Cost-effectiveness and budget impact of decentralising childhood tuberculosis diagnosis: a mathematical modelling study in six high tuberculosis incidence countries

Marc d'Elbée (PhD)^{1,2}, Martin Harker (MSc)³, Nyashadzaishe Mafirakureva (PhD)⁴, Mastula Nanfuka (MD)⁵, Minh Huyen Ton Nu Nguyet (MPH)¹, Jean-Voisin Taguebue (MD)⁶, Raoul Moh (MD)^{7,8}, Celso Khosa (MD)⁹, Ayeshatu Mustapha (MD)¹⁰, Juliet Mwanga-Amumpere (MD)¹¹, Laurence Borand (PharmD)^{12,13}, Sylvie Kwedi Nolna (PhD)¹⁴, Eric Komena (MD)⁸, Saniata Cumbe (MD)⁹, Jacob Mugisha (MD)¹⁵, Naome Natukunda (MPH)¹¹, Tan Eang Mao (MD)¹⁶, Jérôme Wittwer (PhD)¹⁷, Antoine Bénard (MD)¹⁸, Tanguy Bernard (PhD)¹⁹, Hojoon Sohn (PhD)²⁰, Maryline Bonnet (MD)¹⁴, Eric Wobudeya (MD)⁵, Olivier Marcy (MD)¹, Peter J. Dodd (PhD)⁴ for the TB-Speed Health Economics Study Group

Affiliations

¹University of Bordeaux, National Institute for Health and Medical Research (Inserm) UMR 1219, Research Institute for Sustainable Development (IRD) EMR 271, Bordeaux, France

²Ceped UMR 196, Université Paris Cité, Research Institute for Sustainable Development (IRD), Inserm, Paris, France

³TB Modelling Group, TB Centre, and Global Centre for Health Economics, London School of Hygiene and Tropical Medicine, London, United Kingdom

⁴School of Health & Related Research, University of Sheffield, Sheffield, United Kingdom

⁵MU-JHU Care Ltd, MUJHU Research Collaboration, Kampala, Uganda

⁶Mother and Child Center, Chantal Biya Foundation, Yaoundé, Cameroon

⁷Teaching Unit of Dermatology and Infectiology, UFR of Medical Sciences, Félix-Houphouët Boigny University, Abidjan, Côte d'Ivoire

⁸Programme PAC-CI, CHU de Treichville, Abidjan, Côte d'Ivoire

⁹Instituto Nacional de Saúde, Marracuene, Mozambique

¹⁰Ola During Children's Hospital, Freetown, Sierra Leone

¹²Epidemiology and Public Health Unit, Clinical Research Group, Institut Pasteur du Cambodge, Phnom Penh, Cambodia

¹³Center for Tuberculosis Research, Division of Infectious Diseases, Johns Hopkins University School of Medicine, Baltimore, MD USA

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¹¹Epicentre Mbarara Research Centre, Mbarara, Uganda

¹⁵SOLTHIS, Sierra Leone

¹⁶Ministry of Health, Cambodia

¹⁷University of Bordeaux, National Institute for Health and Medical Research UMR 1219,

Bordeaux, France

¹⁸CHU Bordeaux, Service d'information Médicale, USMR & CIC-EC 14-01, Bordeaux, France

¹⁹Bordeaux School of Economics, Bordeaux, France

²⁰Seoul National University College of Medicine, Seoul, South Korea

Corresponding author:

Dr Marc d'Elbée

University of Bordeaux, National Institute for Health and Medical Research (Inserm) UMR 1219, Research Institute for Sustainable Development (IRD) EMR 271, Bordeaux, France

Ceped UMR 196, Université Paris Cité, Research Institute for Sustainable Development (IRD), Inserm, Paris, France

Email: marc.delbee@u-bordeaux.fr

Phone number: +33 (0)6 71 86 01 98

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Abstract (230/250 words max)

Background

The burden of childhood tuberculosis remains high globally, largely due to under-diagnosis.

Decentralising childhood tuberculosis diagnosis services to lower health system levels could

improve case detection, but there is little empirically based evidence on cost-effectiveness or

budget impact.

Methods

We assessed the cost-effectiveness and budget impact of decentralising a comprehensive

diagnosis package for childhood tuberculosis to district hospitals (DH-focused) or primary

health centres (PHC-focused) compared to standard of care (SOC) in Cambodia, Cameroon,

Côte d'Ivoire, Mozambique, Sierra Leone, and Uganda (NCT04038632). A mathematical

model was developed to assess the health and economic outcomes of the intervention from a

health system perspective. Estimated outcomes were tuberculosis cases, deaths, disability-

adjusted life years and incremental cost-effectiveness ratios (ICERs). We also calculated the

budget impact of nationwide implementation.

Findings

For the DH-focused strategy versus SOC, ICERs ranged between \$263 (Cambodia) and \$342

(Côte d'Ivoire) per disability-adjusted life-year (DALY) averted. For the PHC-focused strategy

versus SOC, ICERs ranged between \$477 (Cambodia) and \$599 (Côte d'Ivoire) per DALY

averted. Results were sensitive to TB prevalence and the discount rate used. The additional

costs of implementing the DH-focused strategy ranged between \$13M (Cambodia) and \$50M

(Mozambique), and between \$14M (Sierra Leone) and \$135M (Uganda) for the PHC-focused

strategy.

Interpretation

The DH-focused strategy may be cost-effective in some countries, depending on the cost-

effectiveness threshold used for policy making. Either intervention would require substantial

3

early investment.

Funding

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Research in context

Evidence before the study

We searched the PubMed database using (tuberculosis) AND ((paediatric) OR (pediatric) OR (child*)) AND ((Costs and Cost Analysis) OR ("budget impact analysis")) between January 2000 and February 2023, without language restrictions.

We found 16 articles assessing the cost-effectiveness of a broad range of interventions for child tuberculosis such as vaccine programs, the use of GeneXpert machines, systematic screening, directly observed therapy, short-course preventive therapy, addition of sampling (stool) method to national algorithms, isoniazid preventive therapy delivery, and contact tracing. We did not find any budget impact analysis on tuberculosis care intervention for children. A study published by Thompson et al in 2023 investigated the costs and cost-effectiveness of a decentralised molecular testing strategy for adult tuberculosis in Uganda. It did not include chest radiography as part of the diagnostic strategy. Overall, the authors recommended decentralised Xpert testing, however, the study addressed adult patients, and no budget impact analysis was conducted to inform scale-up.

In 2022, WHO recommended decentralised models of care to deliver tuberculosis services to children. The WHO guideline development group rated the overall certainty of evidence found through systematic review as "very low", with no evidence on cost-effectiveness, and recognised that adequate investment was critical to enable the acceptability, equity and feasibility of decentralised approaches. To date, the TB-Speed Decentralization study is the closest assessment of the intervention recommended by the WHO.

Added value of this study

Unlike all of the published studies, our intervention investigated a decentralised comprehensive package of care for children as developed by the TB-Speed project which includes systematic tuberculosis screening for all sick children <15 years, clinical evaluation, Xpert MTB/RIF Ultra-testing on respiratory and stool samples, and chest radiography for children with presumptive tuberculosis.

This study found that, compared to the standard of care, decentralising a comprehensive diagnosis package for childhood tuberculosis to district hospitals is potentially cost-effective from a health systems perspective while decentralising to primary health centres is unlikely to

4

be cost-effective. Decentralisation would require substantial financial investment in the early implementation phase for equipment purchases.

Implications of all the available evidence

Decentralisation of tuberculosis diagnostic services could be cost-effective in some settings with high prevalence of tuberculosis in children seeking healthcare. The main factors affecting cost-effectiveness are the level of decentralisation (district hospital versus primary health centre), local tuberculosis prevalence, and facility testing volumes. Substantial financial commitment is needed in the early implementation phase. Following WHO recommendations, countries should consider scaling up locally-adapted interventions to improve diagnosis of tuberculosis and other diseases using Xpert machines to improve cost-effectiveness, prioritising areas with highest tuberculosis prevalence, and include such plans when identifying domestic or donor sources of funding.

Figures & tables /5 total

Figure 1. Simplified patient care pathways for the diagnosis and treatment of tuberculosis in children

Figure 2. Screening to diagnosis cascade during the intervention period by decentralisation approach

Figure 3. Cost-effectiveness acceptability curves by country for the district hospital-focused and primary health centre-focused strategies, each compared to the standard of care (in US\$ per DALY averted). DALY: disability-adjusted life year

Table 1. Health impact, costs and cost-effectiveness

Table 2. Projected budget impact of adopting the decentralisation intervention by strategy and by country for years 2022 to 2026

5

Introduction

Tuberculosis mortality remains high in children globally, with 209,000 deaths estimated by the World Health Organization (WHO) for 1.1 million paediatric cases in 2021. Modelling suggests the majority (96%) of these deaths are occurring among children not receiving treatment for tuberculosis.³ In 2021, only 38.5% of childhood tuberculosis cases were reported to the WHO, largely because of substantial underdiagnosis which prevents children from receiving treatment.

Diagnosing tuberculosis in children is challenging largely due to difficulties in collecting expectorated sputum samples and insufficient bacteria in samples to test positive due to the paucibacillary nature of pulmonary tuberculosis in children. ⁴⁻⁶ Alternative specimen collection methods such as induced sputum and gastric aspirate require equipment and experienced personnel, which are often lacking at primary health centre (PHC) level where most sick children seek care, and may be unavailable at district hospital (DH) level. Consequently, most children with presumptive tuberculosis (symptoms suggestive of tuberculosis) do not access appropriate diagnostic tests for tuberculosis, even when seen at DH level. Referrals introduce potential delays, risk losses to follow-up, and do not align with the ambition of providing patient-centred tuberculosis care.8

Developments in molecular diagnostics and novel sample collection procedures provide opportunities to decentralise diagnostic capacity for paediatric tuberculosis to the primary care level. While more costly than smear microscopy, the higher sensitivity, robustness and low training requirements of the WHO-endorsed GeneXpert-operated rapid molecular diagnostic assays for tuberculosis⁹ allow their deployment at PHC level. Stool samples can be collected in young children regardless of setting or equipment compared to respiratory samples and can be used to identify Mycobacterium tuberculosis using Xpert MTB/RIF. 10,11 Nasopharyngeal aspirates (NPA) are easier to collect than gastric aspirate or induced sputum sampling, and in combination with stool samples can provide similar sensitivity to Xpert MTB/RIF testing on two gastric aspirates or induced sputa. 10 The WHO has now recommended these sample collection methods with Xpert MTB/RIF Ultra, the latest generation of molecular tests, for paediatric tuberculosis diagnosis.⁹

The recent revision to the WHO guidelines for child and adolescent tuberculosis recommended decentralised models of care to notably increase case detection in children. However, this recommendation was provisional due to the low quality of evidence, including evidence on

cost-effectiveness, for such approaches. Few published studies evaluate models of care for paediatric tuberculosis, and even fewer include economic evaluation; none to our knowledge has included a budget impact analysis. For many countries with limited resources and high tuberculosis incidence, objectively weighing trade-offs between policy options and considering their affordability is crucial, and is explicitly required when applying for donor support including Global Fund.

The TB-Speed Decentralization study (NCT04038632) used a pre-/post-intervention approach to assess the impact on tuberculosis case detection of decentralising a comprehensive childhood tuberculosis diagnosis package at DH or PHC level. 12 The package included systematic outpatient screening, Xpert MTB/RIF Ultra on stool and NPA samples for those with presumptive tuberculosis, enhanced healthcare worker training and clinical mentoring to improve clinical skills, and digital chest X-ray at DH level. The study was conducted in 12 DHs and 47 PHCs across six high tuberculosis incidence countries: Cambodia, Cameroon, Côte d'Ivoire, Mozambique, Sierra Leone, and Uganda. In each country, two rural or semi-urban districts were randomly allocated to either a PHC-focused or DH-focused decentralisation strategy. In the DH-focused strategy, children with presumptive tuberculosis at PHC level were referred to DH for diagnosis; in the PHC-focused strategy, diagnostic evaluations were performed at PHC, except chest X-ray done at DH if required. Baseline 'pre' data comprised 9 months of retrospective data and at least 3 months of prospective data. Comparator 'post' data were collected for 12 months after introducing the intervention.

In this analysis we sought to fill the evidence gaps on the economic evaluation of real-world strategies for improving paediatric tuberculosis care by undertaking a cost-effectiveness and budget impact analysis of the TB-Speed Decentralization study¹². We used empirical data on cascades of tuberculosis care (screening, costs the clinical assessment, microbiological/radiological testing, and treatment) from the study with mathematical models of patient pathways to compare the health and cost consequences of the different models of care.

Methods

Patient pathways

Conceptual models were developed through iterative consultation with country experts and the TB-Speed Decentralization study team to represent detailed patient care pathways for three comparator arms: a standard of care (SOC), the DH-focused strategy and the PHC-focused strategy (**Figure 1** and **Appendix**). Diagrammatic representations of patient pathways formed the basis of the decision-analytic model structure; accompanying narrative descriptions of activities at each stage informed quantification of resource use. We sought to develop pathways that were general enough to include common elements across all included countries. The SOC pathway represented expert consensus on typical care available in high tuberculosis incidence countries, informed by the baseline facility assessment from the included countries before the start of the TB-Speed Decentralization intervention. DH-focused and PHC-focused pathways reflected the TB-Speed Decentralization protocol, to represent the expected patient pathway if these interventions were implemented widely across the target countries.

Costing approach

Cost data collection

Cost data collection tools were adapted from the Value TB costing tool suite developed in conjunction with the Global Health Cost Consortium (GHCC), with reference to the GHCC/WHO guidance 'Costing Guidelines for Tuberculosis Interventions'. ¹³

Labour costs were sourced from national pay scales and project accounts, medications from the Stop TB partnership Global Drug Facility catalogue, ¹⁴ consumables including materials for diagnostic tests, staff training, and equipment from project accounts, and hospitalisation cost of an inpatient bed day from the WHO-CHOICE unit cost estimates for service delivery. ¹⁵ As this intervention did not change facility infrastructure, we excluded facility-associated overhead costs from the analysis. More information on the scope of the costing can be found in **Appendix**.

Routine health facility-level aggregated data on patient numbers were collected by field research assistants from outpatient, tuberculosis, and laboratory registers during the observation, preparatory and intervention phases.

8

To calculate the proportionate use of major equipment such as X-ray or GeneXpert machines we used the expected lifespan and annual number of uses of each item. These data were obtained through key informants at MSF-Logistique (https://www.msflogistique.org/) for expected lifespan and from laboratory managers for number of uses over the course of 12 months at PHCs and at DHs.

Time and motion study

To value the contribution of labour we conducted a time and motion study to estimate the length of time that staff spent on each patient care task under TB-Speed interventions. Timesheets were used to record the length of tuberculosis-related consultations, appointments and diagnostic tests. Healthcare workers were invited to take part voluntarily and no personal data were collected, only the site name and the role of the healthcare worker. On the self-completed timesheets, participants recorded all activities related to TB-Speed patients and the length of their time they spent on each activity. Participants included doctors, clinical officers, nurses, laboratory technicians and radiographers, at DH and PHC levels in all countries, with 179 participants in total.

Cost analysis

Unit costs were estimated using an ingredient-based costing approach, in which the expected resources required for each child enrolled into the study were listed, costed, and summed to estimate direct health service utilisation and cost varying by patient characteristics and the route taken on the pathway. Services (number and type of diagnostic and treatment procedures, materials, laboratory investigations and medications) provided to patients were valued by multiplying the quantities required by their unit costs. The value of all time spent by staff for each patient was estimated as the product of 'hours spent' and 'hourly labour costs'. Costs were estimated in 2021 US dollars (USD), using a discount rate of 3% for the annualisation of the economic costs of equipment following guidelines. 16,17

Modelling approach

A decision analytic mathematical model was developed in R software to assess the clinical benefits, cost-effectiveness, and budget impact of the intervention from a health system perspective. The attributes of children flowing through the tree were: age (0-4 years or 5-14 years); HIV and antiretroviral treatment status (each positive or negative); and true tuberculosis status (bacteriologically-confirmed tuberculosis, bacteriologically-unconfirmed tuberculosis,

9

not tuberculosis). The attribute 'bacteriologically confirmed tuberculosis' refers to tuberculosis that would be bacteriologically positive under ideal circumstances and with all samples available.

The probabilities of following each route through the tree depended on attributes, and were parameterized using a mixture of literature, TB-Speed Decentralization study data, and expert opinion. Literature was used to parametrize the accuracy of diagnostic algorithms and availability of samples. Study data on child characteristics, level of presentation, and the care cascade were used to calibrate tuberculosis prevalence, the level of initial care seeking (given true tuberculosis status), and the likelihoods of being assessed and being considered to have presumptive tuberculosis. Expert opinion was used to inform unobserved features of typical care such as referral loss to follow-up and the frequency with which certain diagnostic tests were used under standard of care. Modelling of the cascade of care and intervention effect was based on calibration of unobserved parameters to meta-analytic summaries, representing a typical cascade. We undertook sensitivity analyses exploring the influence of tuberculosis prevalence, assumed constant across countries in the model, and of the discount rate applied. See **Appendix** for more details.

Country-specific unit costs associated with resource use at each step of care were accumulated to produce total mean costs. Health benefits for those with tuberculosis were modelled with a previously published approach that used case-fatality ratio from systematic literature reviews to quantify mortality reductions from treating more tuberculosis.³ Country-specific life expectancy from United Nations estimates was used to calculate the mean life-years lost over a lifetime horizon (with and without 3% discounting). We disregarded the contribution of morbidity to disability adjusted life-years (DALYs). All results were calculated using a probabilistic sensitivity analysis with 1,000 replicates.

We report the total and incremental (to SOC) number of children treated for tuberculosis, costs, number of deaths and deaths averted, number of DALYs and DALYs averted, per 100,000 children presenting as outpatients, and the incremental cost-effectiveness ratios (ICERs) for both interventions in each country. ICERs were compared to various options for costeffectiveness thresholds (presented as a range) in each country to assess potential costeffectiveness. 18,19 We complied with the Consolidated Health Economic Evaluation Reporting Standards (CHEERS 2022) reporting guidelines (**Appendix**).²⁰

To project the 5-year (2022-2026) budget impact of adopting these interventions nationally in each country, we fitted a cost function in Excel using parameters related to costs of equipment and its delivery, installation and maintenance, training, supplies, and personnel costs. Input quantities were the national number of DHs, PHCs, eligible healthcare workers, and preintervention (2019) annual rate of tuberculosis notification in children 0-14 years old from the WHO Global TB Programme. See Appendix for additional details. Costs were combined with the scale of deployment to generate the required implementation budget in each year for each country. We assumed a 3-year period (2022-2024) to provide facilities with equipment, to train all eligible staff, and rollout supply chain delivery to all sites. A sensitivity analysis is presented in **Appendix**.

A health economic analysis plan was developed jointly by economists, modellers, international and country investigators. The economic analyses were nested within the main study protocol approved by the WHO Ethical Review Committee, Inserm's ethics review committee (IRB00003888), as well as all national ethics committees prior to the start of the study (See Appendix).

Role of the funding source

The study funders had no role in the study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

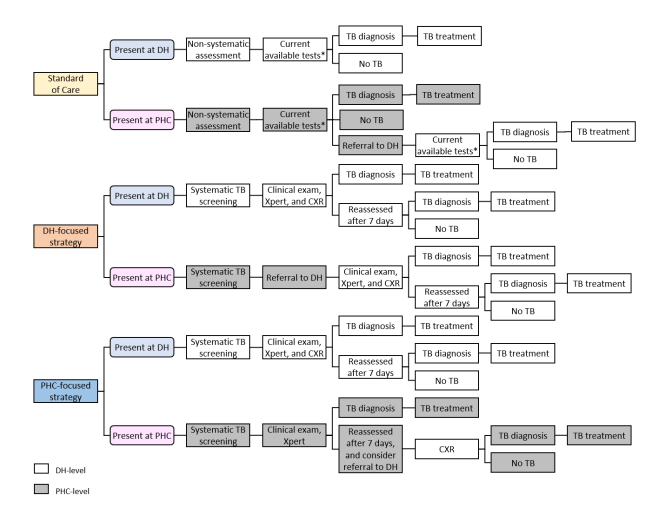


Figure 1. Simplified patient care pathways for the diagnosis and treatment of tuberculosis in children *Clinical exam, and, for a proportion of patients, Xpert on sputum or gastric aspirate, smear microscopy and CXR (in DH) TB: tuberculosis, DH: district hospital, PHC: primary health centre, CXR: chest X-ray

Results

Overall, the modelled cascade of care data showed that the tuberculosis screening rate among children presenting to health facilities increased from 21% (SOC) to 89% (DH-focused and PHC-focused), the rate of children identified with presumptive tuberculosis among screened children remained constant (3%), and the rate of children initiated on treatment among the children identified with presumptive tuberculosis increased from 12% (SOC) to 18% (DHfocused) and 14% (PHC-focused) (Figure 2). Results were similar between 0-4 and 5-14 years age groups (Appendix).

Results from the cost analysis and the time and motion study, as well as costs per child treated disaggregated by arm, cost category, and tuberculosis care stage are presented in the **Appendix**. Compared to 74 (95% uncertainty interval [UI]: 6 to 257) children treated for tuberculosis in the SOC (per 100,000 presenting), we estimated 476 (95% UI: 104 to 1,210) treated in the DHfocused arm and 373 (95% UI: 80 to 925) treated in the PHC-focused arm (Table 1). Compared with the SOC at 44 (6 to 131) deaths per 100,000 presenting, the number of deaths averted was 21 (2 to 69) and 18 (2 to 57), for the DH-focused and PHC-focused interventions, respectively. For the DH-focused intervention costs per 100,000 increased by between \$158K (\$41-357K) in Cambodia and \$190K (\$54-416K) in Côte d'Ivoire, while DALYs averted ranged from 547 (54-1,754) in Sierra Leone to 600 (59-1,916) in Cambodia.

The ICERs compared with the SOC ranged between \$263 (Cambodia) and \$342 (Côte d'Ivoire) per DALY averted for the DH-focused strategy, and between \$477 (Cambodia) and \$599 (Côte d'Ivoire) per DALY averted for the PHC-focused strategy. The DH-focused strategy dominated the PHC-focused strategy, being less expensive and more effective. Sensitivity analysis showed that both paediatric tuberculosis prevalence and discount rate applied on life years had a strong impact on the estimated ICERs (**Appendix**). At a constant TB prevalence of 200/100,000 across age groups, applying no discount rate (0% instead of 3%) led to ICERs decreasing by 57%-60% across countries and strategies. Increasing TB prevalence from 200/100,000 to 500/100,000 (with fixed 3% discounting) led to ICERs decreasing by 55%-56%.

The cost-effectiveness acceptability curves (Figure 3 and Appendix) show the decision uncertainty surrounding the adoption of the strategies, depending on the cost-effectiveness thresholds which decision-makers may select in each country. Using estimated thresholds from Ochalek et al. 18, the highest estimated probabilities of the DH-focused strategy being cost-

effective compared to SOC were for Cambodia (40%-69%) and Côte d'Ivoire (49%-65%) and the lowest for Sierra Leone (0%-1%). For the PHC-focused strategy versus SOC, the highest probabilities were for Cambodia (7%-27%) and Côte d'Ivoire (11%-26%) and the lowest for Cameroon, Sierra Leone and Uganda (<1%).

Additional costs of implementing the decentralisation intervention at DH level over five years were estimated between \$13M in Cambodia and Sierra Leone and \$50M in Mozambique, whereas decentralising to PHC level would cost between \$14M in Sierra Leone and \$135M in Uganda (Table 2). Countries with higher numbers of health facilities to equip (Cambodia, Mozambique, Sierra Leone) and/or higher tuberculosis notification rates (Cambodia, Mozambique, Sierra Leone, Uganda) had higher implementation costs.

The average annual cost of a five-year scale-up plan of the DH-focused intervention ranged between \$3M (Cambodia) and \$10M (Mozambique), corresponding to between 8% (Cambodia) and 54% (Cameroon) of countries' national tuberculosis programme (NTP) budgets in 2021, and, for the PHC-focused intervention, between \$3M (Sierra Leone) and \$27M (Uganda), or 18% (Cambodia) to 164% (Cameroon) of NTP budget. Once the health facilities are equipped, the average annual routine costs (corresponding to year 2026) ranged between \$1.3M (3% NTP budget) in Côte d'Ivoire and \$13.7M (46%) in Mozambique for DH-focused, and between \$0.4M (1%) in Côte d'Ivoire and \$5.1M (17%) in Mozambique for the PHCfocused strategy.

We assumed facilities saw a sustained increase in diagnoses following implementation. We projected an increase in the cumulative number treated for tuberculosis over the five-year implementation period of 194% for the DH-focused, and 24% for the PHC-focused strategy, compared to SOC. Sensitivity analyses suggested that important cost reductions could be achieved through purchase price reduction of the cartridge kits and mucus aspirators, and economies of scale on delivery chain costs (Appendix).

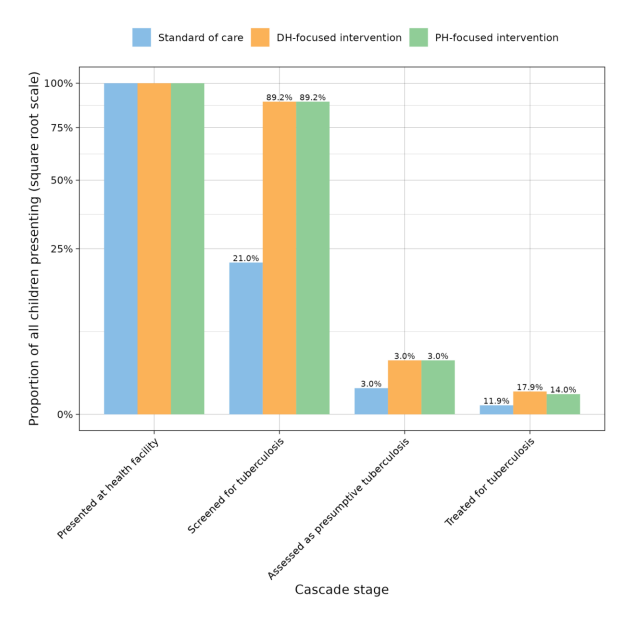


Figure 2. Screening to diagnosis cascades during the intervention period by decentralisation approach. Percentages are bar height as a proportion of the corresponding bar at the previous stage.

 Table 1. Health impact, costs and cost-effectiveness

Arm	Type of outcome	Per 100,000 OPD initial attendances	Cambodia	Cameroon	Côte d'Ivoire	Mozambique	Sierra Leone	Uganda
		Children treated for TB* (95% UI)	74 (6 to 257)					
		Deaths (95% UI)	44 (6 to 131)					
SOC	Total	Costs (95% UI)	10,449 (1,164 to 32,054)	13,430 (1,453 to 41,195)	15,574 (1,797 to 46,247)	10,359 (1,073 to 33,665)	10,164 (1,100 to 32,066)	13,581 (1,575 to 40,263)
		DALYs§ (95% UI)	1,235 (177 to 3,680)	1,162 (166 to 3,461)	1,145 (163 to 3,410)	1,192 (170 to 3,553)	1,126 (160 to 3,354)	1,200 (171 to 3,576)
		Children treated for	476 (104 to					
		TB* (95% UI)	1,210)	1,210)	1,210)	1,210)	1,210)	1,210)
DH- focused	Total	Deaths (95% UI)	23 (3 to 75)					
Tocuseu			168,490	198,455	205,544	181,355	176,682	193,742
		Costs (95% UI)	(48,731 to	(57,910 to	(64,514 to	(49,499 to	(46,093 to	(59,315 to
			375,050)	435,159)	440,572)	409,446)	398,673)	415,612)

		DALYs§ (95% UI)	635 (87 to 2,100)	597 (82 to 1,971)	588 (81 to 1,943)	613 (84 to 2,025)	579 (79 to 1,909)	617 (84 to 2,039)
		Children treated for TB* (95% UI)	402 (71 to 1,032)					
		Deaths averted (95% UI)	21 (2 to 69)					
	Incremental to SOC	Costs (95% UI)	158,040 (40,967 to 357,875)	185,025 (50,236 to 415,776)	189,970 (53,578 to 415,937)	170,996 (41,951 to 389,530)	166,518 (38,986 to 376,362)	180,161 (50,988 to 395,362)
		DALYs§ averted (95% UI)	600 (59 to 1,916)	565 (56 to 1,809)	556 (55 to 1,780)	579 (57 to 1,853)	547 (54 to 1,754)	583 (58 to 1,865)
		ICER¤	263	328	342	295	304	309
PHC- focused	Total	Children treated for TB* (95% UI)	373 (80 to 925)					
Tocuseu		Deaths (95% UI)	26 (4 to 79)					

	Costs (95% UI)	257,022 (71,606 to 583,728)	295,105 (84,794 to 661,929)	303,124 (92,655 to 677,697)	276,615 (73,237 to 639,476)	273,622 (72,754 to 628,440)	286,493 (82,536 to 644,522)
	DALYs§ (95% UI)	717 (102 to 2,212)	675 (97 to 2,077)	665 (95 to 2,047)	693 (99 to 2,133)	654 (94 to 2,012)	697 (100 to 2,147)
	Children treated for TB* (95% UI)	298 (52 to 760)					
	Deaths averted (95% UI)	18 (2 to 57)					
Incremental to SOC	Costs (95% UI)	246,572 (64,623 to 564,187)	281,675 (75,305 to 640,442)	287,550 (81,529 to 637,224)	266,256 (66,963 to 617,904)	263,458 (69,220 to 612,906)	272,912 (75,041 to 616,087)
	DALYs§ averted (95% UI)	517 (52 to 1,608)	487 (49 to 1,516)	480 (48 to 1,493)	500 (50 to 1,554)	472 (47 to 1,470)	503 (50 to 1,564)
	ICER¤	477	578	599	533	558	543

SOC: standard of care, TB: tuberculosis, OPD: patients presenting at outpatient department, DH: district hospital, PHC: primary health centre, DALY: disability-adjusted life year, ICER: incremental cost-effectiveness ratio, UI: uncertainty interval

*We used an average intervention impact, so the number of children treated for tuberculosis by intervention is the same across countries. Costs and life expectancy tables for estimating DALYs are specific to each country.

§Discounted

¤The discrepancy observed between manually calculated and presented ICERs is due to rounding error.

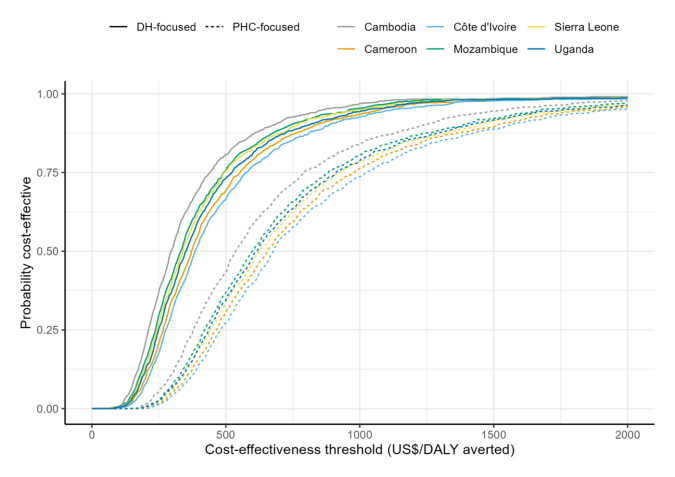


Figure 3. Cost-effectiveness acceptability curves by country for the district hospital-focused and primary health centre-focused strategies, each compared to the standard of care (in US\$ per DALY averted). DALY: disability-adjusted life year

Table 2. Projected budget impact of adopting the decentralisation intervention by strategy and by country for years 2022 to 2026. Costs are in 2021 US\$

	Cambodia	Cameroon	Côte d'Ivoire	Mozambique	Sierra Leone	Uganda			
Country indicators in 2021	Country indicators in 2021								
Population (0-14 years), 2021	5,253,000	11,434,000	11,092,000	14,152,000	3,257,000	21,677,000			
Annual number of OPD attendances by children 0-14 years, 2020	2,177,000	1,478,000	1,199,000	15,138,000	2,767,000	9,676,000			
Annual number of tuberculosis notification for children 0-14 years, 2020	1,849	1,255	1,018	12,856	2,350	8,218			
District hospitals - DH (National)	68	165	150	53	49	163			
Primary Health Centres - PHC (National)	1,141	2,214	3,411	1,566	242	5,155			
Ratio PHC/DH	17	13	23	30	5	32			
National tuberculosis programme budget, 2021 (millions US\$)	34	8	46	30	10	45			
Impact: number of 0-14 year old children	treated for tuberculo	sis over the 5-year in	nplementation period	l					
Standard of care	9,245	6,275	5,090	64,280	11,750	41,090			
DH-focused (% increase from the SOC)	27,217 (+194%)	18,474 (+194%)	14,985 (+194%)	189,240 (+194%)	34,592 (+194%)	120,969 (+194%)			
PHC-focused (% increase from the SOC)	11,467 (+24%)	7,783 (+24%)	6,313 (+24%)	79,728 (+24%)	14,574 (+24%)	50,965 (+24%)			

Budget req	udget requirement related to the decentralisation interventions (incremental costs)												
Interve ntion	Period	Costs (millions US\$)	% total costs										
	2022	2·8 (2·6 to 3·1)	22%	6·1 (5·9 to 6·3)	28%	5.5 (5.3 to 5.8)	28%	6·1 (4·9 to 7·9)	12%	2·4 (2·2 to 2·8)	19%	8·7 (7·8 to 9·9)	18%
	2023	3·2 (2·9 to 3·7)	25%	6·3 (6 to 6·8)	29%	5·8 (5·5 to 6·2)	30%	8·5 (6·5 to 11·8)	17%	2·8 (2·5 to 3·5)	22%	10·5 (8·9 to 13·0)	21%
DII	2024	3.5 (3.1 to 4.2)	27%	6.6 (6.2 to 7.3)	30%	6·0 (5·6 to 6·6)	31%	10·8 (8·0 to 15·7)	21%	3·2 (2·7 to 4·1)	25%	12·2 (10·0 to 16·1)	25%
DH- focused	2025	1.5 (1.0 to 2.4)	12%	1·2 (0·8 to 2·0)	6%	1·0 (0·7 to 1·7)	5%	11·3 (7·8 to 17·6)	22%	2·0 (1·4 to 3·1)	16%	7.9 (5.2 to 13.1)	16%
	2026	1·8 (1·2 to 3·0)	14%	1·5 (1·0 to 2·5)	7%	1·3 (0·8 to 2·2)	7%	13·7 (9·4 to 21·5)	27%	2·4 (1·6 to 3·8)	19%	9.6 (6.3 to 16.2)	20%
	Total 2022-2026	12·8 (10·8 to 16·4)	100%	21·7 (19·9 to 24·9)	100%	19.6 (18.0 to 22.5)	100 %	50·4 (36·5 to 74·4)	100%	12·9 (10·4 to 17·2)	100%	49·0 (38·2 to 68·5)	100 %
PHC-	2022	10·0 (9·6 to 10·4)	32%	21·4 (20·8 to 22·0)	33%	29·9 (29·1 to 30·7)	33%	16·1 (14·7 to 17·7)	27%	4·0 (3·8 to 4·3)	29%	42·4 (40·7 to 44·1)	32%
focused	2023	10·0 (9·6 to 10·5)	32%	21·5 (20·8 to 22·1)	33%	30·0 (29·2 to 30·8)	33%	16·5 (15·0 to 18·3)	28%	4·1 (3·8 to 4·4)	29%	42·7 (40·9 to 44·6)	32%

	2024	10·1 (9·7 to 10·6)	32%	21·5 (20·9 to 22·2)	33%	30·0 (29·2 to 30·9)	33%	16·9 (15·2 to 19·0)	28%	4·2 (3·9 to 4·5)	30%	43·0 (41·1 to 45·1)	32%
	2025	0.6 (0.4 to 0.9)	2%	0·5 (0·3 to 0·7)	1%	0.4 (0.3 to 0.6)	0%	4·7 (3·2 to 6·7)	8%	0.8 (0.5 to 1.1)	6%	3·1 (2·1 to 4·4)	2%
	2026	0·7 (0·4 to 1·0)	2%	0·5 (0·4 to 0·8)	1%	0.4 (0.3 to 0.6)	0%	5·1 (3·5 to 7·3)	9%	0.8 (0.6 to 1.2)	6%	3.4 (2.3 to 4.9)	3%
	Total 2022-2026	31·4 (29·7 to 33·2)	100%	65·4 (63·2 to 67·7)	100%	90·7 (88·0 to 93·6)	100 %	59·3 (51·6 to 69·0)	100%	13.9 (12.6 to 15.6)	100%	134·6 (127·1 to 143·0)	100 %
Interve ntion	Average annual costs	Costs (millions US\$)	% of NTP budge t	Costs (millions US\$)	% of NTP budg et	Costs (millions US\$)	% of NTP budg et						
DH-	Program implementation costs (average of 2022-2026)	2.6 (2.2 to 3.3)	8%	4·3 (4·0 to 5·0)	54%	3.9 (3.6 to 4.5)	9%	10·1 (7·3 to 14·9)	34%	2.6 (2.1 to 3.4)	26%	9.8 (7.6 to 13.7)	22%
focused	Routine costs (2026)	1.8 (1.2 to 3.0)	5%	1·5 (1·0 to 2·5)	19%	1·3 (0·8 to 2·2)	3%	13·7 (9·4 to 21·5)	46%	2·4 (1·6 to 3·8)	24%	9.6 (6.3 to 16.2)	21%
PHC-	Program implementation costs (average of 2022-2026)	6·3 (5·9 to 6·6)	18%	13·1 (12·6 to 13·5)	164%	18·1 (17·6 to 18·7)	39%	11·9 (10·3 to 13·8)	40%	2·8 (2·5 to 3·1)	28%	26·9 (25·4 to 28·6)	60%
focused	Routine costs (2026)	0·7 (0·4 to 1·0)	2%	0·5 (0·4 to 0·8)	6%	0.4 (0.3 to 0.6)	1%	5·1 (3·5 to 7·3)	17%	0.8 (0.6 to 1.2)	8%	3.4 (2.3 to 4.9)	8%

OPD: patients presenting at outpatient department, DH: district hospital, PHC: primary health centre, SOC: standard of care, NTP: National Tuberculosis Program

Discussion

Benchmarked against estimates of country cost-effectiveness thresholds, ¹⁸ this study found that the decentralisation of paediatric tuberculosis services to DH level could be cost-effective compared to the SOC from a health systems perspective in Cambodia and Côte d'Ivoire, whereas decentralisation to PHC level was unlikely to be cost-effective in any country. Ultimately, choice of cost-effectiveness thresholds is a judgement for decision-makers, and other considerations, notably the budget impact, also need to be considered. Sensitivity analysis suggests that decentralisation targeted to geographical areas with very high tuberculosis prevalence would be highly likely to be cost-effective in all countries. Implementation would require substantial financial investment in the early phase, particularly for the PHC-focused intervention.

The PHC-focused strategy was, as expected, more costly than the DH-focused strategy due to spending on diagnostic equipment for a larger number of facilities, but unexpectedly was also less effective than the DH-focused strategy. The reasons for this lower effectiveness (fewer tuberculosis diagnoses) are not clear, but may include differences in staff cadres composition and greater experience in diagnosing paediatric tuberculosis at DH level due to higher patient volumes. The higher tuberculosis diagnosis rate in children seeking care at DH level may indicate care-givers prefer to take more severely ill children to this level first or because children are referred from PHC with more advanced disease. We did not model differences in tuberculosis severity by location which may have favoured the DH-focused strategy. Finally, all children enrolled in DH had a CXR performed while in PHCs, only children with persisting symptoms after 7 days were referred for CXR at DH; this is likely to have contributed to higher tuberculosis detection in the DH-focused strategy. We did not include patient costs in our analysis; the PHC-focused strategy is likely to have reduced costs to patients due to fewer referrals.

We used a single typical SOC as a comparator across the six countries. It was not practical or transparent to represent every variation that exists within and across these countries. In our iterative conceptual modelling, we agreed on a structure clinicians from all countries felt adequately represented patient pathways in their country. However, paediatric tuberculosis care was already partially decentralised to PHC level in Uganda and Mozambique, meaning our modelled SOC is more of an approximation in these settings. Some parameters were derived by calibrating to cascade data, and our estimate of the true tuberculosis prevalence in children

seeking care across all countries was uncertain. In reality tuberculosis prevalence will vary between countries, and higher prevalence values substantially improved cost-effectiveness (Appendix).

We were also limited by the nature of the TB-Speed Decentralization study and data. The pre-/post-intervention primary outcome means results are vulnerable to confounding by factors that changed over time. In particular, the study period overlapped with the COVID-19 pandemic. Restrictions introduced by countries affected transportation (less available and more costly) and these barriers may have increased patient losses during referral to hospitals. Fear of COVID-19 also reduced facility attendance. Facility-level overheads costs were recently reported to be a significant contributor to total costs in adult tuberculosis costing estimates from the Value-TB project (the largest tuberculosis costing effort to date). 21 These were excluded in our study because it was anticipated that the adoption of a new diagnostic strategy without change to the number of health facilities or staff would not significantly affect overheads, but this assumption should be carefully considered alongside roll-out plans. Data collection, as well as our analysis, was focused on children, which ignores the additional benefits from decentralisation of tuberculosis diagnostic capacity in improving the detection of tuberculosis in adults, as well as detection of rifampicin-resistant tuberculosis and other pathogens using the GeneXpert platform.

A major strength of this study is the collection and analysis of primary data to inform costs, impacts, and care cascades in six high tuberculosis incidence countries in different regions of Africa and in South-East Asia. This diversity in economic and health system context suggests the generalisability of these results across comparable settings. Underlying modelling assumptions were based on a number of previously published studies, and applied within a framework that captured considerable complexity in patient pathways, including reassessments and referrals between levels.

To our knowledge, this is the first multi-country study to assess the cost-effectiveness or budget impact of decentralising childhood tuberculosis services, and will add to the evidence base for the interim WHO recommendation on decentralised models of care. Systematic reviews of the cost and cost-effectiveness of tuberculosis screening have found ICERs of between US\$281 and US\$698 per DALY averted in the general population, ²² but for screening in high risk groups ICERs as low as US\$51 per DALY have been reported. ²³ An analysis of the cost-effectiveness of Xpert on stool for paediatric tuberculosis in Ethiopia and Indonesia found ICERs of US\$132

and US\$94 per DALY, respectively.²⁴ However, the majority of these analyses were not based on empirical data from implemented interventions, and restricted to a small component of tuberculosis diagnosis given that the majority of children are clinically diagnosed.

A number of studies have explored the effects of decentralisation of general tuberculosis services and costs. A 2017 study assessed the cost per tuberculosis diagnosis of implementing Xpert testing regardless of age at PHCs in Uganda.²⁵ Average costs per new tuberculosis diagnosis using Xpert averaged US\$119, but were as high as US\$885 in the lowest-volume centre. The authors did not attempt to estimate cost-effectiveness in terms of cost per DALY averted. Thompson and colleagues estimated the cost-effectiveness of an adult tuberculosis diagnostic strategy decentralised in community health centres in Uganda.²⁶ The authors recommended decentralised testing services with ICERs ranging US\$687 per additional treatment initiation in 14 days and US\$1,332 per additional tuberculosis diagnosis. Costeffectiveness notably increased with high testing volumes and lower equipment costs.

TB-Speed Decentralization and other studies have demonstrated the potential for decentralised diagnostic approaches to find more children with tuberculosis. 12,27 Our analysis shows that this is possible in a way that could be considered cost-effective across a range of settings, depending on strategy and tuberculosis prevalence, but that large-scale implementation would still incur substantial costs relative to existing NTP budgets. Countries should consider scaling up locallyadapted interventions to improve tuberculosis diagnosis while monitoring their performance, potentially prioritising areas with the highest tuberculosis prevalence, and including such plans when identifying domestic or donor sources of funding.

Contributors

MH, NM, PD, OM, MB, EW conceived the health economics analysis plan. PD developed the cost-effectiveness model and MDE developed the budget impact model. MDE conducted the cost, the cost-effectiveness and the budget impact analyses. MH coordinated international economic data collection and design of the patient pathways. MDE, PD, MH, NM interpreted economic results. TB, AB, HS provided scientific expertise and guidance.

OM, MB, EW conceived and designed the TB-Speed Decentralization study. MN coordinated and contributed to the implementation of the study protocol. OM, MB, and EW led the study at international level. LB led the study in Cambodia, J-VT led the study in Cameroon, RM led the study in Côte d'Ivoire, CK led the study in Mozambique, JM led the study in Sierra Leone, EW and JM-A led the study in Uganda, MN coordinated study implementation at international level. MHTNN did the statistical analysis of the decentralisation study. OM, MN, MH, MB, EW and MDE contributed to the interpretation of the results.

MDE wrote the first draft and all authors reviewed, edited and approved the final version of the manuscript. MDE, PD, OM, MB, EW were responsible for the decision to submit the manuscript.

TB-Speed Health Economics Study Group

Name	Affiliation
Arlt-Hilares, Doris	MSF-Logistique, Merignac, France
Balestre, Eric	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Research Institute for Sustainable Development EMR 271, Bordeaux, France
Banga, Marie-France	Programme PAC-CI, CHU de Treichville, Abidjan, Côte d'Ivoire
Bénard, Antoine	CHU Bordeaux, Service d'information Médicale, USMR & CIC-EC 14-01, Bordeaux, France

Bernard, Tanguy	Bordeaux School of Economics, Bordeaux, France
Bonnet, Maryline	TransVIHMI, University of Montpellier, IRD /INSERM, Montpellier, France
Borand, Laurence	Center for Tuberculosis Research, Division of Infectious Diseases, Johns Hopkins University School of Medicine, Baltimore, MD USA Epidemiology and Public Health Unit, Clinical Research Group, Institut Pasteur du Cambodge, Phnom Penh, Cambodia
Breton, Guillaume	SOLTHIS, Paris, France
Bunnet, Dim	Epidemiology and Public Health Unit, Clinical Research Group, Institut Pasteur du Cambodge, Phnom Penh, Cambodia
Chateau, Paul-Damien	MSF-Logistique, Merignac, France
Cumbe, Saniata	Instituto Nacional de Saúde, Marracuene, Mozambique
d'Elbée, Marc	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Research Institute for Sustainable Development EMR 271, Bordeaux, France Ceped UMR 196, Université Paris Cité, Research Institute for Sustainable Development (IRD), Inserm, Paris, France
de Lauzanne, Agathe	Epidemiology and Public Health Unit, Clinical Research Group, Institut Pasteur du Cambodge, Phnom Penh, Cambodia

Dodd, Peter J.	School of Health & Related Research, University of Sheffield, Sheffield, United Kingdom
Harker, Martin	TB Modelling Group, TB Centre, and Global Centre for Health Economics, London School of Hygiene and Tropical Medicine, London, United Kingdom
Huyen Ton Nu Nguyet, Minh	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Research Institute for Sustainable Development EMR 271, Bordeaux, France
Kaing, Sanary	Epidemiology and Public Health Unit, Clinical Research Group, Institut Pasteur du Cambodge, Phnom Penh, Cambodia
Khosa, Celso	Instituto Nacional de Saúde, Marracuene, Mozambique
Komena, Eric	Programme PAC-CI, CHU de Treichville, Abidjan, Côte d'Ivoire
Koroma, Monica	SOLTHIS, Sierra Leone
Kwedi Nolna, Sylvie	University of Montpellier, TransVIHMI, IRD /INSERM, Montpellier, France
Mafirakureva, Nyashadzaishe	School of Health & Related Research, University of Sheffield, Sheffield, United Kingdom
Mao, Tan Eang	Cambodia National Tuberculosis Programme

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Marcy, Olivier	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Research Institute for Sustainable Development EMR 271, Bordeaux, France				
Mbang Masson, Douglas	Mother and Child Center, Chantal Biya Foundation, Yaoundé, Cameroon				
Moh, Raoul	Teaching Unit of Dermatology and Infectiology, UFR of Medical Sciences, Félix-Houphouët Boigny University, Abidjan, Côte d'Ivoire Programme PAC-CI, CHU de Treichville, Abidjan, Côte d'Ivoire				
Mugisha, Jacob	SOLTHIS, Sierra Leone				
Mustapha, Ayeshatu	Ola During Children's Hospital, Freetown, Sierra Leone				
Mwanga-Amumpere, Juliet	Epicentre Mbarara Research Centre, Mbarara, Uganda				
Nafuka, Mastula	MU-JHU Care Ltd, MUJHU Research Collaboration, Kampala, Uganda				
Natukunda, Naome	Epicentre Mbarara Research Centre, Mbarara, Uganda				
Orne-Gliemann, Joanna	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Research Institute for Sustainable Development EMR 271, Bordeaux, France				
Ouattara, Eric	University of Bordeaux, National Institute for Health and Medical Research UMR 1219,				

	Research Institute for Sustainable Development EMR 271, Bordeaux, France
Poublan, Julien	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Research Institute for Sustainable Development EMR 271, Bordeaux, France
Sohn, Hojoon	Seoul National University College of Medicine, Seoul, South Korea
Taguebue, Jean-Voisin	Mother and Child Center, Chantal Biya Foundation, Yaoundé, Cameroon
Tulinawe, Immaculate	Epicentre Mbarara Research Centre, Mbarara, Uganda
Voss de Lima, Yara	Instituto Nacional de Saúde, Marracuene, Mozambique
Wittwer, Jérôme	University of Bordeaux, National Institute for Health and Medical Research UMR 1219, Bordeaux, France
Wobudeya, Eric	MU-JHU Care Ltd, MUJHU Research Collaboration, Kampala, Uganda

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Data sharing: Aggregated data for all analyses will be publicly available with the publication on a Github repository under a Creative Commons Attribution (CC BY) licence (URL: https://github.com/petedodd/TBSdecent).

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