify the target condition? | Yes                | Low          | Low                    |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes                | Low          | Low                    |
| **DOMAIN 4: Flow and Timing**                  |                    |              |                        |
| Was there an appropriate interval between index test and reference standard? | Yes                | Low          |                        |
| Did all patients receive the same reference standard? | No                 |              |                        |
| Were all patients included in the analysis?    | Yes                | Low          |                        |
### Nakinyingi 2014

#### Study characteristics

| Patient sampling | Cross-sectional, prospective and consecutive enrolment |
|------------------|--------------------------------------------------------|
| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected adults suspected of tuberculosis; Presented with any of cough, fever, night sweats and weight loss |
| Age | median 33 (IQR 29 to 37) |
| Sex | female: 63% |
| HIV infection | 100% |
| Median CD4 cell count per μL | 152 (IQR 41 to 337) |
| History of tuberculosis | not stated |
| Sample size | 997 |
| Clinical setting | inpatient and outpatient |
| Country | South Africa and Uganda |
| Tuberculosis incidence rate | 567 per 100,000 (South Africa); 201 per 100,000 (Uganda) |
| Number (proportion) of tuberculosis cases in the study | 367 (37%) |

| Index tests | LF-LAM |
|--------------|--------|

| Target condition and reference standard(s) | Target condition: pulmonary tuberculosis, mycobacteraemia |
|---------------------------------------------|----------------------------------------------------------|
| Reference standard | sputum mycobacterial culture (liquid and solid), mycobacterial blood cultures; extrapulmonary specimens tested for all |

| Flow and timing | All patients were included in the analysis |
|-----------------|------------------------------------------|

#### Comparative

#### Notes

The study authors performed sputum induction for individuals who could not expectorate. They excluded individuals without some respiratory specimen available. They attempted to identify extrapulmonary tuberculosis.

#### Methodological quality

| Item | Authors’ judgement | Risk of bias | Applicability concerns |
|------|---------------------|--------------|------------------------|
| DOMAIN 1: Patient Selection |
| Was a consecutive or random sample of patients enrolled? | Yes |
| Was a case-control design avoided? | Yes |
| Did the study avoid inappropriate exclusions? | No |

| High | Low |
|------|-----|

| DOMAIN 2: Index Test All tests |
### Nakiyingi 2014 (Continued)

| Question                                                                 | Yes/No |
|--------------------------------------------------------------------------|--------|
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes    |
| If a threshold was used, was it pre-specified?                           | Yes    |

**DOMAIN 3: Reference Standard**

| Question                                                                 | Yes/No |
|--------------------------------------------------------------------------|--------|
| Is the reference standards likely to correctly classify the target condition? | Yes    |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes    |

**DOMAIN 4: Flow and Timing**

| Question                                                                 | Yes/No |
|--------------------------------------------------------------------------|--------|
| Was there an appropriate interval between index test and reference standard? | Yes    |
| Did all patients receive the same reference standard?                     | Yes    |
| Were all patients included in the analysis?                               | Yes    |

### Pandie 2016

#### Study characteristics

| Characteristic                                                                 | Details |
|------------------------------------------------------------------------------|---------|
| **Patient sampling**                                                         | Cross-sectional, prospective and consecutive enrolment |
| **Patient characteristics and setting**                                     | Presenting signs and symptoms: HIV-infected and HIV-uninfected adults suspected of pericardial tuberculosis. Presented with pericardial effusion. **Age**: median 34 (IQR 29-42) **Sex**, female: 38% **HIV infection**: 74% **Median CD4 cell count per µL**: 139 (IQR 81 to 249) **History of tuberculosis**: not stated **Sample size**: 102 **Clinical setting**: inpatients **Country**: South Africa **Tuberculosis incidence rate**: 567 per 100,000 **Number (proportion) of tuberculosis cases in the study**: 74 (49%) |
| **Index tests**                                                              | LF-LAM  |
| **Target condition and reference standard(s)**                              | Target condition: extrapulmonary tuberculosis (pericardial tuberculosis) and pulmonary tuberculosis |
Pandie 2016 (Continued)

Reference standard: mycobacterial culture (liquid). There was no protocol to ensure a minimum standard of sputum sampling and testing

Flow and timing

Not all patients were included in the analysis

Comparative

Notes

The study authors excluded participants with disseminated tuberculosis if not having pericardial effusion, and excluded patients with pulmonary tuberculosis if no sign of pericardial effusion. Not all patients received the same reference standard. For some patients sputum induction was attempted and sputum examined for mycobacteria. Not all participants were included in the analysis and it is unclear who was excluded. The true negative values reported conflicts and provides uncertainty about the specificity reported.

Methodological quality

| Item                                      | Authors' judgement | Risk of bias | Applicability concerns |
|-------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**           |                    |              |                        |
| Was a consecutive or random sample of pa- | Yes                |              | High                   |
| tients enrolled?                          |                    |              | Low                    |
| Was a case-control design avoided?        | Yes                |              |                        |
| Did the study avoid inappropriate exclu-  | No                 |              |                        |
| sions?                                    |                    |              |                        |
| **DOMAIN 2: Index Test All tests**        |                    |              |                        |
| Were the index test results interpreted  | Yes                |              |                        |
| without knowledge of the results of the  |                    |              |                        |
| reference standard?                       |                    |              | Low                    |
| If a threshold was used, was it pre-spe-  | Yes                |              |                        |
| cified?                                   |                    |              | Low                    |
| **DOMAIN 3: Reference Standard**          |                    |              |                        |
| Is the reference standards likely to cor- | No                 |              |                        |
| rectly classify the target condition?     |                    |              |                        |
| Were the reference standard results in-  | Yes                |              |                        |
| terpreted without knowledge of the results |                    |              | High                   |
| of the index tests?                       |                    |              |                        |
| **DOMAIN 4: Flow and Timing**             |                    |              |                        |
| Was there an appropriate interval between | Yes                |              |                        |
| index test and reference standard?        |                    |              |                        |
### Pandie 2016 (Continued)

| Did all patients receive the same reference standard? | No |
|------------------------------------------------------|----|
| Were all patients included in the analysis?          | No |

**Methodological quality**

| Item                               | Authors’ judgement | Risk of bias | Applicability concerns |
|------------------------------------|--------------------|--------------|------------------------|

**Notes**

The study authors included all patients with symptoms of tuberculosis irrespective of their ability to produce sputum; clinicians chose the reference standard rather than it being directed by the study protocol. The reference standard included testing of pulmonary and extrapulmonary sites by mycobacterial culture.
**Peter 2012a (Continued)**

**DOMAIN 1: Patient Selection**

- Was a consecutive or random sample of patients enrolled? Yes
- Was a case-control design avoided? Yes
- Did the study avoid inappropriate exclusions? Yes

| Low | Low |

**DOMAIN 2: Index Test All tests**

- Were the index test results interpreted without knowledge of the results of the reference standard? Yes
- If a threshold was used, was it pre-specified? Yes

| Low | Low |

**DOMAIN 3: Reference Standard**

- Is the reference standards likely to correctly classify the target condition? No
- Were the reference standard results interpreted without knowledge of the results of the index tests? Yes

| High | High |

**DOMAIN 4: Flow and Timing**

- Was there an appropriate interval between index test and reference standard? Yes
- Did all patients receive the same reference standard? No
- Were all patients included in the analysis? Yes

| High |

**Study characteristics**

| Patient sampling | Cross-sectional, prospective and consecutive enrolment |
|------------------|------------------------------------------------------|
| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected adults suspected of tuberculosis; presenting with any of cough, night sweats, fever or weight loss |
| | Age: median 36 (IQR 30 to 41) |
| | Sex, female: 46% |
HIV infection: 100%
Median CD4 cell count per μL: 210 (IQR 103 to 375)
History of tuberculosis: 26%
Sample size: 569
Clinical setting: outpatient
Country: South Africa, Tanzania, Zambia
Tuberculosis incidence rate: 567 per 100,000 (South Africa); 269 per 100,000 (Tanzania); 361 per 100,000 (Zambia)
Number (proportion) of tuberculosis cases in the study: 181 (32%)

| Index tests        | LF-LAM |
|--------------------|--------|
| Target condition and reference standard(s) | Target condition: pulmonary tuberculosis |
|                     | Reference standard: sputum mycobacterial (liquid) culture |

Flow and timing:
All patients were included in the analysis

Notes:
The study authors excluded individuals unable to expectorate sputum and did not attempt sputum induction. They did not attempt to identify extra-pulmonary tuberculosis.

Methodological quality

| Item                                           | Authors' judgement | Risk of bias | Applicability concerns |
|-----------------------------------------------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection**               |                    |              |                        |
| Was a consecutive or random sample of patients enrolled? | Yes                |              |                        |
| Was a case-control design avoided?            | Yes                |              |                        |
| Did the study avoid inappropriate exclusions? | No                 |              |                        |
|                                               | **High**           | **Low**      |                        |
| **DOMAIN 2: Index Test All tests**            |                    |              |                        |
| Were the index test results interpreted without knowledge of the results of the reference standard? | Yes |              |                        |
| If a threshold was used, was it pre-specified? | Yes |              |                        |
|                                               | **Low**           | **Low**      |                        |
| **DOMAIN 3: Reference Standard**              |                    |              |                        |
### Peter 2015 (Continued)

| Question                                                                 | Response |
|--------------------------------------------------------------------------|----------|
| Is the reference standards likely to correctly classify the target condition? | No       |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes      |
| **DOMAIN 4: Flow and Timing**                                            |          |
| Was there an appropriate interval between index test and reference standard? | Yes      |
| Did all patients receive the same reference standard?                    | Yes      |
| Were all patients included in the analysis?                              | Yes      |

### Low

#### Peter 2016

#### Study characteristics

| Patient sampling | Randomized controlled trial, consecutive prospective enrolment |
|------------------|---------------------------------------------------------------|
| Patient characteristics and setting | Presenting signs and symptoms: HIV-infected adults suspected of tuberculosis; Presenting with any of cough, night sweats, fever, or weight loss; or illness that necessitate admission |
| Age               | median 37 (IQR 30 to 44) |
| Sex               | female: 50% |
| HIV infection     | 100% |
| Median CD4 cell count per μL | 81 (IQR 26 to 198) |
| History of tuberculosis | 27% |
| Sample size       | 1172 |
| Clinical setting  | inpatient |
| Country           | South Africa, Zimbabwe, Zambia |
| Tuberculosis incidence rate | 567 per 100,000 (South Africa); 221 per 100,000 (Zimbabwe); 361 per 100,000 (Zambia) |
| Number (proportion) of tuberculosis cases in the study | 342 (29%) |

#### Index tests

| LF-LAM |

#### Target condition and reference standard(s)

| Target condition: pulmonary tuberculosis, extrapulmonary tuberculosis, mycobacteraemia |
| Reference standard: any of mycobacterial culture (solid and liquid) or nucleic acid amplification test; extrapulmonary specimens collected for those participants where treating clinicians found it indicated |
Flow and timing

All patients were included in the analysis

Comparative

Notes

Health providers selected the sites for testing based on their own clinical suspicion. High concern of applicability because of a lack of study directed extrapulmonary testing. Not all participants received the same reference standard as tuberculosis diagnostics performed at the discretion of the clinician and varied across sites.

Methodological quality

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|------|--------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection** | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | Low | Low |
| Was a case-control design avoided? | Yes | Low | Low |
| Did the study avoid inappropriate exclusions? | Yes | Low | Low |

**DOMAIN 2: Index Test All tests**

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|------|--------------------|--------------|------------------------|
| Were the index test results interpreted without knowledge of the results of the reference standard? | No | Low | Low |
| If a threshold was used, was it pre-specified? | Yes | Low | Low |

**DOMAIN 3: Reference Standard**

| Item | Authors' judgement | Risk of bias | Applicability concerns |
|------|--------------------|--------------|------------------------|
| Is the reference standards likely to correctly classify the target condition? | No | High | High |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Yes | High | High |

**DOMAIN 4: Flow and Timing**

| Item | Authors' judgement | Risk of bias |
|------|--------------------|--------------|
| Was there an appropriate interval between index test and reference standard? | Yes | |
| Did all patients receive the same reference standard? | No | |
| Were all patients included in the analysis? | Yes | |

High
**Study characteristics**

**Patient sampling**
Cohort, prospective and consecutive enrolment with six months follow-up

**Patient characteristics and setting**
Presenting signs and symptoms: irrespective of symptoms. WHO tuberculosis symptom screen positive in 33%. in overall population who could produce sputum signs and symptoms suggestive of tuberculosis, but not specified
- Age: median 34 (IQR 30-41)
- Sex, female: 50%
- HIV infection: 100%
- Median CD4 cell count per μL: 270 (IQR 128 to 443)
- History of tuberculosis: not stated
- Sample size: 517
- Clinical setting: outpatient and inpatients
- Country: Myanmar
- Tuberculosis incidence rate: 358 per 100.000
- Number (proportion) of tuberculosis cases in the study: 54 (12%)

**Index tests**
- LF-LAM

**Target condition and reference standard(s)**
- Target condition: pulmonary tuberculosis
- Reference standard: any of sputum mycobacterial culture (solid and liquid) or nucleic acid amplification test; no extrapulmonary specimens tested. The study did not report if they speciated mycobacteria isolates

**Flow and timing**
The study had specimens collected for index and reference standard tests greater than seven days apart.

**Comparative**

**Notes**
The study authors attempted sputum induction and excluded individuals without some respiratory specimen available. They did not attempt to identify extrapulmonary tuberculosis. The study did not report if they speciated mycobacteria isolates.

**Methodological quality**

| Item | Authors’ judgement | Risk of bias | Applicability concerns |
|------|---------------------|--------------|------------------------|
| **DOMAIN 1: Patient Selection** | | | |
| Was a consecutive or random sample of patients enrolled? | Yes | Low | Low |
| Was a case-control design avoided? | Yes | Low | Low |
| Did the study avoid inappropriate exclusions? | Yes | Low | Low |

**DOMAIN 2: Index Test All tests**
Thit 2017 (Continued)

| Question                                                                 | Yes | Low | Unclear |
|-------------------------------------------------------------------------|-----|-----|---------|
| Were the index test results interpreted without knowledge of the results of the reference standard? |     |     |         |
| If a threshold was used, was it pre-specified?                         | Yes |     |         |

**DOMAIN 3: Reference Standard**

| Question                                                                 | Yes | Low | Unclear |
|-------------------------------------------------------------------------|-----|-----|---------|
| Is the reference standards likely to correctly classify the target condition? | No  |     |         |
| Were the reference standard results interpreted without knowledge of the results of the index tests? | Unclear |     |         |

**DOMAIN 4: Flow and Timing**

| Question                                                                 | Yes | High | Unclear |
|-------------------------------------------------------------------------|-----|------|---------|
| Was there an appropriate interval between index test and reference standard? | No  |      |         |
| Did all patients receive the same reference standard?                    | Yes | High |         |
| Were all patients included in the analysis?                              | Yes |      |         |

**Abbreviations**

LF-LAM: lateral flow urine lipoarabinomannan assay (Alere Determine™ TB lipoarabinomannan);
IQR: interquartile range;
SD: standard deviation;
WHO: World Health Organization

**Characteristics of excluded studies [ordered by study ID]**

| Study       | Reason for exclusion                                           |
|-------------|----------------------------------------------------------------|
| Agha 2013   | Index test not studied                                        |
| Amoudy 1997 | Index test not studied                                        |
| Andrews 2014| Abstract                                                       |
| Balcha 2014 | Did not use manufacturer's current recommended positivity threshold |
| Bisson 2016 | Abstract                                                       |
| Blanc 2018  | Abstract                                                       |
| Boehme 2005 | Index test not studied                                        |
| Boyles 2012 | Index test not studied                                        |
| Broughton 2017 | Study protocol                                               |
| Study                | Reason for exclusion                                        |
|---------------------|------------------------------------------------------------|
| Calligaro 2017      | Insufficient data                                          |
| Chan 2000           | Index test not studied                                     |
| Cho 1997            | Index test not studied                                     |
| Conesa-Botella 2007 | Index test not studied                                     |
| Cox 2015            | Did not use manufacturer’s current recommended positivity threshold |
| d’Elia 2015         | Reference standard not satisfied                           |
| Daley 2009          | Index test not studied                                     |
| Deng 2011           | Index test not studied                                     |
| Dheda 2009          | Index test not studied                                     |
| Dheda 2010          | Index test not studied                                     |
| Drain 2014a         | Did not use manufacturer’s current recommended positivity threshold |
| Drain 2014b         | Abstract                                                   |
| Drain 2015b         | Impact study (no diagnostic accuracy data)                 |
| Drain 2016a         | Did not use manufacturer’s current recommended positivity threshold |
| Drain 2017          | Impact study (no diagnostic accuracy data)                 |
| Elsawy 2012         | Index test not studied                                     |
| Gina 2017           | Reference standard not satisfied                           |
| Gounder 2011        | Index test not studied                                     |
| Grant 2016          | Abstract                                                   |
| Gupta-Wright 2016a  | Not a diagnostic accuracy study                            |
| Gupta-Wright 2016b  | Study protocol, study completed                            |
| Gupta-Wright 2018b  | Abstract                                                   |
| Hamasur 2001        | Index test not studied                                     |
| Hanifa 2015         | Index test not studied                                     |
| Iskandar 2017       | Paediatric population                                      |
| Kashino 2008        | Index test not studied                                     |
| Kerkhoff 2014a      | Insufficient or duplicate data                             |
| Kerkhoff 2014b      | Insufficient or duplicate data                             |
| Study              | Reason for exclusion                                      |
|-------------------|----------------------------------------------------------|
| Kerkhoff 2017     | Second reference (complementary data) to Lawn 2017       |
| Koralnik 2017     | Abstract                                                 |
| Kouchin 2007      | Index test not studied                                   |
| Kroidl 2014       | Index test not studied                                   |
| Kroidl 2015       | Paediatric population                                    |
| LaCourse 2018a    | Paediatric population                                    |
| LaCourse 2018b    | Paediatric population                                    |
| Lawn 2009a        | Index test not studied                                   |
| Lawn 2012a        | Did not use manufacturer’s current recommended positivity threshold |
| Lawn 2012b        | Insufficient or duplicate data                           |
| Lawn 2013a        | Insufficient or duplicate data                           |
| Lawn 2014         | Abstract                                                 |
| Leopold 2017      | Abstract                                                 |
| Love 2016         | Abstract                                                 |
| Manabe 2014       | Insufficient or duplicate data                           |
| Mathabire 2017    | Abstract                                                 |
| Mukundan 2012     | Index test not studied                                   |
| Musarurwa 2018    | Not a diagnostic accuracy study                          |
| Mutetwa 2009      | Index test not studied                                   |
| Nakiyingi 2015    | Second reference (complementary data) to Nakiyingi 2014  |
| Nakiyingi 2015a   | Second reference (complementary data) to Nakiyingi 2014  |
| Nicol 2014        | Paediatric population                                    |
| Patel 2009        | Index test not studied                                   |
| Patel 2010        | Index test not studied                                   |
| Peter 2011        | Insufficient or duplicate data                           |
| Peter 2012b       | Insufficient or duplicate data                           |
| Peter 2012c       | Insufficient or duplicate data                           |
| Peter 2013        | Insufficient or duplicate data                           |
| Study               | Reason for exclusion                                           |
|--------------------|----------------------------------------------------------------|
| Reddy 2018         | Abstract                                                       |
| Reid 2015          | Insufficient data                                              |
| Reither 2009       | Index test not studied                                          |
| Sabur 2017         | Reference standard not satisfied                                |
| Sada 1992          | Index test not studied                                          |
| Sahle 2017         | Did not use manufacturer’s current recommended positivity threshold |
| Savolainen 2013    | Index test not studied                                          |
| Schmidt 2011       | Index test not studied                                          |
| Shah 2009          | Index test not studied                                          |
| Shah 2010          | Index test not studied                                          |
| Shah 2013          | Insufficient or duplicate data                                  |
| Shah 2014          | Insufficient or duplicate data                                  |
| Singh 2011         | Index test not studied                                          |
| Sun 2013           | Insufficient data                                              |
| Suwanpimolkul 2017 | Did not use manufacturer’s current recommended positivity threshold |
| Tessema 2001       | Index test not studied                                          |
| Tessema 2002a      | Index test not studied                                          |
| Tessema 2002b      | Index test not studied                                          |
| Tlali 2014         | Insufficient or duplicate data                                  |
| Van Rie 2013       | Insufficient or duplicate data                                  |
| Wood 2012          | Index test not studied                                          |
| Zijenah 2016       | Second reference (complementary data) to Peter 2016             |

**DATA**

Presented below are all the data for all of the tests entered into the review.
| Test                                                   | No. of studies | No. of participants |
|-------------------------------------------------------|----------------|---------------------|
| 1 Symptomatic adults, all settings                    | 8              | 3449                |
| 2 Symptomatic adults, inpatients                     | 6              | 2253                |
| 3 Symptomatic adults, outpatients                     | 4              | 1196                |
| 4 Symptomatic adults, CD4 > 200, all settings         | 3              | 738                 |
| 5 Symptomatic adults, CD4 ≤ 200, all settings         | 4              | 1825                |
| 6 Symptomatic adults, CD4 ≤ 200, inpatients           | 2              | 1009                |
| 7 Symptomatic adults, CD4 ≤ 200, outpatients          | 1              | 249                 |
| 8 Symptomatic adults, CD4 > 100, all settings         | 4              | 1519                |
| 9 Symptomatic adults, CD4 ≤ 100, all settings         | 5              | 1251                |
| 10 Symptomatic adults, CD4 ≤ 100, inpatients          | 3              | 746                 |
| 11 Symptomatic adults, CD4 ≤ 100, outpatients         | 1              | 121                 |
| 12 Symptomatic adults, CD4 101-200, all settings      | 4              | 586                 |
| 13 Symptomatic adults, CD4 101-200, inpatients        | 2              | 275                 |
| 14 Symptomatic adults, CD4 101-200, outpatients       | 1              | 128                 |
| 17 Unselected adults, all settings                    | 7              | 3365                |
| 18 Unselected adults, inpatients                      | 3              | 537                 |
| 19 Unselected adults, outpatients                     | 6              | 2828                |
| 20 Unselected adults, CD4 > 200, all settings         | 1              | 156                 |
| 21 Unselected adults, CD4 ≤ 200, all settings         | 2              | 706                 |
| 22 Unselected adults, CD4 ≤ 200, inpatients           | 1              | 54                  |
| 23 Unselected adults, CD4 ≤ 200, outpatients          | 2              | 652                 |
| 24 Unselected adults, CD4 > 100, all settings         | 4              | 952                 |
| 25 Unselected adults, CD4 ≤ 100, all settings         | 3              | 417                 |
| 26 Unselected adults, CD4 ≤ 100, inpatients           | 2              | 200                 |
| 27 Unselected adults, CD4 ≤ 100, outpatients          | 2              | 217                 |
| 28 Unselected adults, CD4 101-200, all settings       | 1              | 103                 |
| 29 Unselected adults, CD4 101-200, inpatients         | 1              | 9                   |
| Test                                                                 | No. of studies | No. of participants |
|----------------------------------------------------------------------|----------------|---------------------|
| 30 Unselected adults, CD4 101-200, outpatients                       | 1              | 94                  |

Test 1. Symptomatic adults, all settings.

Test 2. Symptomatic adults, inpatients.

Test 3. Symptomatic adults, outpatients.

Test 4. Symptomatic adults, CD4 > 200, all settings.

Test 5. Symptomatic adults, CD4 ≤ 200, all settings.

Test 6. Symptomatic adults, CD4 ≤ 200, inpatients.

Test 7. Symptomatic adults, CD4 ≤ 200, outpatients.

Test 8. Symptomatic adults, CD4 > 100, all settings.

Test 9. Symptomatic adults, CD4 ≤ 100, all settings.

Test 10. Symptomatic adults, CD4 ≤ 100, inpatients.

Test 11. Symptomatic adults, CD4 ≤ 100, outpatients.

Test 12. Symptomatic adults, CD4 101-200, all settings.
Test 13. Symptomatic adults, CD4 101-200, inpatients.

Test 14. Symptomatic adults, CD4 101-200, outpatients.

Test 17. Unselected adults, all settings.

Test 18. Unselected adults, inpatients.

Test 19. Unselected adults, outpatients.

Test 20. Unselected adults, CD4 > 200, all settings.

Test 21. Unselected adults, CD4 ≤ 200, all settings.

Test 22. Unselected adults, CD4 ≤ 200, inpatients.

Test 23. Unselected adults, CD4 ≤ 200, outpatients.

Test 24. Unselected adults, CD4 > 100, all settings.

Test 25. Unselected adults, CD4 ≤ 100, all settings.

Test 26. Unselected adults, CD4 ≤ 100, inpatients.

Test 27. Unselected adults, CD4 ≤ 100, outpatients.
Test 28. Unselected adults, CD4 101-200, all settings.

Test 29. Unselected adults, CD4 101-200, inpatients.

Test 30. Unselected adults, CD4 101-200, outpatients.
### ADDITIONAL TABLES

**Table 1. Summary characteristics of included studies**

| Study     | Characteristics                                                                 | Participants (% symptom) | Setting          | Median CD4 cell count per μL (IQR) | TB prevalence % (n/N) | Did the study avoid inappropriate exclusion? | Specimens collected | High-quality reference standarda | Unique study characteristics |
|-----------|---------------------------------------------------------------------------------|--------------------------|-----------------|-----------------------------------|-----------------------|-----------------------------------------------|----------------------|-------------------------------|--------------------------------|
| HIV-positive adults with signs and symptoms of TB                                                                                             |                          |                 |                                   |                       |                                               |                      |                               |                                |
| Drain 2016                                                                                                                                  | Symptomatic: Two of four TB-related symptoms (cough, fever weight loss, night sweat) for > 2 weeks; smear microscopy negative x 2 | Outpatient           | 168 (89 to 256)               | 63% (57/90)             | No                                       | Pulmonary samples    | No                             | Adults (> 18 years); HIV positive (93.2%); targeting a relatively well outpatient population; Karnofsky performance score > 50 |
| Huerga 2017                                                                                                                                  | Symptomatic: Cough > 2 weeks or any cough and one of weight loss, night sweats or fever; severely ill; CD4< 200 or BMI below 17 | Outpatients (33%); Inpatients (67%) | 109 (43 to 214)               | 57% (156/275)          | No                                       | Pulmonary samples; urine Xpert only for patients without sputum available | No                             | Adults (> 15 years); LAM-guided treatment; excluded many participants from analysisb |
| Juma 2017                                                                                                                                     | Symptomatic: Suggestive of extrapulmonary TB, not specified | Inpatients           | Not stated                    | 33% (29/67)            | No                                       | Extrapulmonary samples only, no sputum samples | No                             | Adults (> 14 years), HIV-positive (68%); excluded patients with concomitants active pulmonary TB |
| Nakiyingi 2014                                                                                                                                | Symptomatic: Any of cough, fever weight loss, night sweat | Outpatients (45%); Inpatients (55%) | 152 (41 to 337)               | 37% (367/997)          | No                                       | Pulmonary samples; Blood culture for all | Yes                            | Adults (> 18 years); multisite; large sample size. |
| Pandie 2016                                                                                                                                   | Symptomatic: Presence of a pericardial effusion and suspected of pericardial TB | Inpatients           | 139 (81 to 249)               | 95% (36/38)            | No                                       | Extrapulmonary samples (pericardial effusion); pulmonary samples for some | No                             | Adults (> 18 years); HIV-positive (74%); excluded participants from analysis affecting specificityc |
### Table 1. Summary characteristics of included studies (Continued)

| Study         | Population | Symptomatic: | Sample Size | Sensitivity | Algorithm Defined | Pulmonary Samples | Extrapulmonary Samples | Study Design |
|---------------|------------|--------------|-------------|-------------|-------------------|-------------------|--------------------------|--------------|
| Peter 2012a  | Symptomatic: Inpatients | 90 (47 to 197) | 48% (116/241) | Yes | Clinically relevant pulmonary samples; clinically relevant extrapulmonary samples. No study defined algorithm. | No | Adults (> 18 years); multisite; TB diagnostic work-up was not standardized but up to clinical judgements |
| Peter 2015   | Symptomatic: Outpatients | 210 (103 to 375) | 32% (181/569) | No | Pulmonary samples | No | Adults (> 18 years); multisite; nested within a randomized, parallel-arm trial. |
| Peter 2016   | Symptomatic: Inpatients | 81 (26 to 198) | 29% (342/1172) | Yes | Pulmonary samples; clinically relevant extrapulmonary samples. No study defined algorithm. | No | Adults (> 18 years); multisite; LAM arm of a randomized controlled trial. |

### HIV-positive adults irrespective of signs and symptoms of TB

| Study         | Population | Symptomatic: | Sample Size | Sensitivity | Algorithm Defined | Pulmonary Samples | Extrapulmonary Samples | Study Design |
|---------------|------------|--------------|-------------|-------------|-------------------|-------------------|--------------------------|--------------|
| Bjerrum 2015  | Unselected: Outpatients | 127 (35 to 256) | 12% (55/469) | No | Pulmonary samples | No | Adults (> 18 years); majority symptomatic. |
| Drain 2015a  | Unselected: Outpatient | 248 (107 to 379) | 17% (54/320) | No | Pulmonary samples | No | Adults (> 18 years) |
| Floridia 2017 | Unselected: Outpatient | 278 (142 to 395) | 9% (90/972) | No | Pulmonary samples | No | Adults (> 15 years); LAM-guided treatment. |
| Hanifa 2016  | Unselected: Outpatient | 111 (56 to 161) | 9% (40/408) | Yes | Pulmonary samples; blood culture for all | Yes | Adults (> 18 years); CD4 < 200; reference standard included any sample taken within six months from enrolment. |
| LaCourse 2016 | Unselected: Outpatient | 437 (342 to 565) | 1% (3/266) | No | Pulmonary samples | No | Pregnant women (> 16 years) attending ANC; healthy popula- |
**Table 1. Summary characteristics of included studies (Continued)**

| Study | Unselected: | Analysis Type | Participants | Particpants with TB (%) | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) | Reference Standard | Adults (> 18 years). Included many samples from different sites |
|-------|-------------|---------------|--------------|-------------------------|-----------------------------|------------------------------|----------------|-------------------------------------------------|
| Lawn 2017 | Unselected: | Inpatients | 149 (55-312) | 33% (139/413) | Yes | Pulmonary samples; Blood culture for all; Clinically relevant extrapulmonary samples | Yes | Adults (> 18 years). Included many samples from different sites |
| Thit 2017 | Unselected: | Outpatients (90%); Inpatients (10%) | 270 (128 to 443) | 10% (54/517) | Yes | Pulmonary samples | No | Conducted in Myanmar. Adults (median age 34). Reference standard included samples taken within six months from enrolment. |

**Abbreviations:** LF-LAM: lateral flow urine lipoarabinomannan assay (Alere Determine™ TB lipoarabinomannan assay); ANC: antenatal clinic; BMI: body mass index; IQR: interquartile range; TB: tuberculosis; Xpert: Xpert MTB/RIF

a For a microbiological reference standard, we considered a higher quality reference standard to be one in which two or more specimen types were evaluated for TB diagnosis in all participants as part of a defined standardized study algorithm.

b Huerga 2017 excluded participants from analysis if missing Xpert results or culture contaminated for any of the samples in the absence of a positive result; overall samples size 474 (156 with TB); 275 included in analysis (156 with TB).

c Pandie 2016 excluded a large number of non-TB participants from analysis; Overall samples size 102 (36 with TB); 38 included for analysis (36 TB cases).

**Table 2. LF-LAM pooled sensitivity and specificity for TB diagnosis, by study population**

| Type of analysis | Symptomatic participants | Unselected participants |
|------------------|--------------------------|-------------------------|
|                  | Studies (total participants) | Participants with TB (%) | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) | Studies (total participants) | Participants with TB (%) | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) |
| Overall accuracy | 8 studies (3449) | 1277 (37%) | 42% (31% to 55%) | 91% (85% to 95%) | 7 studies (3365) | 432 (13%) | 35% (22% to 50%) | 95% (89% to 98%) |
| By setting | | | | | | | | |
| Inpatient | 6 studies (2253) | 868 (39%) | 52% (40% to 64%) | 87% (78% to 93%) | 3 studies (537) | 159 (30%) | 62% (41% to 83%) | 84% (48% to 96%) |
| Outpatient | 4 studies | 409 | 29% | 96% | 6 studies | 273 | 31% | 95% |
### Table 2. LF-LAM pooled sensitivity and specificity for TB diagnosis, by study population (Continued)

| By CD4 cell | (1196) | (34%) | (17% to 47%) | (91% to 99%) | (2828) | (10%) | (18% to 47%) | (87% to 99%) |
|---|---|---|---|---|---|---|---|---|
| CD4 > 200 | 3 studies | 163 | 16% (8% to 31%) | 94% (81% to 97%) | 1 study | 11 | Not applicable | Not applicable |
| | (738) | (22%) | | | (156) | (7%) | | |
| CD4 ≤ 200 | 4 studies | 722 | 45% (31% to 61%) | 89% (77% to 94%) | 2 studies | 82 | 26% (9% to 56%) | 96% (87% to 98%) |
| | (1825) | (40%) | | | (706) | (12%) | | |
| CD4 > 100 | 4 studies | 425 | 17% (10% to 27%) | 95% (89% to 98%) | 4 studies | 115 | 20% (10% to 35%) | 98% (95% to 99%) |
| | (1519) | (28%) | | | (952) | (12%) | | |
| CD4 ≤ 100 | 4 studies | 512 | 54% (38% to 69%) | 88% (77% to 94%) | 3 studies | 130 | 47% (40% to 64%) | 90% (77% to 96%) |
| | (1239) | (41%) | | | (417) | (31%) | | |
| CD4 101-199 | 4 studies | 210 | 24% (14% to 38%) | 90% (77 to 96) | 1 study | 13 | Not applicable | Not applicable |
| | (586) | (36%) | | | (103) | (13%) | | |
| By CD4 and setting | (Continued) | | | | | | | |
| CD4 ≤ 200 | 2 studies | 348 | 54% | 80% | 1 study | 14 | Not applicable | Not applicable |
| inpatients | (1009) | (34%) | (34% to 73%) | (58% to 91%) | (54) | (26%) | | |
| CD4 ≤ 100 | 2 studies | 270 | 61% | 81% | 2 studies | 84 | 57% | 90% |
| inpatients | (734) | (37%) | (40% to 78%) | (61% to 91%) | (200) | (42%) | (33% to 79%) | (69% to 97%) |
| CD4 101-199 | 2 studies | 78 | 32% | 81% | 1 study | 4 | Not applicable | Not applicable |
| inpatients | (275) | (28%) | (16% to 57%) | (55% to 92%) | (9) | (44%) | | |
| CD4 ≤ 200 | 1 study | 97 | Not applicable | Not applicable | 2 studies | 68 | 21% | 96% |
| outpatients | (249) | (39%) | | | (652) | (10%) | (8% to 48%) | (89% to 99%) |
| CD4 ≤ 100 | 1 study | 48 | Not applicable | Not applicable | 2 studies | 46 | 40% | 87% |
| outpatients | (121) | (40%) | | | (217) | (21%) | (20 to 64) | (68 to 94) |
Table 2. LF-LAM pooled sensitivity and specificity for TB diagnosis, by study population  (Continued)

| CD4 101-199 | 1 study\(^h\) | 51 | Not applicable | Not applicable | 1 study\(^e\) | 9 | Not applicable | Not applicable |
|-------------|----------------|----|----------------|----------------|-------------|---|----------------|----------------|
| outpatients | 128            | (40%) | Not applicable | Not applicable | (94)        | (10%) | Not applicable | Not applicable |

**Abbreviations:** LF-LAM: lateral flow urine lipoarabinomannan assay (Alere Determine™ TB lipoarabinomannan assay); Crl: credible interval; TB: tuberculosis.

\(^a\)Bjerrum 2015, Sensitivity 27% (6% to 61%); Specificity 99% (96% to 100%); \(^b\)Bjerrum 2015, Sensitivity 38% (14% to 68%); Specificity 99% (94% to 100%); \(^c\)Bjerrum 2015, Sensitivity 64% (35% to 87%); Specificity 82% (67% to 93%); \(^d\)Bjerrum 2015, Sensitivity 75% (19% to 99%); Specificity 100% (48% to 100%); \(^e\)Bjerrum 2015, Sensitivity 22% (3% to 60%); Specificity 99% (94% to 100%); \(^f\)Peter 2015, Sensitivity 24% (16% to 33%); Specificity 94% (89% to 97%); \(^g\)Peter 2015, Sensitivity 30% (18% to 46%); Specificity 93% (85% to 98%); \(^h\)Peter 2015, Sensitivity 18% (8% to 31%); Specificity 95% (87% to 99%).
### Table 3. Sensitivity analyses, LF-LAM

| Type of analysis                                                                 | Studies participants |    |    | Studies participants |    |    |
|---------------------------------------------------------------------------------|----------------------|----|----|----------------------|----|----|
|                                                                            | Symptomatic participants | Unselected participants |
|                                                                            | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) |
| Including studies at low risk of patient selection bias                        | 2 studies 1413        | 48% (29% to 67%)             | 82% (61% to 92%)             | 3 studies 1338                 | 39% (17% to 66%)             | 93% (69% to 98%)             |
| Including studies at low risk of reference standard bias                       | 1 study 997           | Insufficient data            | Insufficient data            | 2 studies 821                   | 24% (8% to 53%)              | 99% (95% to 99%)             |
| Including studies that used only fresh specimens                               | 4 studies 2511        | 52% (38% to 68%)             | 91% (82% to 95%)             | 5 studies 2544                   | 41% (26% to 56%)             | 93% (82% to 97%)             |
| Re-classification of Bjerrum 2015; Lawn 2017 as studies among symptomatic participant (> 80% participants symptomatic) | 10 studies 4331       | 42% (33% to 52%)             | 93% (88% to 96%)             | 5 studies 2483                   | 31% (16% to 50%)             | 94% (84% to 98%)             |

**Abbreviations:** LF-LAM: lateral flow urine lipoarabinomannan assay (Alere Determine™ TB lipoarabinomannan assay); CrI: Credible Interval

### Table 4. Original Alere LAM review, main findings

| Type of analysis                                                                 | Symptomatic participants |    |    | Unselected participants |
|---------------------------------------------------------------------------------|--------------------------|----|----|--------------------------|
|                                                                            | Studies (total participants) | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) | Studies (total participants) | Pooled sensitivity (95% CrI) | Pooled specificity (95% CrI) |
| Overall accuracy                                                                | 5 studies (2313)         | 45% (29% to 63%)             | 92% (80% to 97%)             | 3 studies a (1055)           | 30% (20% to 43%)             | 94% (86% to 97%)             |
| By setting                                                                      |                          |                            |                              |                           |                             |                              |
| Inpatient                                                                       | 4 studies (1299)         | 53% (38% to 70%)             | 90% (73% to 96%)             | Insufficient data           |                             |                              |
| Outpatient                                                                       | Insufficient data        |                             |                               | Insufficient data           |                             |                              |

**Abbreviations:** LF-LAM: lateral flow urine lipoarabinomannan assay (Alere Determine™ TB lipoarabinomannan assay); CrI: credible interval

Main findings from the original review on LF-LAM accuracy (Shah 2016).

aReported at grade 1 on the previous reference scale card with five bands that is outdated and no longer the recommended positivity threshold.
APPENDICES

Appendix 1. Alere Determine TB LAM Ag test

Figure 10. (A) Alere™ TB LAM Ag test. To the sample pad (white pad marked by the arrow symbols) 60 µL of urine is applied and visualized bands are read 25 minutes later. (B) Updated Reference Scale Card accompanying test strips to 'grade' the test result and Alere positivity. Copyright © [2019] [Abbott Inc]: reproduced with permission.

Appendix 2. Reference card grading of Alere Determine TB LAM Ag test

Figure 11.
Appendix 3. Minimum specifications for a point-of-care TB diagnostic test

| Test specification                        | Minimum required value                                                                 |
|-------------------------------------------|----------------------------------------------------------------------------------------|
| Medical decision                          | Treatment initiation                                                                   |
| Sensitivity: adults (for pulmonary tuberculosis only; regardless of HIV status) | Pulmonary tuberculosis
  - 95% for smear positive, culture positive
  - 60% to 80% for smear negative, culture positive
  (detection of extrapulmonary tuberculosis being a preferred but not minimal requirement) |
| Sensitivity: children (including extrapulmonary tuberculosis; regardless of HIV status) | 80% compared to culture of any specimen and
  - 60% of probable tuberculosis (noting problem of lack of a gold standard) |
| Specificity: adults                       | 95% compared to culture                                                               |
| Specificity: children                     | 90% for culture-negative probable tuberculosis (noting problem of lack of a gold standard)
  - 95% compared to culture |
| Time to results                           | 3 hours maximum (patient must receive results the same day)                           |
### (Continued)

(desirable would be < 15 minutes)

| Throughput               | 20 tests/day minimum, by one laboratory technician |
|--------------------------|---------------------------------------------------|
| Specimen type            | Adults: urine, oral, breath, venous blood, sputum |
|                          | (desired: non sputum-based sample type and use of finger prick instead of venous blood) |
|                          | Children: urine, oral, capillary blood (finger or heel prick) |
| Sample preparation       | • 3 steps maximum                                  |
|                          | • Safe: biosafety level 1                           |
|                          | • Ability to use approximate volumes (that is, no need for precise pipetting) |
|                          | • Preparation that is not highly time sensitive    |
| Number of samples        | 1 sample per test                                   |
| Readout                  | • Easy-to-read, unambiguous, simple "yes", "no", or "invalid" answer |
|                          | • Readable for at least one hour                    |
| Waste disposal           | • Simple burning or sharps disposal; no glass component |
|                          | • Environmentally acceptable disposal               |
| Controls                 | • Positive control included in test kit             |
|                          | • Quality control simpler and easier than with sputum smear microscopy |
| Reagents                 | • All reagents in self-contained kit                |
|                          | • Kit contains sample collection device and water (if needed) |
| Storage/stability required| • Shelf life of 24 months, including reagents       |
|                          | • Stable at 30°C, and at higher temperatures for shorter time periods |
|                          | • Stable in high humidity environments              |
| Instrumentation          | • If instrument needed, no maintenance required     |
|                          | • Instrument works in tropical conditions           |
|                          | • Acceptable replacement cost                       |
|                          | • Fits in backpack                                  |
|                          | • Shock resistant                                   |
| Power requirement        | Can work on battery                                 |
| Training                 | • 1 day maximum training time                       |
|                          | • Can be performed by any health worker             |
| Cost                     | < USD 10 per test after scale-up                    |

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Abbreviations: USD, United States Dollar.

**Appendix 4. Detailed search strategies**

**MEDLINE (PubMed)**
Search

#9 Search (#3) AND (#7) AND #8
#8 Search test OR assay OR antigen OR Ag OR lateral flow assay* OR urine antigen OR point of care Field: Title/Abstract
#7 Search (#4) OR #5) OR #6
#6 Search LAM; Field: Title/Abstract
#5 Search "lipoarabinomannan" [Supplementary Concept]
#4 Search lipoarabinomannan ; Field: Title/Abstract
#3 Search (#1) OR #2
#2 Search tuberculosis Or TB Field: Title/Abstract
#1 Search ("Tuberculosis"[Mesh]) OR "Mycobacterium tuberculosis"[Mesh]

Embase 1947-Present, updated daily

Search strategy:

1 tuberculosis.mp. or tuberculosis/ or Mycobacterium tuberculosis/ (115438)
2 limit 1 to yr="2014 -Current" (8833)
3 lipoarabinomannan.mp. or lipoarabinomannan/ (775)
4 LAM.mp. (4928)
5 limit 4 to yr="2014 -Current" (500)
6 3 or 5 (1252)
7 2 and 6 (79)
8 (test or assay or antigen or Ag or lateral flow assay* or urine antigen or point of care).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword] (2733052)
9 limit 8 to yr="2014 -Current" (223771)
10 7 and 9 (46)

Cochrane Central Register of Controlled Trials : Issue 4 of 12, April 2018

ID Search
#1 tuberculosis:ti,ab,kw (Word variations have been searched)
#2 TB:ti, ab, kw
#3 MeSH descriptor: [Mycobacterium tuberculosis] explode all trees
#4 MeSH descriptor: [Tuberculosis] explode all trees
#5 #1 or #2 or #3 or #4
#6 LAM:ti,ab,kw
#7 lipoarabinomannan:ti,ab,kw
#8 #6 or #7
#9 #5 and #8

**Web of Science Core Collection - Indexes: SCI-EXPANDED, CPCI-S, Biosis previews**

**TOPIC**: (tuberculosis OR TB OR mycobacterium) AND **TOPIC**: (lipoarabinomannan OR LAM) AND **TOPIC**: (test OR assay OR antigen OR Ag OR lateral flow assay* OR urine antigen OR point of care)

**SCOPUS**

( TITLE-ABS-KEY (tuberculosis OR TB) AND TITLE-ABS-KEY (lipoarabinomannan OR LAM) AND (test OR diagnos* OR urine OR assay))

**CIDG Specialized Register, LILACS, Proquest dissertations, Current Controlled trials, WHO trials register**

Tuberculosis AND (lipoarabinomannan OR LAM)

Appendix 5. Data collection form

---

**LF-LAM** - Lateral flow urine lipoarabinomannan assay for diagnosing active tuberculosis in people living with HIV

**Data form**

### I. STUDY IDENTIFICATION

|   |   |
|---|---|
| 1 | First author |
| 2 | Corresponding author and email |
| 3 | Title of study |
| 4 | Year of publication |
| 5 | Year of study start |
| 6 | Language if other than English |

### II. STUDY DETAILS

|   |   |
|---|---|
| 7 | Population |
| 8 | In which country or countries was the study conducted? |
| 9 | Country World Bank Classification (income) |

1. Adults (15 years of age)
2. Children and adolescents
3. Both adults, children and adolescents
4. Other
   If other, describe

1. Low income
2. Lower-middle income
3. Upper-middle-income
4. High income
5. Other combination
6. Unknown/Not reported
   If other, describe:
|   | Country WHO classification for high TB burden country (WHO Global Tuberculosis Report 2018) | 1. Yes, part of the High TB/HIV burden list  
2. No, not part of the High TB/HIV burden list |
|---|---|---|
|   | Study design | 1. Randomized controlled trial  
2. Cross-sectional  
3. Cohort  
7. Other, specify  
9. Could not tell  
If other, describe: |
|   | Was a case-control design avoided? | 1. Yes  
2. No  
9. Unclear |

### III. PATIENT SELECTION

|   | What was the manner of participant selection into the study? | 1. Consecutive  
2. Random  
3. Convenience  
7. Other, specify  
9. Unknown/Not Reported/Unclear  
If other, describe: |
|---|---|---|
|   | Direction of study data collection | 1. Prospective  
2. Retrospective  
9. Unknown/Not reported |
|   | Please select the statement that best describes the selection of participants into the study. | 1. HIV-positive participants with signs or symptoms suggestive of active tuberculosis were tested using LF-LAM. Please provide study definition of ‘signs and symptoms’:  
2. A preferred target population of HIV-positive individuals, irrespective of signs and symptoms of tuberculosis, were tested using LF-LAM. Please specify target population:  
3. Both 1 and 2  
4. Neither 1 nor 2. This is what was done: |
|   | Sample size | 1. ______  
9. Unknown/Not reported |
|   | Did the study avoid inappropriate exclusions? | 1. Yes  
2. No  
9. Unknown/Not reported/Unclear |
|   | Could the selection of patients have introduced bias? | 1. High risk  
2. Low risk  
9. Unclear risk |

### IV. PATIENT CHARACTERISTICS AND SETTING

|   | Presenting signs and symptoms | List |
|---|---|---|
|   | Age (years) | If age is reported in median indicate IQR |
If age is reported in mean indicate SD

|   |   |
|---|---|
| 21 | Age of all study participants, Range |
|   | Upper |
|   | Lower |
| 22 | HIV infection (%) |
| 23 | Participants included of female sex (%) |
| 24 | CD4 |
|   | __________ |
|   | If CD4 is reported in median indicate IQR |
|   | If CD4 is reported in mean indicate SD |
| 25 | Number (percent) of tuberculosis cases in the study (%) |
| 26 | What was the target condition? |
|   | 1. Pulmonary tuberculosis |
|   | 2. Extra pulmonary tuberculosis |
|   | 3. Mycobacteraemia |
|   | 4. Both 1 and 2 |
|   | 5. Any of 1,2,3 |
|   | 7. Other, specify |
| 27 | Did the study include patients with prior tuberculosis history? |
|   | 1. Yes |
|   | 2. No |
|   | 9. Unknown/Not reported |
|   | If yes, what is the % __________ |
|   | Specify the numerator/denominator _____/_____ |
| 28 | What was the clinical setting of the study? |
|   | 1. Outpatient |
|   | 2. Inpatient |
|   | 3. Both out-patient and in-patient |
|   | 7. Other, describe: |
|   | 9. Unknown/Not reported |
| 29 | How would you describe the health facility where the study took place? |
|   | 1. Primary care clinic, stand alone |
|   | 2. Primary care clinic, connected to a referral hospital |
|   | 3. Referral hospital |
|   | 7. Other, describe: |
|   | 9. Unknown/Not reported |
| 30 | Are there concerns that the included patients and setting do not match the review question? |
|   | 1. High concern |
|   | 2. Low concern |
|   | 9. Unclear concern |
| 31 | NOTES ON CHARACTERISTICS |

V. INDEX TEST

|   |   |
|---|---|
| 32 | Was a LF-LAM threshold used to define positivity that was pre-specified in the primary analysis? |
|   | 1. Yes, Grade 1/5 |
|   | 2. Yes, Grade 2/5 |
|   | 3. Yes, Grade 1/4 |
|   | 4. Yes, Grade 2/4 |
|   | 5. No |
|   | 7. Other, specify: |
| 33 | What LF-LAM threshold was used to define positivity for data extraction? |
|   | 1. Grade 2/5 |
|   | 2. Grade 1/4 |
|   |   |
|---|---|
|34| Are their concerns about index test conduct or interpretation differing from review question? | 1. High concern 2. Low concern 9. Unclear concern |
|35| Was LF-LAM performed on fresh or stored urine? | 1. Fresh 2. Stored, specify type of storage (e.g. frozen) 3. Both fresh and stored 9. Unknown/Not reported |
|36| Was LF-LAM result interpreted without knowledge of the result of the reference standard result? | 1. Yes 2. No 9. Unknown/Not reported/Unclear |
|37| Were there any LF-LAM results that were invalid (no bar in control window)? | 1. Yes a. Specify number of invalid tests: _____ b. Were invalid tests repeated (yes/no): _____ 2. No 9. Unknown/Not reported |
|38| Could the conduct or interpretation of the index test have introduced bias? | 1. High risk 2. Low risk 9. Unclear risk |
|39| NOTES ON INDEX TEST |

**VI. REFERENCE STANDARD**

|   |   |
|---|---|
|40| For the diagnosis of pulmonary tuberculosis, what reference standard was used to identify tuberculosis and not tuberculosis? | 1. Sputum: solid culture 2. Sputum: liquid culture 3. Sputum: both solid and liquid culture 4. Nucleic acid amplification test, specify 5. Any of culture or nucleic amplification test, specify 7. Other, specify |
|41| Was sputum induction performed for individuals unable to produce expectorated sputum? | 1. Yes Specify N/% requiring sputum induction _____ 2. No |
|42| Were patients without sputum specimens (for example, no expectorated, no induced sputum) included in this study? | 1. Yes Specify N/% included without sputum _____ 2. No Specify N/% excluded due to lack of sputum _____ |
|43| Were non-pulmonary specimens evaluated to allow diagnosis of extrapulmonary tuberculosis? | 1. All participants received testing of non-pulmonary specimens, please specify sites/fluids: 2. Some participants received testing of non-pulmonary specimens, please specify which patients were tested, and sites/fluids: 3. Extrapulmonary tuberculosis was not evaluated 7. Other, please specify: |
|44| For the diagnosis of extrapulmonary tuberculosis, what tests were used to identify tuberculosis and not tuberculosis (circle all that apply)? | 1. Solid culture 2. Liquid culture 3. Both solid and liquid culture 4. Nucleic acid amplification test, specify 7. Other, specify: __________________________ |
(Continued)

| No. | Question                                                                 | Yes | No | Unclear |
|-----|--------------------------------------------------------------------------|-----|----|---------|
| 45  | Did the study speciate mycobacteria isolated in culture?                 | 1. Yes | 2. No | 9. Unknown/Not reported |
| 46  | Was the reference standard likely to correctly classify the target condition | 1. Yes | 2. No | 9. Unclear |
| 47  | Was the reference standard result interpreted without knowledge of the result of LF-LAM? | 1. Yes | 2. No | 9. Unclear |
| 48  | How many sputum specimens were obtained in order to detect pulmonary tuberculosis? | 1. Single | 2. Multiple | 8. Not applicable |
| 49  | How many specimens from fluid (sites) other than sputum were obtained to detect extrapulmonary tuberculosis? | 1. Single | 2. Multiple | 8. Not applicable |
| 50  | Could the reference standard, its conduct, or its interpretation have introduced bias? | 1. High risk | 2. Low risk | 9. Unclear risk |
| 51  | Are there concerns that the target condition as defined by the reference standard does not match the question? | 1. High concern | 2. Low concern | 9. Unclear concern |

VII. FLOW AND TIMING

| No. | Question                                                                 | Yes | No | Unclear |
|-----|--------------------------------------------------------------------------|-----|----|---------|
| 53  | Was there appropriate interval between index test and reference standard | 1. Yes, specimens collected at the same time. | 2. No, specimens collected greater than 7 days apart | 9. Unclear |
| 54  | Did all patients receive a reference standard?                           | 1. Yes | 2. No | 9. Unclear |
| 55  | Did all patients receive the same reference standard?                    | 1. Yes | 2. No (answer no if clinicians chose sample types, or other differences in reference standards between patients) | 9. Unclear |
| 56  | Were all participants included in the analysis?                          | 1. Yes | 2. No | 9. Unclear |
| 57  | Could the patient flow have introduced bias?                             | 1. High risk | 2. Low risk | 9. Unclear risk |

NOTES ON FLOW AND TIMING

Lateral flow urine lipoarabinomannan assay for detecting active tuberculosis in people living with HIV (Review)
Abbreviations: IQR, interquartile range; SD, standard deviation.

VIII. TABLES: Tuberculosis detection against a microbiological reference standard

**Tuberculosis** is defined as positive culture or NAAT from sputum or any body fluid or site.

**Not Tuberculosis** is defined as negative cultures or NAATs from sputum or any body fluid or site.

(Table example; provide additional tables for each of the applicable questions)

| LAM result | Tuberculosis | Not tuberculosis | Total |
|------------|--------------|-----------------|-------|
| Positive   |              |                 |       |
| Negative   |              |                 |       |
| Total      |              |                 |       |

Provide additional tables for each of the questions:

What is the diagnostic accuracy of LF-LAM for the diagnosis of tuberculosis in all HIV-positive adults with advanced HIV disease and signs and symptoms of tuberculosis?

a. in inpatient setting CD4 ≤ 200
b. in outpatient setting CD4 ≤ 200
c. in all settings CD4 ≤ 200
d. in all settings CD4 > 200
e. in inpatient setting CD4 ≤ 100
f. in outpatient setting CD4 ≤ 100
g. in all settings CD4 ≤ 100
h. in all settings CD4 > 100
i. in inpatients settings CD4 101 to 200
j. in outpatient settings CD4 101 to 200
k. in all settings CD4 101 to 199

What is the diagnostic accuracy of LF-LAM for the diagnosis of tuberculosis in adults with advanced HIV disease irrespective of signs and symptoms of tuberculosis?

a. in all settings CD4 > 200
b. in all settings CD4 > 100
c. in inpatients settings CD4 101 to 200
d. in outpatient settings CD4 101 to 200
e. in all settings CD4 101 to 199

Appendix 6. QUADAS-2

**Domain 1: patient selection**

**Risk of bias: could the selection of patients have introduced bias?**

**Signaling question 1: Was a consecutive or random sample of patients enrolled?**

We answered ‘yes’ if the study enrolled a consecutive or random sample of eligible participants; ‘no’ if the study selected participants by convenience; and ‘unclear’ if the study did not report the manner of participant selection or we could not tell.

**Signaling question 2: Was a case-control design avoided?**

We answered ‘yes’ to all included studies given that we are excluding case-control study designs.

**Signaling question 3: Did the study avoid inappropriate exclusion?**

We answered ‘yes’ to studies which included all HIV-positive participants and participants who were unable to produce sputum (expectorated or induced). We answered ‘no’ if studies excluded participants who could not produce sputum (i.e. there were no attempts...
at sputum induction or patients could not produce sputum despite sputum induction and were excluded). We also answered ‘no’ if studies excluded patients presumed to have extrapulmonary tuberculosis. We scored ‘unclear’ if we could not tell.

Applicability: Are there concerns that the included patients and setting do not match the review question?

We were interested in how LF-LAM performs in patients whose urine specimens were evaluated as they would be in routine practice. We expected to judge ‘low concern’ for most studies since we planned to determine test accuracy both for patients with signs and symptoms of tuberculosis and patients investigated for tuberculosis irrespective of signs and symptoms for tuberculosis.

For LF-LAM used as a tuberculosis diagnostic test among patients with signs and symptoms of tuberculosis, we judged ‘high concern’ if the study participants did not resemble people with presumed HIV/tuberculosis; ‘low concern’ if the study population did resemble a population with presumed HIV/tuberculosis, and ‘unclear concern’, if we could not tell.

For LF-LAM used as a tuberculosis diagnostic test among patients that were investigated for tuberculosis irrespective of signs and symptoms of tuberculosis, we judged ‘low concern’ for studies in which the LF-LAM was performed uniformly within the predetermined study target populations of HIV-infected individuals, ‘high concern’ if LF-LAM was not performed uniformly within the predetermined study target populations of HIV-infected individuals, and ‘unclear concern’ if we could not tell. We judged ‘high concern’ if the study participants did not resemble people with presumed HIV/tuberculosis coinfection.

Domain 2: index test

Risk of bias: could the conduct or interpretation of the index test have introduced bias?

Signaling question 1: were the index test results interpreted without knowledge of the results of the reference standard?

We answered ‘yes’ if the study interpreted the result of LF-LAM blinded to the result of the reference standard; we answered ‘no’ if the study did not interpret the result of LF-LAM blinded to the result of the reference standard. We answered ‘yes’ for studies in which LF-LAM was performed on fresh specimens, since reference standard results would be unavailable at the time of test interpretation. We answered ‘unclear’ if stored specimens were tested and we could not tell if the index test results were interpreted without knowledge of the reference standard results.

Signaling question 2: if a LF-LAM threshold was used to define positivity, was it prespecified?

We answered ‘yes’ if the threshold was prespecified in the study or by the authors, ‘no’ if the threshold was not prespecified, and ‘unclear’ if we could not determine if the threshold was prespecified or not.

Applicability: are there concerns that the index test, its conduct, or its interpretation differ from the review question?

If index test methods vary from those specified in the review question, concerns about applicability may exist. We judged ‘high concern’ if the test procedure was inconsistent with the manufacturer recommendations, ‘low concern’ if the test procedure was consistent with the manufacturer recommendations, and ‘unclear concern’ if we could not tell. In cases where the primary study defined grade 1 of 5 as the positivity threshold, but where we were able to extract data at the manufacturer’s currently recommended positivity threshold, we judged ‘low concern’ for applicability.

Domain 3: reference standard

Risk of bias: could the reference standard, its conduct, or its interpretation have introduced bias?

Signaling question 1: is the reference standard likely to correctly classify the target condition?

HIV-infected tuberculosis patients may have pulmonary tuberculosis, extrapulmonary tuberculosis, or both pulmonary and extrapulmonary tuberculosis. A microbiological reference standard, primarily culture, is considered the gold standard for tuberculosis. Owing to the difficulties in diagnosing HIV-associated tuberculosis, it is recommended that multiple cultures from sputum and other specimens be evaluated.

We answered ‘yes’ when appropriate specimens were obtained for the diagnosis of HIV-associated tuberculosis. For presumed pulmonary tuberculosis, sputum specimens should be obtained for culture, NAAT, or both culture and NAAT. If the patient cannot produce sputum, induced sputum should be performed. For presumed extrapulmonary tuberculosis, specimens should be consistent with Standard 4 of the International Standards for Tuberculosis Care which states: “For all patients, including children, suspected of having extrapulmonary tuberculosis, appropriate specimens from the suspected sites of involvement should be obtained for microbiological and histological examination” (TB CARE I 2014). We answered yes if multiple specimens were collected from different sites for extrapulmonary tuberculosis. An Xpert® MTB/RIF test is recommended as the preferred initial microbiological test for suspected tuberculosis meningitis because of the need for a rapid diagnosis. We also answered ‘yes’ if studies followed a standardized approach of collecting appropriate specimens from “suspected sites of involvement”, for example, blood or lymph nodes on all patients.

We answered ‘no’ when the reference standard was restricted to sputum specimens or the reference standard was restricted to extrapulmonary specimens (for example, urine, blood, etc.). We also answered ‘no’ if a consistent approach was not followed for all patients.
(for example, some but not all patients with presumed tuberculosis lymphadenitis receive lymph node tissue sampling). We answered ‘unclear’ if we could not tell.

**Signaling question 2: were the reference standard results interpreted without knowledge of the results of the index test?**

We answered ‘yes’ if the study interpreted the result of the reference standard blinded to the result of LF-LAM, or if the reference standard result was reported on an automated instrument; ‘no’ if the study did not interpret the result of the reference standard blinded to the result of LF-LAM, and ‘unclear’ if we could not tell.

**Applicability: are there concerns that the target condition as defined by the reference standard does not match the question?**

In general, we thought there was low concern for almost included studies based on the current definitions of the reference standard. We judged ‘high concern’ if included studies did not speciate mycobacteria isolated in culture, ‘low concern’ if speciation was performed, and ‘unclear’ if we could not tell. We also judged high concern if there was no protocol to ensure a minimum standard of testing with a reference standard.

**Domain 4: Flow and timing**

**Risk of bias: could the patient flow have introduced bias?**

**Signaling question 1: was there an appropriate interval between the index test and reference standard?**

We expected urine specimens for LF-LAM and the reference standards to be obtained at the same time and answered ‘yes’ for all studies that meet this criterion, or if index and reference standard tests were performed on specimens collected no greater than seven days apart. We chose seven days as a time period during which either treatment of tuberculosis or natural progression of tuberculosis without treatment could impact test results. We answered ‘no’ if specimens were collected for index and reference standard tests greater than seven days apart, and ‘unclear’ if we could not tell.

**Signaling question 2: did all patients receive the same reference standard?**

We answered ‘yes’ if all participants in the study received the reference standard to confirm tuberculosis, and ‘unclear’ if we could not tell.

**Signaling question 3: were all patients included in the analysis?**

We determined the answer to this question by comparing the number of participants enrolled in the study with the number of participants included in the two-by-two tables. We answered ‘yes’ if all participants enrolled in the study were tested with results presented and accounted for. We answered ‘no’ if participants meeting enrolment criteria were not tested or results were not presented, and ‘unclear’ if we could not tell.

**Judgements for 'Risk of bias' assessments:**

If we answered all signalling questions for a domain "yes", then we judged risk of bias as "low".

If we answered all or most signalling questions for a domain "no", then we judged risk of bias as "high".

If we answered only one signalling question for a domain "no", we discussed further the "risk of bias" judgement.

If we answered all or most signalling questions for a domain "unclear", then we judged risk of bias as "unclear".

If we answered only one signalling question for a domain "unclear", we discussed further the "risk of bias" judgement for the domain.

**Appendix 7. Statistical appendix**

We list here the OpenBUGS program used to fit the bivariate meta-analysis models for estimating the accuracy of the index test. In the subsections below, we first describe the likelihood and prior distribution for the model followed by the OpenBUGS program.

As is usual with Bayesian models, initial values must be provided for all unknown parameters. We selected three independent sets of initial values for the parameters using the in-built ModelGenInit() function within OpenBUGS. The Gelman-Rubin statistic within the OpenBUGS program was used to assess convergence. We did not observe any convergence problems for the analyses presented. We treated the first 10,000 iterations as burn-in iterations and dropped them. We obtained summary statistics based on a total of 150,000 iterations resulting from the three separate chains.

**A. Estimation of index test accuracy**

**Notation:** in the $i$-th study the cells in the cross-tabulation between the index and reference tests are denoted by $TP_i$, $FP_i$, $TN_i$, $FN_i$. The sensitivity in $i$-th study is denoted by $s_{ei}$ and the specificity by $sp_i$.

We denote the Binomial probability distribution with sample size $N$ and probability $p$ as $\text{Binomial}(p,N)$, the Bivariate Normal probability distribution with mean vector $\mu$ and variance-covariance matrix $\text{TAU}$ as $\text{BVN}((\mu, \text{TAU}))$, the univariate Normal distribution with mean $m$ and variance $tau^2$ by $N(m, tau^2)$ and the Uniform probability distribution between $a$ and $b$ by Uniform$(a,b)$. Note that logit refers to log odds.

**Likelihood:**

Within studies:

\[ TP_i \sim \text{Binomial}(TPR_i, TP_i + FN_i) \text{, and} \]
\[ FP_i \sim \text{Binomial}(FPR_i, TN_i + FP_i) \]

Between studies:

The bivariate vector \((\text{logit}(TPR_i), \text{logit}(FPR_i)) \sim \text{BVN}(\mu = (\mu_1, \mu_2), \tau)\) where

\[ \tau \text{ is a 2 X 2 matrix with entries} \]
\[ \tau[1,1] = \text{variance of logit}(TPR_i) = \tau_1^2, \]
\[ \tau[2,2] = \text{variance of logit}(FPR_i) = \tau_2^2 \text{ and} \]
\[ \tau[1,2] = \tau[2,1] = \text{covariance between logit}(TPR_i) \text{ and logit}(FPR_i) = \rho \times \tau_1 \times \tau_2 \]

and \(\rho\) is the correlation between logit\((TPR_i)\) and logit\((FPR_i)\) across studies.

The pooled sensitivity is given by \(1/(1+\exp(-\mu_1))\), and the pooled specificity is given by \(1/(1+\exp(-\mu_2))\).

Prior distributions:

\[ \mu_1 \text{ and } \mu_2 \sim N(\mu = 0, \tau_2^2 = 4), \]
\[ \rho \sim \text{Uniform}(-1, 1) \]
\[ (1/\tau_1^2) \text{ and } (1/\tau_2^2) \sim \text{Gamma}(\text{shape}=2, \text{rate}=0.5) \]

A.1 OpenBUGS program for estimating a bivariate hierarchical meta-analysis model for sensitivity and specificity of the index test.

Observed data must be provided for \(L\) (the number of studies), and TP, FN, FP and TN in each study.

```sql
model {
  for(i in 1:L) {  # L is the number of studies in the Meta-analysis
    # Likelihood
    pos[i]<-TP[i]+FN[i]
    neg[i]<-TN[i]+FP[i]
    TP[i] ~ dbin(TPR[i],pos[i])
    FP[i] ~ dbin(FPR[i],neg[i])
    logit(TPR[i]) ~ dnorm(mu[1], T[1,1])
    logit(FPR[i]) ~ dnorm(mu[2], T[2,2])
    se[i] <- 1-TPR[i]
    }  # Prior Distributions
```
mu[1] ~ dnorm(0,0.25)
mu[2] ~ dnorm(0,0.25)
T[1:2,1:2] <- inverse(TAU[1:2,1:2])

# Between-study variance-covariance matrix
TAU[1,1] <- tau[1]*tau[1]
TAU[2,2] <- tau[2]*tau[2]
TAU[1,2] <- rho*tau[1]*tau[2]
TAU[2,1] <- rho*tau[1]*tau[2]

# prec is the between-study precision in the logit(sensitivity) and logit(specificity)
# rho is the correlation between logit(sensitivity) and logit(specificity) across studies
prec[1] ~ dgamma(2,0.5)
prec[2] ~ dgamma(2,0.5)
rho ~ dunif(-1,1)
tau[1] <- pow(prec[1],-0.5)
tau[2] <- pow(prec[2],-0.5)

# Pooled sensitivity and specificity
Pooled_S <- 1/(1+exp(-mu[1]))
Pooled_C <- 1/(1+exp(-mu[2]))

# Predicted sensitivity and specificity in a new study
l.new[1:2] ~ dmnorm(mu[],T[1,])
sens.new <- 1/(1+exp(-l.new[1]))
spec.new <- 1/(1+exp(-l.new[2]))

# An R function (SROC_BUGS) for creating a joint summary plot of sensitivity and specificity of the index test
# using results from the OpenBUGS program above can be found in the DTAplots R package
# available for download at https://www.nandinidendukuri.com/software/diagnostic-meta-analysis

Appendix 8. Summary plot of LF-LAM sensitivity and specificity for tuberculosis detection in symptomatic participants.

Figure 12.
Figure 12. Summary plot of LF-LAM sensitivity and specificity for tuberculosis detection in symptomatic participants. The blue circles represent individual study estimates for sensitivity and specificity for the studies among symptomatic participants. The size of the circle is proportional to the sample size of the study. The filled black circle is the pooled estimate for sensitivity and specificity. The solid red line marks the 95% credible region around the summary estimate, the dashed red line marks the 95% prediction region.

Appendix 9. Summary plot of LF-LAM sensitivity and specificity for tuberculosis detection in unselected participants.

Figure 13.
Figure 13. Summary plot of LF-LAM sensitivity and specificity for tuberculosis detection in unselected participants. The blue circles represent individual study estimates for sensitivity and specificity for the studies among unselected participants. The size of the circle is proportional to the sample size of the study. The filled black circle is the pooled estimate for sensitivity and specificity. The solid red line marks the 95% credible region around the summary estimate, the dashed red line marks the 95% prediction region.

Appendix 10. Unselected participants with CD4 > 200 and CD4 ≤ 200

Figure 14.
Figure 14. Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies among unselected participants, by CD4 strata (CD4 > 200 and CD4 ≤ 200) and health setting. The individual studies are ordered by decreasing sensitivity. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).

Unselected adults, CD4 > 200, all settings

| Study    | TP | FP | FN | TN | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|----------|----|----|----|----|----------------------|----------------------|----------------------|----------------------|
| Ejieren  | 3  | 1  | 8  | 144| 0.27 [0.06, 0.61]    | 0.99 [0.96, 1.00]    |                      |                      |

Unselected adults, CD4 ≤ 200, all settings

| Study    | TP | FP | FN | TN | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|----------|----|----|----|----|----------------------|----------------------|----------------------|----------------------|
| Ejieren  | 10 | 10 | 23 | 237| 0.45 [0.30, 0.61]    | 0.93 [0.96, 0.96]    |                      |                      |
| Hanifa   | 3  | 5  | 37 | 363| 0.07 [0.02, 0.20]    | 0.96 [0.97, 1.00]    |                      |                      |

Unselected adults, CD4 ≤ 200, inpatients

| Study    | TP | FP | FN | TN | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|----------|----|----|----|----|----------------------|----------------------|----------------------|----------------------|
| Ejieren  | 9  | 7  | 6  | 33 | 0.64 [0.35, 0.87]    | 0.82 [0.87, 0.93]    |                      |                      |

Unselected adults, CD4 ≤ 200, outpatients

| Study    | TP | FP | FN | TN | Sensitivity (95% CI) | Specificity (95% CI) | Sensitivity (95% CI) | Specificity (95% CI) |
|----------|----|----|----|----|----------------------|----------------------|----------------------|----------------------|
| Ejieren  | 10 | 12 | 16 | 204| 0.36 [0.19, 0.56]    | 0.94 [0.90, 0.97]    |                      |                      |
| Hanifa   | 3  | 5  | 37 | 363| 0.07 [0.02, 0.20]    | 0.96 [0.97, 1.00]    |                      |                      |

Appendix 11. Unselected participants with CD4 > 100 and CD4 ≤ 100

Figure 15.
Figure 15. Forest plots of LF-LAM sensitivity and specificity for tuberculosis against a microbiological reference standard for studies among unselected participants, by CD4 strata (CD4 > 100 and CD4 ≤ 100) and health setting. TP = True Positive; FP = False Positive; FN = False Negative; TN = True Negative. Between brackets are the 95% confidence interval (CI) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line).

Appendix 12. Characteristics of studies among children with HIV

| Study   | Country       | Age of enrolment                      | Presence of tuberculosis symptoms? | Setting               | Tuberculosis prevalence % (n/N) | Percent children with advanced or severe immunosuppression |
|---------|---------------|---------------------------------------|------------------------------------|-----------------------|--------------------------------|----------------------------------------------------------|
| Kroidl 2015 | Tanzania    | 6 weeks to 14 years, median 8.8 years (IQR 3.9 to 9.5)* | Tuberculosis symptoms            | Outpatient           | 40% (12/30)                      | 65%                                                      |
| LaCourse 2018a | Kenya     | ≤ 12 years, median 24 months (IQR 13 to 58) | Irrespective of tuberculosis symptoms | Inpatient            | 7% (9/130)                      | 70%                                                      |
| Nicol 2014   | South Africa| ≤ 15 years, median 42.5 months (IQR 19.1 to 66.3)* | Tuberculosis symptoms            | Inpatient and outpatient | 22% (23/160)                   | 53%                                                      |
aFor all participants, including HIV-positive and HIV-negative children.

IQR: interquartile range.

WHAT'S NEW

| Date            | Event                                      | Description                                                                                                                                 |
|-----------------|--------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------|
| 9 October 2019  | New citation required but conclusions      | We included 15 studies in this review update: nine new studies and six studies from the original review (Shah 2016) that met the inclusion criteria. The findings in this update are consistent with those reported previously (Shah 2016). |
| 9 October 2019  | New search has been performed              | A new search was performed. The review authors identified 15 unique studies, integrating nine new studies to six of those included in the previous Cochrane Review (Shah 2016). We used the terminology of lateral flow urine lipoarabinomannan assay (LF-LAM). The Alere Determine™ TB LAM Ag assay was the only LF-LAM test commercially available at the time of the review. New tests coming into the market, such as the FujiLAM, was not commercialized nor with any published data as of May 2018. We evaluated performance of LF-LAM only at grade 1 on the update reference scale card (scale of 1 to 4) equivalent to grade 2 on the old reference scale card (scale 1 to 5). We evaluated LF-LAM accuracy against a microbiological reference standard and did not include analysis against a composite reference standard. We grouped the analysis by studies among 'Symptomatic' and 'Unselected' participants instead of the previous terminology of 'Diagnosis' and 'Screening'. We changed the QUADAS judgements for some of the previously included studies. |

CONTRIBUTIONS OF AUTHORS

MS and SB reviewed articles for inclusion and extracted data. MS, SB, IS, ND, and KRS analysed the data. MS, SB, IS, ND, CMD, and KRS interpreted the analyses. SB, MS, and KRS drafted the manuscript in adults. RRN reviewed articles with impact data and drafted the summary sections on patient outcome data. MK, SB, MS, and KRS drafted the GRADE tables. ND and IS drafted the statistical analysis section and the statistical appendix (Appendix 7). All authors provided critical revisions to the manuscript. All review authors read and approved the final manuscript draft.

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RN is a board member of TB Proof and has participated on an advisory board for insmed, but these are not thought to represent a conflict of interest.

CMD is employed by FIND, a Swiss non-profit organization and WHO Collaborating Centre for Diagnosis. FIND provided funding for an initial assessment of data available for the review.

The review authors have no financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the review apart from those disclosed.
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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

Differences between review and review update
To further investigate for heterogeneity in data, we carried out additional subgroup analyses for the following CD4 strata: CD4 > 200; CD4 > 100 and CD4 101 to 200 cells/μL. We also performed a subgroup analysis by tuberculosis prevalence in the study population (pre-test probability). In the review, we also estimated the predicted sensitivity and specificity in a future study together with their 95% credible intervals (CrI), which we did not mention in the protocol.

INDEX TERMS

Medical Subject Headings (MeSH)
Biomarkers [urine]; CD4 Lymphocyte Count; HIV Seropositivity [*complications]; Lipopolysaccharides [*urine]; Point-of-Care Systems; Randomized Controlled Trials as Topic; Sensitivity and Specificity; Tuberculosis [*diagnosis]; Tuberculosis, Pulmonary [*diagnosis]

MeSH check words
Adult; Humans