Tuberculosis screening costs and cost-effectiveness in high-risk groups: a systematic review

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Abstract

Background: Systematic screening for active tuberculosis (TB) is a strategy which requires the health system to seek out individuals, rather than waiting for individuals to self-present with symptoms (i.e., passive case finding). Our review aimed to summarize the current economic evidence and understand the costs and cost-effectiveness of systematic screening approaches among high-risk groups and settings.

Methods: We conducted a systematic review on economic evaluations of screening for TB disease targeting persons with clinical and/or structural risk factors, such as persons living with HIV (PLHIV) or persons experiencing homelessness. We searched three databases for studies published between January 1, 2010 and February 1, 2020. Studies were included if they reported cost and a key outcome measure. Owing to considerable heterogeneity in settings and type of screening strategy, we synthesized data descriptively.

Results: A total of 27 articles were included in our review; 19/27 (70%) took place in high TB burden countries. Seventeen studies took place among persons with clinical risk factors, including 14 among PLHIV, while 13 studies were among persons with structural risk factors. Nine studies reported incremental cost-effectiveness ratios (ICERs) ranging from US$51 to $1980 per disability-adjusted life year (DALY) averted. Screening was most cost-effective among PLHIV. Among persons with clinical and structural risk factors there was limited evidence, but screening was generally not shown to be cost-effective.

Conclusions: Studies showed that screening is most likely to be cost-effective in a high TB prevalence population. Our review highlights that to reach the “missing millions” TB programmes should focus on simple, cheaper initial screening tools (i.e., symptom screen and CXR) followed by molecular diagnostic tools (i.e., Xpert®) among the highest risk groups in the local setting (i.e., PLHIV, urban slums). Programmatic costs greatly impact cost-effectiveness thus future research should provide both fixed and variable costs of screening interventions to improve comparability.

Keywords: Tuberculosis, Economic evaluation, Systematic review, Cost-effectiveness, Tuberculosis in HIV-infected, Tuberculosis control

Introduction

The 2020 Global Tuberculosis (TB) Report noted 3 million undiagnosed TB cases annually worldwide [1]. There is an urgent need to strengthen TB programme efforts to improve TB case detection of these missing millions. These efforts involve scaling up systematic screening programmes targeting household contacts and other
Economic evaluations may help in ensuring that limited resources are used wisely. The current published literature includes systematic reviews of screening interventions and their economic impact but focused on specific populations (i.e., persons experiencing homelessness or incarceration, PLHIV, immigrants or other high-risk groups) [12–17]. These reviews highlight that although systematic screening and active case finding is recommended in high-risk groups, there is a need for clearer guidance on which specific tools and screening algorithms or strategies are cost-effective, essentially highlighting the gap in knowledge that still exists despite WHO’s endorsement of systematic screening. Two systematic reviews of ICF among PLHIV showed significant variability across countries and target groups of patients, but highlighted that ICF was cost-saving compared to PCF in high TB/HIV burden countries, though authors noted the lack of standardized methods for cost data collection [5, 17]. The other reviews of the cost of TB screening were focused on a specific population and setting (i.e., immigrants in low-TB burden settings or contacts in Eastern Europe) [13–15] and recommended systematic screening for high-risk groups, but noted that there was limited data which was heterogeneous and of low quality. Therefore, our objective was to comprehensively synthesize economic evaluations of systematic screening for TB disease to inform a guideline development meeting leading to the updated guidance on TB screening, “WHO consolidated guidelines on tuberculosis. Module 2: screening—systematic screening for tuberculosis disease” [3] (see full list of PICO questions in the Additional file 1). Our study aimed to provide an up-to-date and comprehensive review of the economic evidence for all systematic screening interventions that was not limited to one sub-population, high-risk group, or setting.

Methods

Protocol

We performed a systematic review of the published literature on economic evaluations for TB screening with a focus on high-risk populations of persons with clinical risk factors such as PLHIV, diabetes or other respiratory diseases. We also focused on persons with structural risk factors, defined by the WHO as, “the circumstances in which people are born, grow up, live, work and age,” [18] including persons experiencing homelessness or residing in prisons, miners, elderly or indigenous persons. We sought to understand costs, cost-effectiveness, and affordability of screening approaches in key high-risk populations from the health system perspective. We performed this review according to the PRISMA guidelines (see Additional file 1) [19, 20].
Information sources
We searched three online databases: Ovid, EMBASE and Scopus for new studies published within the past ten years (i.e., January 1, 2010 through February 1, 2020). This review was intended to inform an update to the current WHO guidelines. Technologies used for screening and diagnosis of TB have significantly improved in recent years, such as the development of rapid molecular tests for TB including Xpert® MTB/RIF (Xpert) and other diagnostics that were not available before 2010. A search strategy was developed to identify cost and cost-effectiveness studies of systematic screening in high-risk groups. We reviewed citations of all eligible articles, guidelines, and reviews for additional studies (see Additional file 1 for search terms).

Study selection
Studies were included if they evaluated any type of systematic screening activities among persons with clinical or structural risk factors and included costs. Our search terms were designed to broadly capture any economic evaluations or studies including costs and an outcome, such as disability-adjusted life years (DALYs) or quality-adjusted life years (QALYs) and was not limited to cost-effectiveness or cost-utility analyses. Studies including utilities, such as DALYs, without costs were not included. Relevant studies were identified through electronic searches of the online databases, and duplicates were removed. Articles were excluded if they did not evaluate screening activities, or were reviews, letters, or opinion pieces.

Data collection
Articles were excluded if they only screened for latent TB infection, did not report costs per person screened or diagnosed or did not report the costs for the screening intervention separately from standard care (i.e., PCF). Studies were excluded if they evaluated screening activities in the general population, among contacts or children, since these groups were included in a separate manuscript (accepted, in press). Full text review was done independently by two reviewers (HA and BE) on remaining articles that met predetermined inclusion criteria, with all disagreements resolved by discussion with a third reviewer (AZ). No language filter was applied. Assessment of the quality of each economic evaluation and study quality was guided by the Consensus Health Economic Criteria (CHEC) [21, 22] and CHEERS checklist, respectively (see Additional file 1: Tables S1 and S2) [21].

Data extraction
The study data elements extracted from each study included: primary research question, country and setting, year of study, patient population, clinical setting, type of intervention, comparison diagnostic scenarios, economic analysis perspective, analytic time horizon, type of economic evaluation, source of costing, primary and secondary outcome measures, type of model, sensitivity and uncertainty analyses performed and willingness-to-pay (WTP) threshold. The WHO’s World Health Report guidelines on Choosing Interventions that are Cost-Effective (CHOICE) are the most commonly referenced WTP threshold among cost-effectiveness studies, particularly in low- and middle-income countries (LMICs) and are typically based on 1–3 times the country gross domestic product (GDP) per capita [23]. However, the use of these GDP-based thresholds has been challenged by many experts, the thresholds are considered overly simplistic and too easily attained when an intervention is effective [24]. Another key criticism of GDP-based WTP thresholds is the lack of their value and usefulness in assessing the trade-offs that decision-makers face in allocating limited healthcare resources [24]. Although the WHO no longer endorses GDP-based WTP thresholds [25], the challenge remains for clinicians, programme managers and researchers to determine the best metric for assessing value and reporting outcomes for cost-effectiveness and affordability of healthcare interventions. For the purposes of this review and to enable comparisons across currently published studies, we have included the GDP-based WTP thresholds used by study authors. However, we support the need for better decision-making tools for resource allocation in the local context, bearing in mind opportunity costs and the burden of disease.

Model parameters were extracted including epidemiologic, treatment and outcome parameters. Key outcomes included: cost per patient diagnosed, and incremental cost-effectiveness ratio (ICER) per utility measure (e.g., DALY averted). Costs are presented in United States Dollars (USD) adjusted to 2019 [27, 28]. Given the heterogeneity of study setting, year and type of screening strategies employed, there was no plan to calculate global estimates or pool data.

Results
Study selection
We identified a total of 3481 articles through database searching (Fig. 1). After duplicate removal, we screened 2318 citations by title and abstract for inclusion. Of these, we assessed 145 full-text publications against our inclusion criteria and excluded 118 publications. Exclusions were mainly due to the wrong intervention or no
economic evaluation. A total of 27 studies were identified for inclusion in the review [11, 29–53].

Study characteristics
Study characteristics for the 27 included studies are summarized in Tables 1 and 2; 19/27 (70%) of studies were conducted in high TB burden countries. Some studies included stratified analyses among multiple high-risk populations and thus contributed results to multiple categories (i.e., clinical and/or structural risk factors). Seventeen studies included persons with clinical risk factors; [32, 34, 38] fourteen among PLHIV, the majority of which (12/14) were conducted in Sub-Saharan Africa (SSA) [29–32, 37, 39–42, 44, 45, 49, 52, 53]. Thirteen studies included persons with structural risk factors (i.e., migrants, persons experiencing homelessness, or miners) and were from a range of countries such as Belgium, Cambodia, China, Russia, South Africa, and Zimbabwe [11, 32, 33, 35, 36, 38, 43, 46–48, 51, 54].

Study findings
We present the study findings stratified by high-risk subgroup, including persons with clinical and structural risk factors and PLHIV, as well as by screening tool used below.
| Study, year       | Country setting | Study population                        | Screening interventions and diagnostic tools | Reference screening strategies and diagnostic tools | Analysis perspective |
|-------------------|-----------------|-----------------------------------------|----------------------------------------------|---------------------------------------------------|----------------------|
| Abimbola et al. 2012 | South Africa    | PLHIV initiating ART                    | 1) WHO 4SS; SSM; SM (−) followed by CXR; CXR (−) followed by culture; 2) WHO 4SS; Xpert for diagnosis | Standard practice: SSM; SM (−) followed by CXR | Health system |
| Adelman et al. 2017 | Ethiopia        | PLHIV in HIV clinics                    | WHO 4SS; Xpert for diagnostic test of PLHIV with positive symptom screening (i.e. at least 1 symptom); Xpert (+) then DST/culture | Current recommended practice: 4SS; the SM and/or clinical diagnosis of TB | Health system |
| Andrews et al. 2012 | South Africa    | ARV naïve, PLHIV                         | Initial WHO 4SS then: 1) SSM (2 samples)—Xpert in TB-symptomatic; 2) SSM (2)—Xpert in all; 3) Culture (2)—Xpert in TB-symptomatic; 4) Culture (2)—Xpert in all; 5) Xpert (1) in TB-symptomatic; 6) Xpert (1) in all; 7) Xpert (2) in TB-symptomatic; 8) Xpert (2) in all | No TB screening | Health system |
| Bassett et al. 2010 | South Africa    | PLHIV initiating ART                    | ICF: WHO 4SS and CXR, then all PLHIV provided sputum for QIAmp (PCR) and culture | Symptoms (cough > 2 weeks) | Health system |
| Bogdanova et al. 2019 | Russian Federation | General population, PLHIV, homeless, migrants, chronic medical conditions | 1) Contact tracing done then CXR and SSM for diagnosis; 2) Mass screening in hospitals using CXR and SSM; 3) Mass screening in TB dispensary using mobile CXR and SSM | PCF including CXR and SSM | Health system |
| James et al. 2017 | Cambodia        | poor, urban residents; elderly living in rural areas | ACF interventions: 1) HOPE—Door-to-door screening WHO 4SS, symptomatic patients SSM (LED), then Xpert and culture; 2) CATA—Door-to-door screening older patients WHO 4SS, if symptomatic referred for CXR screening if abnormal then Xpert | N/A | Health system |
| Ji et al. 2020    | China           | Diabetic patients                       | ACF intervention—Patients received regular physical exams for 3 years; Diabetes patients all WHO 4SS, then positive symptom had CXR, then SSM, SM(−) then culture for TB confirmation | N/A | Health system |
| Study, year  | Country setting         | Study population                                                                 | Screening interventions and diagnostic tools                                                                 |
|-------------|-------------------------|----------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------|
| Jit et al. 2011 | United Kingdom          | Hard to reach individuals                                                        | Find and Treat service: 48 mobile CXR screening units offering: 1) mobile CXR; 2) enhanced case management; 3) referral for loss-to-follow-up |
| Jo et al. 2020  | Cambodia, Tajikistan     | Cambodia—elderly, vulnerable groups in rural areas Tajikistan—detention centers and diabetic patients | Cambodia—Community sensitization and training of CHWs, door-to-door WHO 4SS by CHWs, all patients had mobile CXR, abnormal CXR then Xpert Tajikistan—Community sensitization and training of TB staff, CHWs used mobile-phone questionnaire at health facilities WHO 4SS followed by CXR, SSM then Xpert based on SSM results |
| Karki et al. 2017 | Papua New Guinea        | General population in rural villages                                             | Outreach visits to villages: Systematic 4SS throughout villages, if symptomatic then SSM                     |
| Kranzer et al. 2012 | South Africa            | Peri-urban population attending mobile testing                                   | Mobile HIV testing van added TB testing: WHO 4SS for all HIV(-), if symptomatic then SSM, all HIV + SSM and referral to TB clinic for evaluation including CXR |
| Macheke et al. 2019 | Zimbabwe                | High-risk groups: PLHIV, contacts, miners, HCW, prisoners, elderly               | Zimbabwe ACF: WHO 4SS, then all have CXR, if either 4SS or CXR positive then Xpert with clinical diagnosis, if needed |
| Maheswaran et al. 2012 | Sub-Saharan Africa    | PLHIV                                                                            | 1) Any classic symptom; 2) 2+ classic symptoms; 3) CXR on all; 4) SSM on all; 5) 4SS, if positive then CXR; 6) 4SS, if positive then SSM; 7) 4SS, if positive then SSM then CXR; 8) 4SS if positive then CXR then SSM |
| Murray et al. 2016 | Uganda                 | General population                                                              | WHO 4SS, if cough > 2 weeks then triage testing with CXR or CRP then Xpert for diagnosis                   |

**Reference screening strategies and diagnostic tools**

PCF – people passively presenting to hospital

N/A

WHO algorithms:

WHO2b: 4SS, if positive then Xpert and clinical diagnosis, if needed

WHO2d: 4SS, if positive then CXR, then Xpert then clinical diagnosis

WHO3b: CXR, then Xpert and clinical diagnosis, if needed

Chronic cough > 2 weeks

PCF (baseline scenario): WHO 4SS, no triage testing, if symptomatic then Xpert

**Analysis perspective**

Health system
| Study, year | Country setting | Study population | Screening interventions and diagnostic tools | Reference screening strategies and diagnostic tools | Analysis perspective |
|-------------|-----------------|-----------------|----------------------------------------------|------------------------------------------------|---------------------|
| Orlando et al. 2018 | Mozambique | PLHIV | 1) Xpert: Xpert for all participants; 2) LAM: urine TB-LAM in all patients with CD4 < 200, Xpert in all patients with CD4 > 200 and TB-LAM(-) with CD4 < 200 | Standard: WHO 4SS, if symptomatic then SSM | Health system |
| Reddy et al. 2019 | Malawi, South Africa | Hospitalized PLHIV | KCF intervention: Sputum Xpert, urine LAM and urine Xpert; Modified intervention: Sputum Xpert and TB-LAM | Standard of care: Xpert | Health system |
| Sekandi et al. 2015 | Uganda | HHC, urban population | 1) PCF + ACF: HCW perform door-to-door chronic cough surveys (> 2 weeks) then collect 2 sputum for SSM, then return test results to the patient at their home; 2) PCF + HHC investigations (HCI): HCW perform WHO 4SS to HHC in home; child contacts and those unable to produce sputum diagnosed using CXR | PCF: Self-referral or presenting to health facility; chronic cough (> 2 weeks). WHO 4SS then SSM, if unable to produce sputum then diagnosed using CXR | Health system, societal perspective |
| Shah et al. 2017 | Peru | HHC (low HIV incidence setting) | 1) ACF. PCF + HCW visits to screen HHC using WHO 4SS and SSM, 2) PCF + Xpert; 3) ACF + Xpert: HCW visits to screen HHC using WHO 4SS and Xpert for diagnosis | PCF: Self-referral or presenting to health facility, TB diagnosis using SSM and clinical evaluation | Health system |
| Shah et al. 2009 | Ethiopia | PLHIV (18+ years) in VCT clinic | 1) WHO 4 SS then SSM, if SM() then CXR; 2) CXR for all PLHIV at entry, if CXR(+)(+) then SSM | N/A | Health system |
| Shah et al. 2008 | Vietnam | PLHIV | All PLHIV screened using CXR (all diagnosed HIV + before CXR), confirmed diagnosis with SSM | N/A | Health system |
| Smit et al. 2017 | Belgium | High risk groups and contacts (asylum seekers and migrants) | Systematic screening in high-risk groups including WHO 4SS and CXR; Follow-up of asylum seekers with abnormal CXR, supplemental CXR and periodic screening (6/12 months) after arrival | N/A | Health system |
| Sohn et al. 2019 | India | Rural, tribal population | CHW visits to homes with TB education, screening: WHO 4SS and SSM, CHW return with results | N/A | Health system |
| Study, year         | Country setting  | Study population | Screening interventions and diagnostic tools                                                                 | Reference screening strategies and diagnostic tools | Analysis perspective |
|---------------------|------------------|------------------|-------------------------------------------------------------------------------------------------------------|------------------------------------------------------|-----------------------|
| Winestsky et al. 2012 | Russian Federation | Prisoners        | 1) CXR screening; 2) WHO 4SS; 3) Annual Xpert screening; 4) WHO 4SS + CXR; 5) CXR + Xpert; 6) WHO 4SS + Xpert; 7) WHO 4SS + CXR + Xpert | PCF: No screening (self-referral)                      | Health system          |
| Yoon et al. 2019    | Uganda           | PLHIV in two HIV clinics | Novel ICF algorithms: 1) WHO 4SS + TB-LAM + Xpert; 2) WHO 4SS + TB-LAM + Xpert + culture; 3) POC CRP + Xpert; 4) POC CRP + TB-LAM + Xpert; 5) POC CRP + TB-LAM + Xpert + culture | Current ICF: WHO 4SS, if symptomatic then Xpert for diagnosis | Health system          |
| Zhang et al. 2016   | China            | High risk groups including elderly (65+) | 1) WHO A1: WHO 4SS, if cough > 2 weeks then CXR, if CXR(+) then SSM; 2) WHO A1B: WHO 4SS, if cough > 2 weeks then CXR, if CXR(+) then SSM; 3) WHO A2: WHO 4SS, if any TB symptoms then CXR, if CXR(+) then SSM; 4) WHO A3: Screening with CXR, if CXR(+) then SSM | N/A                                                  | Health system          |
| Zishiri et al. 2014 | South Africa     | Correctional facility inmates | WHO 4SS questionnaire for all newly admitted inmates, 1 + symptoms then diagnosed with Xpert | N/A                                                  | Health system          |
| Zwerling et al. 2015 | Malawi           | Newly diagnosed PLHIV in rural Malawi | Initial WHO 4SS, if 1 or more TB symptoms, then LED or Xpert; If initial 4SS negative, then patient asked to return 1 month later for second test if still symptomatic; Diagnosis with SSM, LED and Xpert | Standard of care (i.e. discretion of treating physician) using SSM | Health system          |

4SS four symptom screen, ACF active case finding, ART antiretroviral therapy, CE cost effective, CHW community health worker, CRP C reactive protein, CXR chest x-ray, DALY disability adjusted life year, DST drug-susceptibility testing, GDP gross domestic product, GNI gross national income, HCW healthcare workers, HHC household contacts, HIV human immunodeficiency virus, ICER incremental cost-effectiveness ratio, ICF intensified case finding, PCF passive case finding, PLHIV people living with HIV, SSM sputum smear microscopy, TB tuberculosis, USD United States dollars, WTP willingness to pay threshold, Xpert GeneXpert MTB/RIF, YLS year of life saved
| Study, year | Type of economic evaluation | Empirical study or modelling | Time horizon | Source of costing | Primary outcome | Sensitivity analysis | Key scenarios | WTP threshold |
|------------|-----------------------------|-----------------------------|--------------|------------------|----------------|---------------------|---------------|--------------|
| Abimbola et al. 2012 | Decision analytic model | Modelling | 6 months after ART initiation | South Africa specific published literature | ICER ($/TB death averted) | One-way and probabilistic | Mortality rates, cost inputs | Per-capita South African 2012 GDP ($5,678 USD) |
| Adelman et al. 2017 | Decision analytic model | Modelling | 10 years | Parent study results, AHIR HIV clinic, published literature | ICER ($/DALY averted) | One-way | Varied model inputs based on ranges in the literature | Per-capita Ethiopian 2014 GDP ($505 USD) |
| Andrews et al. 2012 | Decision analytic model | Modelling | Lifetime | Cape Town AIDS Cohort, unit costs from hospitals, published literature from South Africa | ICER ($/YLS) | One-way and two-way | Varied key parameters and costs | Per-capita South African 2010 GDP ($7,100 USD) |
| Bassett et al. 2010 | Cost analysis | Empirical | N/A | HIV clinic data (McCord hospital) | Cost ($/TB case identified) | N/A | Subset of patients with cough at study entry | N/A |
| Bogdanova et al. 2019 | Cost analysis | Empirical | N/A | Finance department, medical insurance fund | Cost ($/TB case detected) | N/A | N/A | N/A |
| James et al. 2017 | Cost analysis | Empirical | N/A | CENAT, National TB and Leprosy program, TB REACH | Cost ($/TB case detected) | N/A | N/A | N/A |
| Ji et al. 2020 | Cost-effectiveness analysis; cost-utility and cost benefit analyses | Modelling | 3 years | National Essential Public Health Program (NEPHS) | ICER ($/DALY gained) | One-way | Varied key parameters, and ROC curve | 3 times per capita 2016 GDP ($CNY not stated) |
| Jit et al. 2011 | Decision analytic model | Modelling | N/A | Find and Treat program records, NICE guidelines | ICER ($/QALY gained) | One-way | Varied all key parameters | 20–30,000 GBP per QALY |
| Jo et al. 2020 | Cost analysis | Empirical | 2 years | TB REACH budgets and financial reports, program data | Cost ($/TB case identified) | N/A | N/A | N/A |
| Karki et al. 2017 | Cost analysis | Empirical | N/A | Program expenditures and budget | Cost ($/TB case detected) | N/A | N/A | N/A |
| Kranzer et al. 2012 | Cost analysis | Empirical | 20 months | Study data and published literature | Cost ($/TB case detected) | One-way | Varied staff salaries and discount rates | N/A |
| MacEachera et al. 2019 | Cost analysis | Empirical | N/A | ZimACF project data | Cost ($/case diagnosed) | One-way | Varied all unit costs | N/A |
| Maheswaran et al. 2012 | Cost-effectiveness analysis | Modelling | 2 years | Published data | ICER ($/QALY) | Probabilistic | VOI used as alternative to univariate SA | Per capita 2010 SSA GNI ($2167 USD) |
| Murray et al. 2016 | Decision analytic model | Modelling | Lifetime | Published literature | ICER ($/life year gained) | One-way, multi-way, and probabilistic | | Per capita 2014 Ugandan GNI ($680 USD) |
| Study, year | Type of economic evaluation | Empirical study or modelling | Time horizon | Source of costing | Primary outcome | Sensitivity analysis | Key scenarios | WTP threshold |
|-------------|-----------------------------|-----------------------------|--------------|------------------|----------------|---------------------|--------------|---------------|
| Orlando et al. 2018 | Cost-effectiveness analysis | Modelling | 1 year | DREAM program, Global Fund data, published literature | ICER ($/DALY averted) | One-way | Varied all key parameters | Per capita 2016 Mozambique GDP ($382 USD) |
| Reddy et al. 2019 | Cost-effectiveness analysis | Modelling | 2 and 5 years, lifetime | STAMP trial, published literature | ICER ($/Year of life saved (YLS)) | One-way, multi-way | Varied all key parameters | Malawi ($750 USD); South Africa ($940 USD) |
| Sekandi et al. 2015 | Cost-effectiveness analysis | Modelling | 1.5 years | Uganda NTP, program data and published literature | ICER ($/additional TB case detected) | One-way | Varied costs and probabilities | Twice per capita 2013 Ugandan GDP ($1102 USD) |
| Shah et al. 2017 | Cost-effectiveness analysis | Modelling | N/A | Peru NTP data, published literature | ICER ($/DALY averted) | One-way | Varied all model parameters | Per capita 2014 Peruvian GNI ($6300 USD) |
| Shah et al. 2009 | Cost analysis | Empirical | N/A | Program data | Cost ($/TB case diagnosed) | N/A | N/A | N/A |
| Shah et al. 2008 | Cost analysis | Empirical | N/A | Provincial TB and HIV/AIDS program data | Cost ($/TB case diagnosed) | N/A | N/A | N/A |
| Smit et al. 2017 | Cost-effectiveness analysis | Empirical | 1 year | Flemish Association for Respiratory Health and TB Control | ICER (Euros/TB case detected) | One-way | Varied costs and number of cases detected | N/A |
| Sohn et al. 2019 | Cost analysis | Empirical | 1 year | Asha Kalp program finance and operations data | Cost ($/TB case detected) | One-way | Top-down and bottom-up cost estimates provided | N/A |
| Winestsky et al. 2012 | Decision analytic model | Modelling | 10 years | Primary data in prisons, published literature from Russia, Tajikistan and Latvia | ICER ($/QALY gained) | One-way, selected two-way, situational analyses | Varied all key parameters, and plausible situational analyses | Varied WTP thresholds |
| Yoon et al. 2019 | Cost analysis | Empirical | N/A | Program data | Cost ($/TB case diagnosed) | N/A | N/A | N/A |
| Zhang et al. 2016 | Cost analysis | Empirical | N/A | Program data | Cost ($/TB case diagnosed) | N/A | N/A | N/A |
| Zishiri et al. 2014 | Cost analysis | Empirical | N/A | Department of Correctional Services finance data, published literature | Cost ($/TB case identified) | One-way | Changed base-case parameters | N/A |
Table 2 (continued)

| Study, year     | Type of economic evaluation | Empirical study or modelling | Time horizon                     | Source of costing                                      | Primary outcome               | Sensitivity analysis                      | Key scenarios                          | WTP threshold                             |
|-----------------|-----------------------------|------------------------------|----------------------------------|--------------------------------------------------------|-------------------------------|-------------------------------------------|----------------------------------------|------------------------------------------|
| Zwerling et al. 2015 | Decision analytic model     | Modelling                    | Lifetime (assumed 59.2 years)    | Cost and operational analysis of four study sites, published literature | ICER ($/DALY averted)         | One-way, two-way and probabilistic uncertainty analysis | Varied all key parameters              | Per capita 2010 average SSA GDP ($1417 USD) |

4SS four symptom screen, ACF active case finding, ART antiretroviral therapy, CE cost effective, CRP C reactive protein, CXR chest x-ray, DALY disability adjusted life years, GDP gross domestic product, GNI gross national income, HCW healthcare workers, HIV human immunodeficiency virus, ICER incremental cost-effectiveness ratio, ICF intensified case finding, PCF passive case finding, PLHIV people living with HIV, TB tuberculosis, USD United States dollars, WTP willingness to pay threshold, Xpert GeneXpert MTB/RIF, YLS year of life saved.
Persons with clinical risk factors
Three studies provided cost-effectiveness data for individuals with the following clinical risk factors: diabetes mellitus, chronic respiratory disease and fibrotic lesions (Table 3) [32, 34, 38].

Diabetes and other medical co-morbidities
Bogdanova et al. assessed screening with chest X-ray (CXR) in Russia, and stratified results among diabetic patients as well as persons with other medical comorbidities (i.e., respiratory disease, gastro-intestinal, or fibrotic chest lesions); [32] reported costs ranged from US$11,648 to $105,754 per TB case diagnosed [32]. In Zimbabwe, Machekera et al. found that screening diabetic patients with CXR followed by Xpert® cost US$2191 per person diagnosed with TB [38]. Ji et al. reported that routine screening with CXR among Chinese patients with diabetes was considered highly cost-effective, as compared to POC, with an ICER of US$288 per DALY averted [34]. Machekera et al. included personnel and laboratory costs and had a large number needed to screen which drove up the cost per person diagnosed, while Ji et al. did not include the overhead costs for diabetic patient care.

People living with HIV (PLHIV)
There was significant heterogeneity of screening and diagnostic tools used among the 14 studies reporting the cost and cost-effectiveness of programmes in PLHIV (Table 4). Studies often considered multiple diagnostic algorithms or tools. Seven studies used molecular rapid diagnostic tests (i.e., Xpert®MTB/RIF) as an initial screening tool; [29–31, 40–42, 49, 52, 53] six studies used CXR to screen PLHIV in the outpatient setting; [31, 32, 39, 40, 44, 45] two studies used C-reactive protein (CRP) to screen; [40, 49] and one study used sputum smear microscopy (SSM) alone for screening [37].

Screening PLHIV with molecular rapid diagnostics
All seven studies that assessed screening in PLHIV using Xpert® MTB/RIF as an initial test were conducted in Sub-Saharan Africa, with 6/7 concluding screening was cost-effective. Two studies among PLHIV in South Africa conducted initial WHO-recommended four symptom screen (W4SS), including screening for cough of any duration, weight loss, fever or night sweats [55], followed by screening with Xpert® and found screening to be cost-effective with ICERS of US$324 per additional TB patient diagnosed and US$48,542 per TB death averted [29, 31]. Andrews et al. used two Xpert® to screen all PLHIV at a clinic in South Africa which was found to be cost-effective with an ICER of $4,096 per year of life saved (YLS) (WTP threshold: $7100 per YLS). [30] Among PLHIV initiating anti-retroviral therapy (ART), a combination of HIV treatment medications, in Mozambique, Orlando et al. showed that W4SS followed by screening with either Xpert® alone or Xpert® and lateral flow urine lipoarabinomannan assay (LF-LAM) was cost-effective, compared to W4SS and SSM, with ICERS ranging from US$37 to $51 per DALY averted [41]. Reddy et al. modeled the impact of a screening intervention with TB-LAM, urine and sputum Xpert® in hospitalized PLHIV, regardless of symptoms, in Malawi and South Africa [42]. The intervention was cost-effective, compared to sputum Xpert® alone (standard of care), with reported ICERS of US$450 and US$840 per YLS in Malawi and South Africa (WTP threshold: $750 and $940 per YLS, respectively).

Zwerling et al. found a randomized controlled trial using point of care (POC) Xpert® to screen PLHIV in rural Malawi was not cost-effective, due to low-test volumes (i.e., 50 tests/year, ICER of US$1980 per DALY averted, Uncertainty Range (UR): $1544–$3552). Zwerling et al. showed that Xpert® could be cost-effective at higher-test volumes (1000 tests/year, ICERs of US$398 per DALY averted, UR: $80–$1682) [WTP threshold: 3 × GDP per capita of Malawi (US$254)].

Screening PLHIV with CXR
Four of the six studies (67%) reporting on the use of CXR to screen PLHIV were conducted in SSA. The cost per person diagnosed with TB ranged from US$106 to $570 [31, 32, 44, 45]. Murray et al. found that community screening for cough followed by CXR in Uganda was cost-effective with an ICER of US$536 per year of life gained (YLG) (UR: $176–$2514) [40] Maheswaran et al. showed the W4SS followed by CXR in SSA was cost-effective with an ICER of US$6245 per QALY (UR: $6245–$19,581) [WTP threshold: 3 × GDP per capita of Malawi (US$2167)] [39].

Screening PLHIV with CRP
Among ART-naïve HIV clinic patients in Uganda, Yoon et al. found POC CRP followed by Xpert® for diagnosis [49] algorithms were cost-saving compared to W4SS followed by Xpert® (standard of care), ranging from US$69 to $92 per TB patient diagnosed. Murray et al. showed that CRP was cost-effective as a triage test in PLHIV in Uganda, compared to Xpert®, with an ICER of $517 per YLG (UR: $176–$2514) [WTP threshold: $3500 per DALY averted]. [39].

Screening among PLHIV using SSM
Kranzer et al. performed an intervention of adding TB symptom screening and SSM to an existing mobile HIV testing clinic in South Africa, [37] with an average cost of US$762 per TB case diagnosed [37].
Table 3  TB screening algorithm costs among persons with clinical and structural risk factors

| First author, Year | Country | Screening algorithm | Source of unit costs | Type of unit costs | Average cost of screening per person | Average cost of diagnosis per person |
|--------------------|---------|---------------------|----------------------|--------------------|--------------------------------------|-------------------------------------|
|                    |         |                     |                      |                    |                                      |                                    |
| Persons with structural risk factors (N = 16) |
| Migrants, refugees, internally displaced persons (IDPs) |
| Bogdanova 2019 Russia | Mass CXR screening | SSM Reported | ✓ ✓ ✓ | $4 per migrant screened | $834 per TB case diagnosed |
| Smit 2017 Belgium | WHO-4SS, CXR | NR Calculated | ✓ ✓ | $18 per migrant screened | $506,025 per migrant diagnosed with TB |
| Homeless persons and intravenous drug users (IDUs) |
| Bogdanova 2019 Russia | Mass CXR screening | SSM Reported | ✓ ✓ ✓ | $4-$13 | $793 per TB case diagnosed |
| Jit 2011 United Kingdom | Mobile CXR screening for IDUs and homeless | NR Reported | ✓ ✓ ✓ ✓ | | $9837 per QALY gained (UR: $6,302-$27,666) |
| Persons who live in urban slums |
| Sekandi 2015 Uganda | WHO-4SS for all HHC | SSM and CXR Reported | ✓ ✓ ✓ | NR | $1371 per additional TB case diagnosed |
| Shah 2017 Peru | Household visits for WHO-4SS for all HHCC | Xpert MTB/RIF Reported | ✓ ✓ ✓ ✓ ✓ | $32 per person screened | $3244 per DALY averted |
| James 2017 Cambodia | Door-to-door WHO 4SS | SSM and Xpert Reported | ✓ ✓ | NR | $268 per TB case diagnosed |
Table 3 (continued)

| First author, Year | Country | Screening algorithm | Source of unit costs | Type of unit costs | Average cost\(^1\) of screening per person | Average cost\(^1\) of diagnosis per person |
|---------------------|---------|---------------------|---------------------|-------------------|-----------------------------------------|-----------------------------------------|
| Members of tribal or indigenous populations | | | | | | |
| Sohn 2019 | India | Household visits for TB education and screening | SSM | Reported | ✓ | ✓ | ✓ | ✓ | < $1 per person screened | $3–5 per TB case diagnosed |
| Persons residing in prisons | | | | | | |
| Machekera 2019 | Zimbabwe | WHO 4SS, CXR | Xpert MTB/ RIF | Calculated | ✓ | ✓ | ✓ | ✓ | $14 per person screened | $460 per TB case diagnosed |
| Smit 2017 | Belgium | WHO 4SS, CXR | NR | Calculated | ✓ | ✓ | ✓ | ✓ | $18 per person screened | $14,034 per TB case diagnosed |
| Winetsky 2017 | Former Soviet Union | WHO 4SS alone | NR | Reported | ✓ | ✓ | ✓ | ✓ | $3 per person screened | $538 per QALY gained |
| Zishiri 2014 | South Africa | WHO 4SS | Xpert MTB/ RIF | Reported | ✓ | ✓ | ✓ | ✓ | $33 per person screened | $1,423 TB case diagnosed |
| Elderly (55+) | | | | | | |
| Jo 2020 | Cambodia (> 55) | Symptom screen | Xpert, SSM, CXR | Reported | ✓ | ✓ | ✓ | ✓ | < $1 per person screened | $406 per TB case diagnosed |
| James 2017 | Cambodia (> 55) | Symptom screen | Xpert, CXR | Calculated | ✓ | ✓ | ✓ | ✓ | < $2 per person screened | $340 per TB case diagnosed |
| Zhang 2017 | China (> 65) | Door-to-door symptom screening or CXR | CXR, SSM, culture | Reported | ✓ | ✓ | ✓ | ✓ | NR | $74–$315 per TB case diagnosed |
| People living in areas with limited access to healthcare (remote, isolated, hard-to-reach areas) | | | | | | |
| Karki 2017 | Papua New Guinea | Community-wide symptom screening | SSM | Reported | ✓ | ✓ | ✓ | NR | $158 per TB case diagnosed |
Table 3 (continued)

| First author, Year | Country | Screening algorithm | Source of unit costs | Type of unit costs | Average cost$ of screening per person | Average cost$ of diagnosis per person |
|--------------------|---------|---------------------|---------------------|--------------------|--------------------------------------|-------------------------------------|
| Jo 2020            | Cambodia| Symptom screen      | Xpert, SSM, CXR     | ✓                  | < $1 per person screened             | $406 per TB case diagnosed          |
|                    |         |                     |                     |                    |                                      |                                     |
| Miners (i.e., workers with silica exposure) | Machekera Zimbabwe 2019 | WHO 4SS, CXR | Xpert MTB/ RIF | Calculated | ✓ | $14 per person screened | $404 per miner diagnosed |
| Persons with clinical risk factors (N=3)$ |                   |                     |                     |                    |                                      |                                     |
| Diabetes mellitus  | Bogdanova 2019 | CXR | SSM | Reported | ✓ | ✓ | ✓ | ✓ | NR | $21,780 per TB case diagnosed |
| Ji 2020            | China    | CXR | SSM | Calculated | ✓ | ✓ | ✓ | ✓ | < $2 per person screened | $288 per DALY averted |
|                    | Machekera Zimbabwe 2019 | CXR | Xpert | Reported | ✓ | ✓ | ✓ | ✓ | NR | $2191 per TB case diagnosed |
| Respiratory disease | Bogdanova 2019 | CXR | SSM | Reported | ✓ | ✓ | ✓ | ✓ | NR | $32,746 per TB case diagnosed |
| Gastro-intestinal, genito-urinary, steroid use or fibrotic chest lesions | Bogdanova 2019 | CXR | SSM | Reported | ✓ | ✓ | ✓ | ✓ | NR | $11,648- $105,754 per TB case diagnosed |

$ There is limited evidence on screening among persons with structural risk factors, including migrants (n=2), homeless persons and IDUs (n=2), persons who live in urban slums (n=3), members of tribal or indigenous populations (n=1), persons residing in prisons (4), elderly (5), people living in remote areas (n=2) and miners (1). The costs of screening among persons with structural risk factors ranged from $1–33 per person screened and $3–$506,025 per TB case diagnosed. Screening programs were found to be cost-effective in persons living in urban slums and homeless, with reported ICERs of $3244 per DALY averted and $9837 per QALY, respectively. Screening was not shown to be cost-effective in migrants with an ICER of $506,025 per migrant diagnosed. In the Former Soviet Union, screening persons residing in prisons was found to be cost-effective with an ICER of $538 per QALY gained. Door-to-door screening of the elderly in China was shown to be cost-effective with ICERs ranging from US$74 to $315 per additional TB patient diagnosed.

$ There is limited evidence on screening among persons with clinical risk factors, primarily from one study in Russia. One study in Russia demonstrated a cost ranging from $11,648 to $105,754 for systematic screening among persons with various clinical risk factors (i.e., gastro-intestinal, respiratory disease, genito-urinary, steroid use and fibrotic chest lesions). Two studies conducted in Russia and Zimbabwe reported the cost per person diagnosed with diabetes mellitus (DM) which ranged from $2191 to $21,780. A cost-effectiveness analysis of patients with DM in China found systematic screening using CXR was cost-effective with an ICER of $288 per DALY averted.

1 Costs in 2019 USD unless stated otherwise

A/F active case finding, HHC household contacts, SSM sputum smear microscopy, Xpert GeneXpert, WHO 4SS four symptom screen, mWRDs molecular WHO-approved rapid diagnostics, CXR chest x-ray, NR not reported

✓✓ indicates cost component was explicitly included in unit cost calculation
Table 4  TB Screening Algorithm Costs in PLHIV

| First author, Year | Country       | Screening algorithm | Source of Unit Costs | Type of unit costs | Average cost \(^1,2\) of the screening algorithm per person | Average cost \(^1,2\) of diagnosis per person |
|-------------------|---------------|---------------------|---------------------|--------------------|----------------------------------------------------------|---------------------------------------------|
| Bassett 2010      | South Africa  | WHO 4SS, sputum     | Xpert MTB/RIF       | ✓ ✓ ✓              | $52 per person screened                                   | $324 per additional TB case diagnosed       |
| Andrews 2012      | South Africa  | Two Xpers for everyone | Xpert MTB/RIF      | ✓ ✓ ✓              | $37 per person screened                                   | $4,096 per YLS                              |
| Abimbola 2012     | South Africa  | WHO 4SS             | Xpert MTB/RIF       | ✓ ✓                | $26 per person screened                                   | $48,542 per death averted                   |
| Zwerling 2015     | Malawi        | WHO 4SS, Xpert      | Xpert MTB/RIF       | ✓ ✓ ✓              | $50 tests/year: $116 per person screened                 | $1,980 per DALY averted                     |
|                   |               |                     |                     |                    | (UR: $1,544-$3,552)                                       |                                             |
|                   |               |                     |                     |                    | 1000 tests/year: $18 per person screened                 | $398 per DALY averted (UR: $80–1682)        |
| Adelman 2017      | Ethiopia      | WHO 4SS, Xpert      | Xpert MTB/RIF       | ✓ ✓ ✓              | NR                                                       | $18 per person screened                    |
| Orlando 2018      | Mozambique    | Screening with Xpert | Xpert MTB/RIF       | ✓ ✓                | $10 per person screened                                   | $37 per DALY averted                        |
| Reddy 2019        | South Africa  | Sputum/urine Xpert, TB-LAM | Xpert MTB/RIF | ✓ ✓                | $31 per person screened* (Range: $11-$73)               | $802 per YLS                               |
|                   | Malawi        |                     |                     |                    | $72 per person screened* (Range: $18-$105)               | $596 per YLS                               |

PLHIV—using molecular rapid diagnostic tests (i.e., Xpert MTB/RIF) for screening (N = 7)\(^a\)

PLHIV—using CXR for screening (N = 6)\(^b\)

Shah 2008 Vietnam Monthly home visits to PLHIV with a CXR voucher NR Calculated from study data ✓ ✓ ✓ ✓ ✓ $7-$11 per person screened $115 per TB case diagnosed

Shah 2009 Ethiopia CXR for all PLHIV at entry to the clinic SSM on all positive CXR Calculated from study data ✓ ✓ $6 per person screened $106 per TB case diagnosed

Bassett 2010 South Africa WHO 4SS, CXR Xpert MTB/RIF Reported by study ✓ ✓ ✓ $52 per person screened $324 per additional TB case diagnosed (UR: $6,245-$19,581)

Maheswaran 2012 Sub-Saharan Africa (SSA) Symptom screen for any symptom SSM then CXR if smear negative Reported by study ✓ ✓ $11 per person screened (Range: $10-$29) $6,245 per QALY (UR: $6,245-$19,581)
In Sub-Saharan Africa, the average cost per person screened using rapid molecular diagnostic tests ranged from $3 to $116. Screening programs were found to be cost-effective in 6/7 (86%) of studies, with the following ICERs reported: $324 per additional TB case diagnosed; $37-$79 per DALY averted; $596–$4096 per YLS; and $517 per YLG. However, one study in Malawi found that screening with Xpert was not cost-effective at low test volumes, with an ICER of $1,980 per DALY averted. Cost-effectiveness depended on the volume of tests conducted annually and prevalence of TB.

Limited data was available for screening among outpatient PLHIV using CRPs from two studies both of which were of good quality. The average cost for test costs alone of outpatient PLHIV in Uganda screened using CRP with Xpert, TB-LAM and culture for diagnosis ranged from $13 to $34 per person screened and was shown to be cost-effective with an ICER of $517 per YLS or $69–$92 per additional TB case diagnosed in Uganda.

Limited data was available from one study conducted at an HIV testing clinic in South Africa for screening using SSM alone. This study reported an average cost of $762 per TB case diagnosed at a mobile testing clinic.

CRP: C-reactive protein, CXR: chest x-ray, mWRDs: molecular WHO-approved rapid diagnostics, NR: not reported. PLHIV: people living with HIV, POC: point-of-care, WHO 4SS: four symptom screen, Xpert: GeneXpert, YLG: year of life gained, YLS: year of life saved.

Table 4 (continued)

| First author, Year | Country | Screening algorithm | Source of Unit Costs | Type of unit costs | Average cost of the screening algorithm per person | Average cost of diagnosis per person |
|-------------------|---------|---------------------|---------------------|-------------------|---------------------------------------------|-------------------------------------|
| Murray 2016       | Uganda  | Screening for cough, CXR triage test | Reported by study | ✓ ✓ ✓ | $22 per person screened | $536 per YLG (UR: $176-$2,514) |
| Bogdanova 2019    | Russian Federation | Mass CXR screening programs in PLHIV | Calculated from study data | ✓ ✓ ✓ ✓ | $4-$7 per person screened | $570 per TB case diagnosed |
| Yoon 2019         | Uganda  | Screening for cough, CXR triage test | Reported by study | ✓ ✓ | $21 per person screened | $517 per YLG (UR: $194-$1,535) |
| Murray 2016       | Uganda  | Screening for cough, CXR triage test | Reported by study | ✓ ✓ | $21 per person screened | $517 per YLG (UR: $194-$1,535) |
| Kranzer 2012      | South Africa | WHO 4SS; SSM at a mobile HIV testing clinic | Reported by study | ✓ ✓ ✓ | NR | $762 per TB case diagnosed |

1 Costs in 2019 USD unless stated otherwise
2 All costs reported in 2019 USD unless stated otherwise
Persons with structural risk factors
Thirteen studies provided cost effectiveness data for individuals with structural risk factors including (Table 3): migrants, [32, 47] persons experiencing homelessness, [11, 32] persons who live in urban slums, [33, 43, 46] members of indigenous populations, [54] persons residing in prison, [38, 47, 48, 51] elderly, [33, 35, 50] people living in remote areas, [35, 36] and miners [38].

Migrants, persons experiencing homelessness and intravenous drug users (IDUs)
Bogdanova et al. assessed the cost of screening with CXR in Russia in multiple subgroups and found a cost of US$834 and US$793 per migrant and homeless person diagnosed with TB, respectively [32]. Jit et al. demonstrated that a programme using mobile screening vans among persons experiencing homelessness and IDUs in London, United Kingdom was cost-effective, compared to PCF, with an ICER of USD $9837 (UR: $6301–$28,666) per QALY gained (WTP threshold: £20,000–£30,000 per QALY gained set by NICE standards) [11]. However, in Belgium, Smit et al. showed CXR screening among migrants from high TB incidence countries was not cost-effective, compared with PCF, with an ICER of $506,025 (95% UR: $90,686–$2,040,006) per additional TB case diagnosed [47].

Persons who live in urban slums
Three studies examined door-to-door screening interventions in urban slums. James et al. reported that door-to-door symptom screening in Cambodian slums using CXR followed by Xpert® for diagnosis cost US$268 per TB case diagnosed [33]. Shah et al. demonstrated that household visits to screen all contacts of persons with TB in an urban slum in Lima, Peru was cost-effective, compared to PCF, with an ICER of US$3244 per DALY averted [WTP threshold: 2014 per capita GNI for Peru (US$6360)]. [46] However, Sekandi et al. found that door-to-door symptom screening of all household contacts, followed by SSM and CXR in urban Uganda, was not cost-effective compared to PCF with an ICER of US$1371 per additional TB case diagnosed [WTP threshold: twice the 2012 Ugandan per capita GDP (US$551)]. [43].

Members of indigenous populations
Sohn et al. reported a cost of US$3–$5 per person detected with TB for a programme of home visits by community health workers (CHWs) to indigenous persons in rural India. This study included the costs for CHWs time to travel to homes, conduct screening and diagnostic visits, provide directly observed therapy (DOT) and care, as well as the laboratory and administrative services, [54] but did not account for routine TB care and medications provided by the Indian government.

Persons residing in prisons
Four studies provided direct evidence for the costs of screening among persons residing in prisons [38, 47, 48, 51]. Zishiri et al. found W4SS followed by Xpert® for all persons residing in prison in South Africa cost US$1423 per person diagnosed. [51] While in Zimbabwe, Machekera et al. reported that an intervention of W4SS and CXR followed by Xpert® cost US$460 per person diagnosed [38]. Winetsky et al. performed a dynamic transmission model of TB among persons residing in prisons in the Former Soviet Union and demonstrated that annual screening with Xpert® was cost-effective compared with mass CXR screening, with an ICER of US$538 per QALY gained [WTP threshold: per capita GDP of Tajikistan (US$1900)]. [48] Smit et al. demonstrated that systematic screening of persons residing in prisons using W4SS followed by CXR in Belgium was cost-effective, compared to PCF, with an ICER of US$14,034 (95% UR: $10,898–$18,033) per additional case of TB detected. [47].

Elderly
Two studies were conducted in Cambodia of door-to-door screening using the W4SS followed by CXR and Xpert® for diagnosis among the elderly (55+), and reported average costs ranging from US$340 to $406 per person diagnosed with TB [33, 35]. Zhang et al. found that door-to-door symptom screening and CXR among the elderly in China (65 +) had costs ranging from US$74–$315 per TB case diagnosed [50]. Among elderly persons in China, additional risk factors (i.e., male, tobacco use or close TB contact) were associated with higher average costs per patient diagnosed [50].

People living in remote areas
In Papua New Guinea, Karki et al. reported that an intervention using SSM to screen all villagers with symptoms of TB cost US$158 per person detected with TB. [36] A second intervention, conducted by Jo et al. in remote Cambodia, included house-to-house symptom screening followed by mobile CXR and reported a cost of US$406 per person diagnosed with TB [35].

Miners
There was limited evidence for screening in miners from one study in Zimbabwe. [38] Machekera et al. demonstrated that a screening algorithm of CXR followed by Xpert® among those with positive CXR was cost saving, with a cost of US$404 compared to US$576 per person detected.
diagnosed with an algorithm screening everyone with W4SS and CXR.

Discussion
Our review of the published literature identified 27 studies of systematic screening among high-risk groups for TB, such as PLHIV, miners, persons residing in prisons, and the elderly. Our review found that systematic screening approaches are most likely to be cost-effective in settings and or populations with high prevalence of TB, such as PLHIV, persons residing in prisons and urban slum dwellers. Studies demonstrated that initial screening with more costly diagnostic tests was not cost-effective in high-risk groups, except among PLHIV. Simple, inexpensive initial screening methods (i.e., W4SS or CXR) followed by molecular diagnostic tests (i.e., Xpert®) among the highest risk groups in the local setting are the most cost-effective approaches to systematic screening. In high TB prevalence settings, door-to-door symptom screening in densely populated areas (i.e., slums), was generally shown to be cost-effective. However, mobile CXR units were not cost-effective due to high programmatic costs, particularly when interventions were targeting hard to reach populations (i.e., persons experiencing homelessness or IDUs). There was limited evidence identified for each high-risk group included in this review, thus caution should be used when extrapolating from a small number of studies.

Our review found the most evidence for cost-effectiveness of screening programmes among PLHIV. Despite varying screening strategies (i.e., W4SS, CRP, CXR, Xpert®, MTB/RIF) and patient settings (i.e., in-patient or outpatient clinic), the majority 9/10 (90%) of PLHIV studies that calculated an ICER found screening interventions to be cost-effective among PLHIV using an author determined WTP threshold. Key drivers of costs among PLHIV included: annual test volume and diagnostic test costs, [52] underlying prevalence of TB and HIV, [42, 52] clinic setting, [32, 44, 45] and programmatic costs included (i.e., transportation, mobile van units, or staffing costs) [31, 32, 55]. In sensitivity analyses of low-TB prevalence settings among PLHIV, screening strategies with simple tools, such as W4SS and SSM, were cost-saving compared to more expensive tools such as CXR or Xpert®, [39] highlighting the importance of the local setting.

Screening interventions in high TB prevalence settings, such as urban slums or among persons residing in prisons, increased identification and diagnosis of people with TB and were shown to be cost-effective [46, 48]. Equally important was ensuring proper follow-up to avoid treatment failure or loss post-screening [44]. However, among groups with other medical conditions, high programmatic screening costs, coupled with low TB prevalence meant screening interventions were not cost-effective, but was limited to two studies [32, 38].

Systematic screening is an expensive undertaking, particularly compared to PCF, when it involves mobilizing staff to go into the community. Evidence from this review suggests that community-based interventions [43] had higher costs compared to systematic screening programmes targeting persons presenting to healthcare facilities [35]. The use of artificial intelligence (AI) to inform screening tools through the use of CAD software is an exciting prospect for improving the efficiency and affordability of screening from standard CXR. However, our review did not find any studies which used CAD software in the context of a systematic TB screening programme. Indirect evidence, not presented in this review, has shown that the unit costs for each CXR read with CAD software are likely to be small but require significant investment in equipment and maintenance costs as well as the purchase of CAD software. These new technologies are still being evaluated in many programmes and highlight the need for costing and cost-effectiveness studies to inform their use in the programmatic setting.

The WTP thresholds, which are noted a priori by study authors and based on WHO recommendations, determine whether a given ICER is considered cost-effective. Among the papers reviewed in this analysis, many used either a country’s GDP, or twice the GDP, as the WTP threshold, however there was significant heterogeneity due to the range of country settings. This variability makes comparisons challenging, particularly since setting a higher threshold increases the likelihood of an intervention being considered cost-effective. Careful attention should be paid to the WTP threshold employed by study authors when interpreting cost-effectiveness. Furthermore, there are concerns about using GDP per capita as the basis of determining cost-effectiveness, particularly in LMICs where there are more stringent resources constraints [23]. A key concern is that using a threshold that is too low (i.e., GDP per capita) may result in health systems choosing not to adopt an intervention that would generate net health benefits because the threshold does not take into account health opportunity costs [56]. Current efforts to develop country-specific estimates that account for opportunity costs, as well as updated data on population and economic growth, are underway and aim to provide better options for informing decision-making and resource allocation for health interventions [24, 25, 56].

Limitations
The heterogeneity around reporting of costs and costing components made comparisons across
studies challenging. For instance, some costing analyses accounted for all operational and personnel-related costs, and thus reported higher total costs, while other analyses report only direct costs related to diagnostics testing [11, 38]. There were no standardized screening algorithms, even across similar high-risk groups, and employed different standards of care (i.e., PCF or symptom screen alone or with CXR) which limits comparability of studies and generalizability to other settings. Not all studies describe cost components in the same manner [11] and comparisons across studies is further impeded by a range of primary outcomes from cost per case diagnosed, $/DALYs averted, $/QALY, or $/YLS. Thus, even among the studies that do calculate an ICER, direct comparison is not necessarily appropriate. No included studies assessed the impacts of screening on earlier case detection and proper TB treatment, but this is an area that merits additional evidence to better understand the impact on cost and cost-effectiveness of preventing additional disease transmission.

Our study was restricted to the published literature and thus is likely impacted by publication bias towards those interventions that were shown to be cost-effective; programmes that were not deemed cost-effective may not have been published. A recent Task Force Report from the Professional Society for Health Economics and Outcomes Research (ISPOR) suggests benchmarking approaches, such as reviewing trial protocols, to better explore the potential for publication bias but also notes the need to develop new approaches to assess publication bias [57]. Further, more recent economic evaluations of screening interventions, particularly for novel diagnostic tests such as Xpert® may have been more likely to demonstrate cost effectiveness than earlier studies due to consistently decreasing test costs.

Our review highlights key gaps in the existing economic evidence, namely the need for more studies on the costs and cost-effectiveness of systematic TB screening programmes from Latin America and Asia, since the majority of included studies took place in SSA. Access to various screening and diagnostic tools was not consistent across study settings. Increased efforts should be made to ensure availability of newer diagnostic technologies to TB programmes globally. In addition, standardization of systematic screening interventions, along with fixed and variable costs included in economic analyses of programmes, is needed for better evidence generation and comparability across studies.

Conclusions
The COVID-19 pandemic has dramatically impacted TB services globally. Modelling has shown that COVID-19 related restrictions and interruptions to TB programmes may result in an increase in TB incidence up to 6.3 million, and mortality to 1.5 million, by 2025 [58, 59]. Our review is the first to summarize the economic evidence for systematic screening for TB disease among high-risk groups. Our review highlights that to reach the “missing millions”, and address the setbacks to TB services due to the COVID-19 pandemic, TB programmes should focus on simple, cheaper initial screening tools (i.e., symptom screen and CXR) followed by molecular diagnostic tools (i.e., Xpert®) among the highest risk groups in the local setting (i.e., PLHIV, urban slums). Programmatic costs greatly impact cost-effectiveness thus future research should provide both fixed and variable costs of screening interventions to improve comparability. COVID-19 has dramatically increased the number of digital applications for contact screening, as well as other video or tele-health options for service delivery. Expanded use of such digital technologies can be leveraged for TB screening to improve identification and treatment options for patients globally.

Abbreviations
WASS: WHO-recommended four symptom screen; CRP: C-reactive protein; CXR: Chest x-ray; DALY: Disability-adjusted life year; DM: Diabetes mellitus; DOT: Directly-observed therapy; GDP: Gross domestic product; GNI: Gross national income; ICER: Incremental cost-effectiveness ratio; PCF: Passive case finding; PLHIV: People living with HIV; POC: Point of care; QALY: Quality-adjusted life year; SSA: Sub-Saharan Africa; SSM: Sputum smear microscopy; TB: Tuberculosis; WHO: World Health Organization; UR: Uncertainty range; USD: United States dollar; WTP: Willingness-to-pay; Xpert®, GeneXpert®, YLG: Year of life gained; YLS: Year of life saved.

Supplementary Information
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Additional file 1. Annex S1. Key PICO questions and full search terms used in systematic review.

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Authors’ contributions
AZ and HA wrote the study protocol. All authors (HA, BE and AZ) contributed to data extraction, analysis and interpretation. HA wrote the first version of the manuscript. All authors (HA, BE, and AZ) reviewed the manuscript and accepted the final version of the paper. All authors read and approved the final manuscript.

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Availability of data and materials
Data sharing is not applicable to this article as no datasets were generated or analyzed during the current study.
Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

We declare no competing interests.

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References

1. World Health Organization. Global tuberculosis report 2020. Geneva: WHO; 2020.
2. Golub JE, Mohan CJ, Comstock GW, et al. Active case finding of tuberculosis: historical perspective and future prospects. Int J Tuberc Lung Dis. 2005;9(11):1183–203.
3. WHO consolidated guidelines on tuberculosis: Module 2: screening—systematic screening for tuberculosis disease. Geneva:2021.
4. Fox GJ, Johnston JC, Nguyen TA, et al. Active case-finding in contacts of people with TB. Int J Tuberc Lung Dis. 2021;25(2):95–105. https://doi.org/10.5558/ijtld.20.0656.
5. Kranzer K, Houben RM, Glynn JR, et al. Yield of HIV-associated tuberculosis during intensified case finding in resource-limited settings: a systematic review and meta-analysis. Lancet Infect Dis. 2010;10(2):93–102. https://doi.org/10.1016/S1473-3099(09)70326-3.
6. Kuupiel D, Bawontou V, Mashamba-Thompson TP. Mapping evidence on tuberculosis active case finding policies, strategies, and interventions for tuberculosis key populations: a systematic scoping review protocol. Syst Rev. 2019;8(1):162. https://doi.org/10.1186/s13643-019-1098-1.
7. Singh M, Sagili KD, Tripathy JP, et al. Are treatment outcomes of patients with tuberculosis detected by active case finding different from those detected by passive case finding? J Glob Infect Dis. 2020;12(1):28–33. https://doi.org/10.4103/jgid.jgid_66_19.
8. Shriram V, Sinhari R, Gayatri T, et al. Active case finding for Tuberculosis among migrant brick kiln workers in South India. Indian J Tuberc. 2020;67(1):38–42. https://doi.org/10.1016/j.ijtb.2019.09.003.
9. Jeon CY, Murray MB. Diabetes mellitus increases the risk of active tuberculosis: a systematic review of 13 observational studies. PLoS Med. 2008;5(7):e152. https://doi.org/10.1371/journal.pmed.0050152.
10. Sawert H, Kongsin S, Payanandana V, et al. Costs and benefits of improving tuberculosis control: the case of Thailand. Soc Sci Med. 1997;44(2):1805–16. https://doi.org/10.1016/s0747-9563(96)00289-4.
11. Jit M, Stagg HR, Aldridge RW, et al. Dedicated outreach service for hard to reach patients with tuberculosis in London: observational study and economic evaluation. BMJ. 2011;343:d5376. https://doi.org/10.1136/bmj.d5376.
12. Silva EN, Pereira A, de Araujo WN, et al. A systematic review of economic evaluations of interventions to tackle tuberculosis in homeless people. Rev Panam Salud Publica. 2018;42:40. https://doi.org/10.26633/RPS.2018.40.
13. Tavochi L, Vroling H, Maddedu G, et al. Active case finding for communicable diseases in prison settings: increasing testing coverage and uptake among the prison population in the European union/European economic area. Epidemiol Rev. 2018;40(1):105–20. https://doi.org/10.1093/epirev/mxy001.
14. Fox GJ, Barry SE, Britton WJ, et al. Contact investigation for tuberculosis: a systematic review and meta-analysis. Eur Respir J. 2013;41(1):140–56. https://doi.org/10.1183/09031996.00070812.
15. Zennor D, Southern J, van Hest R, et al. Active case finding for tuberculosis among high-risk groups in low-incidence countries. Int J Tuberc Lung Dis. 2013;17(5):573–82. https://doi.org/10.5588/ijtld.12.0920.
16. Dasgupta K, Menates D. Cost-effectiveness of tuberculosis control strategies among immigrants and refugees. Eur Respir J. 2005;25(6):1107–16. https://doi.org/10.1183/09031936.05.0004004.
17. de Siqueira-Filho NT, Legood R, Cavalcanti A, et al. Cost of tuberculosis diagnosis and treatment in patients with HIV: a systematic literature review. Value Health. 2018;21(4):482–90. https://doi.org/10.1016/j.jval.2017.09.003.
18. World Health Organization. Social determinants of health. Key concepts 2013. https://www.who.int/news-room/q-a-detail/social-determinants-of-health-key-concepts accessed November 23 2020.
19. Liberati A, Altman DG, Tetzlaff J, et al. The PRISMA statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: explanation and elaboration. Ann Intern Med. 2009;151(4):W65-94. https://doi.org/10.7326/0003-4819-151-4-200908180-00136.
20. Moher D, Shamseer L, Clarke M, et al. Preferred reporting items for systematic review and meta-analysis protocols (PRISMA-P) 2015 statement. Syst Rev. 2015;4:1. https://doi.org/10.1186/2046-4053-4-1.
21. Husereau D, Drummond M, Petrosu S, et al. Consolidated health economic evaluation reporting standards (CHEERS) statement. BMJ. 2013;346:f1049. https://doi.org/10.1136/bmj.f1049.
22. Husereau D, Drummond M, Petrosu S, et al. Consolidated health economic evaluation reporting standards (CHEERS)—explanation and elaboration: a report of the ISPOR health economic evaluation publication guidelines good reporting practices task force. Value Health. 2013;16(2):231–50. https://doi.org/10.1016/j.jval.2013.02.002.
23. Leech AA, Kim DD, Cohen JT, et al. Use and misuse of cost-effectiveness analysis thresholds in low- and middle-income countries: trends in cost-per-DALY studies. Value Health. 2018;21(7):759–61. https://doi.org/10.1016/j.jval.2017.12.016.
24. Maurice E, Larson B, Kazi DS, et al. Thresholds for the cost-effectiveness of interventions: alternative approaches. Bull World Health Organ. 2015;93(2):118–24. https://doi.org/10.2471/BLT.14.138206.
25. Bertram MV, Lauer JA, De Joncheere K, et al. Cost-effectiveness thresholds: pros and cons. Bull World Health Organ. 2016;94(12):925–30. https://doi.org/10.2471/BLT.15.164418.
26. Organization WH. New cost-effectiveness updates from WHO-CHOICE Geneva: World Health Organization; https://www.who.int/news-room/ feature-stories/detail/new-cost-effectiveness-updates-from-who-choice accessed August 13, 2021.
27. Turner HC, Lauer JA, Tran BX, et al. Adjusting for inflation and currency changes within health economic studies. Value Health. 2019;22(9):1026–32. https://doi.org/10.1016/j.jval.2019.03.021.
28. World Data Inflation Rate Calculator Oldenburg, Germany. https://www.worlddata.info/ accessed June 29 2020.
29. Abimbola TO, Marston BJ, Date AA, et al. Cost-effectiveness of tuberculosis diagnostic strategies to reduce early mortality among persons with advanced HIV infection initiating antiretroviral therapy. J Acquir Immune Defic Syndr. 2012;60(1):e1-7. https://doi.org/10.1097/QAIb013e318246538f.
30. Andrews JR, Lawn SD, Rusu C, et al. The cost-effectiveness of routine tuberculosis screening with Xpert MTB/RIF prior to initiation of antiretroviral therapy: a model-based analysis. AIDS. 2012;26(8):987–95. https://doi.org/10.1097/QAD.0b013e3283224d47.
31. Bassett NV, Wang B, Chetty S, et al. Intensive tuberculosis screening for HIV-infected patients starting antiretroviral therapy in Durban, South Africa Clin Infect Dis. 2010;51(7):823–9. https://doi.org/10.1086/656282.
32. Bogdanova E, Mariandyszhev O, Hindenaker SC, et al. Mass screening for active case finding of pulmonary tuberculosis in the Russian Federation: how to save costs. Int J Tuberc Lung Dis. 2019;23(7):830–7. https://doi.org/10.5588/ijtld.18.0449.
33. James R, Khim K, Boudrene L, et al. Tuberculosis active case finding in Cambodia: a pragmatic, cost-effectiveness comparison of three implementation models. BMC Infect Dis. 2017;17(1):580. https://doi.org/10.1186/s12879-017-2670-y.
34. Ji Y, Cao H, Liu Q, et al. Screening for pulmonary tuberculosis in high-risk groups of diabetic patients. Int J Infect Dis. 2020;93:84–9. https://doi.org/10.1016/j.ijid.2020.01.019.
35. Jo Y, Mirzoeva F, Chry M, et al. Standardized framework for evaluating costs of active case-finding programs: an analysis of two programs in Cambodia and Tajikistan. PLoS ONE. 2020;15(1):e0228216. https://doi.org/10.1371/journal.pone.0228216.

36. Karki B, Kettle G, Botokon I, Jr, et al. Active community-based case finding for tuberculosis with limited resources. Asia Pac J Public Health. 2017;29(1):17–27. https://doi.org/10.1177/105535916683497.

37. Kranzer K, Laxom SD, Meyer-Rath G, et al. Feasibility, yield, and cost of active tuberculosis case finding linked to a mobile HIV service in Cape Town, South Africa: a cross-sectional study. PLoS Med. 2012;9(8):e1001281. https://doi.org/10.1371/journal.pmed.1001281.

38. Machekeza SM, Wilkinson E, Hinderaker SG, et al. A comparison of the yield and relative cost of active tuberculosis case-finding algorithms in Zimbabwe. Public Health Action. 2019;9(2):63–8. https://doi.org/10.5588/pha.18.00098.

39. Maheswaran H, Barton P. Intensive case finding and isoniazid preventative therapy in HIV infected individuals in Africa: economic model and value of information analysis. PLoS ONE. 2012;7(1):e30457. https://doi.org/10.1371/journal.pone.0030457.

40. Murray M, Cattamanchi A, Denkinger C, et al. Cost-effectiveness of triage testing for facility-based systematic screening of tuberculosis among Ugandan adults. BMJ Glob Health. 2016;1(4):e000064. https://doi.org/10.1136/bmjgh-2016-000064.

41. Orlando S, Truboli I, Cicciacci F, et al. Delayed diagnosis and treatment of tuberculosis in HIV+ patients in Mozambique: a cost-effectiveness analysis of screening protocols based on four symptom screening, smear microscopy, urine LAM test and Xpert MTB/RIF. PLoS ONE. 2018;13(7):e0200523. https://doi.org/10.1371/journal.pone.0200523.

42. Reddy KP, Gupta-Wright A, Felding KL, et al. Cost-effectiveness of urine-based tuberculosis screening in hospitalised patients with HIV in Africa: a microsimulation modelling study. Lancet Glob Health. 2019;7(2):e200–8. https://doi.org/10.1016/S2214-109X(18)30436-4.

43. Sekandi JN, Dobbin K, Oloya J, et al. Cost-effectiveness analysis of community active case finding and household contact investigation for tuberculosis case detection in urban Africa. PLoS ONE. 2015;10(2):e0117009. https://doi.org/10.1371/journal.pone.0117009.

44. Shah NS, Semitile FC, Asege L, et al. Yield and efficiency of novel intensified tuberculosis case-finding algorithms for people living with HIV. Am J Respir Crit Care Med. 2019;199(5):643–50. https://doi.org/10.1164/rccm.201803-0490OC.

45. Shah S, Demissie M, Lambert L, et al. Intensified tuberculosis case finding among HIV-Infected persons from a voluntary counseling and testing center in Addis Ababa. Ethiopia J Acquir Immune Defic Syndr. 2009;50(5):537–45. https://doi.org/10.1097/QAI.0b013e3181968349.

46. Shah L, Rojas M, Mori O, et al. Cost-effectiveness of active case-finding of household contacts of pulmonary tuberculosis patients in a low HIV, tuberculosis-endemic urban area of Lima. Peru Epidemiol Infect. 2017;145(6):1107–17. https://doi.org/10.1017/S0950268816003186.

47. Smit GS, Apers L, Arrazola de Oate W, et al. Cost-effectiveness of screening for active cases of tuberculosis in Flanders, Belgium. Bull World Health Organ. 2017;95(1):27–35. https://doi.org/10.2471/BLT.16.169583.

48. Winetsy DE, Negoescu DM, DeMarchis EH, et al. Screening and rapid molecular diagnosis of tuberculosis in prisons in Russia and Eastern Europe: a cost-effectiveness analysis. PLoS Med. 2012;9(11):e1001348. https://doi.org/10.1371/journal.pmed.1001348.

49. Yoon C, Semitile FC, Asege L, et al. Yield and efficiency of novel intensified tuberculosis case-finding algorithms for people living with HIV. Am J Respir Crit Care Med. 2019;199(5):643–50. https://doi.org/10.1164/rccm.201803-0490OC.

50. Yoon C, Semitile FC, Asege L, et al. Yield and efficiency of novel intensified tuberculosis case-finding algorithms for people living with HIV. Am J Respir Crit Care Med. 2019;199(5):643–50. https://doi.org/10.1164/rccm.201803-0490OC.

51. Zishiri V, Charalambous S, Shah MR, et al. Implementing a large-scale systematic tuberculosis screening program in correctional facilities in South Africa. Open Forum Infect Dis. 2015;2(1):ofu121. https://doi.org/10.1093/ofid/ofu121.

52. Zwerling AA, Sahu M, Ngwira LG, et al. Screening for tuberculosis among adults newly diagnosed with HIV in sub-saharan Africa: a cost-effectiveness analysis. J Acquir Immune Defic Syndr. 2015;70(1):83–90. https://doi.org/10.1097/QAI.0000000000001072.

53. Adelman MW, McFarland DA, Tsegaye M, et al. Cost-effectiveness of WHO-recommended algorithms for TB case finding at Ethiopian HIV-clinics. Open Forum Infect Dis. 2018;5(1):ofx269. https://doi.org/10.1093/ofid/ofx269.

54. Soin H, Vyas A, Puri L, et al. Costs and operation management of community outreach program for tuberculosis in tribal populations in India. Public Health Action. 2019;9(2):58–62. https://doi.org/10.5588/pha.18.0091.

55. Systematic Screening for Active Tuberculosis: Principles and Recommendations. Geneva2013.

56. Ozahulek J, Claxton K, Lomas J, et al. Valuing health outcomes: developing better defaults based on health opportunity costs. Expert Rev Pharmacoecon Outcomes Res. 2020. https://doi.org/10.1080/14737167.2020.1812387.

57. Mandrik OL, Severens JLH, Bardach A, et al. Critical appraisal of systematic reviews with costs and cost-effectiveness outcomes: an ISPOR good practices task force report. Value Health. 2021;24(4):463–72. https://doi.org/10.1016/j.jval.2021.01.002.

58. Cilloni L, Fu H, Vesga JF, et al. The potential impact of the COVID-19 pandemic on the tuberculosis epidemic: a modelling analysis. EClinical Medicine. 2020;28:100603. https://doi.org/10.1016/j.eclinm.2020.100603.

59. Sahu S, Ditiu L, Sachdeva KS, et al. Recovering from the impact of the COVID-19 pandemic and accelerating to achieving the United Nations general assembly tuberculosis targets. Int J Infect Dis. 2021. https://doi.org/10.1016/j.ijid.2021.02.078[publishedOnlineFirst:2021/03/16].

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