Closing gaps in the tuberculosis care cascade: an action-oriented research agenda

Ramnath Subbaraman\textsuperscript{a,b,*}, Tulip Jhaverib, Ruvandhi R. Nathavitharana\textsuperscript{c}

\textsuperscript{a} Department of Public Health and Community Medicine and Center for Global Public Health, Tufts University School of Medicine, Boston, USA
\textsuperscript{b} Division of Geographic Medicine and Infectious Diseases, Tufts Medical Center, Boston, USA
\textsuperscript{c} Division of Infectious Diseases, Beth Israel Deaconess Medical Center and Harvard Medical School, Boston, USA

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ABSTRACT

The care cascade—which evaluates outcomes across stages of patient engagement in a health system—is an important framework for assessing quality of tuberculosis (TB) care. In recent years, there has been progress in measuring care cascades in high TB burden countries; however, there are still shortcomings in our knowledge of how to reduce poor patient outcomes. In this paper, we outline a research agenda for understanding why patients fall through the cracks in the care cascade. The pathway for evidence generation will require new systematic reviews, observational cohort studies, intervention development and testing, and continuous quality improvement initiatives embedded within national TB programs. Certain gaps, such as pretreatment loss to follow-up and post-treatment disease recurrence, should be a priority given a relative paucity of high-quality research to understand and address poor outcomes. Research on interventions to reduce death and loss to follow-up during treatment should move beyond a focus on monitoring (or observation) strategies, to address patient needs including psychosocial and nutritional support. While key research questions vary for each gap, some patient populations may experience disparities across multiple stages of care and should be a priority for research, including men, individuals with a prior treatment history, and individuals with drug-resistant TB. Closing gaps in the care cascade will require investments in a bold and innovative action-oriented research agenda.

1. Introduction

The care cascade evaluates patient outcomes for a disease across stages of care. National-level care cascade analyses have identified that large numbers of individuals with active tuberculosis (TB) experience poor outcomes at critical points in health system engagement, highlighting foundational problems in quality of TB care [1,2]. We recently outlined guidelines for estimating the number of individuals with active TB in a population who successfully reach (or drop out at) different care cascade stages [3]. While such analyses help quantify gaps in care delivery, they do not illuminate why patients fall through the cracks—information that is critical for developing interventions to improve outcomes in TB programs.

Reasons for poor outcomes—and interventions to address these problems—may vary at each care cascade stage. Closing gaps in the care cascade may require interventions at the level of the population or health system (including the private sector), at the level of TB diagnostic and treatment centers, and at the level of the TB patient-health provider interaction. Rectifying gaps at different scales will require diverse interventions—potentially including large-scale public education, increased access to health facilities, initiatives in the private sector, integration of new diagnostic and monitoring technologies, and interventions to address patients’ psychosocial needs. In addition, some patients may be at higher risk for poor outcomes, thereby meriting greater attention and specialized interventions. In light of these complexities, in this manuscript, we outline an agenda to start answering key questions regarding poor patient outcomes in the TB care cascade.

2. Frameworks and research questions

2.1. Framework for the TB care cascade

We previously described a care cascade model for individuals with active TB, in which each stage contains a step (number of individuals who reach that point in care) and a gap (those with poor outcomes, quantified as the difference between steps) (Fig. 1) [3]. Key gaps

* Corresponding author at: Department of Public Health and Community Medicine, Tufts University School of Medicine, 136 Harrison Ave., MV237, Boston, MA 02111, USA.

E-mail address: ramnath.subbaraman@tufts.edu (R. Subbaraman).

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include: individuals with active TB in the population who do not reach health facilities and access a TB diagnostic test (Gap 1), those who access locations where diagnostic tests are available but do not get successfully diagnosed (Gap 2), those successfully diagnosed who do not get registered in treatment (Gap 3), those who start therapy but do not achieve treatment success (Gap 4), and those who finish therapy but experience death or TB recurrence within a year (Gap 5). We describe a research agenda to address each of these gaps below.

2.2. Research questions

The research agenda below is guided by three broad questions. First, **who is disproportionately falling out of the TB care cascade?** Understanding the types of individuals who are at higher risk for poor outcomes at each stage may help to develop and refine interventions that focus on these specific populations, although we acknowledge that quality of care can and should be improved for all people with TB. Risk of dropping out of care may vary by demographics (e.g., age, gender), type of tuberculosis (e.g., pulmonary, extrapulmonary, prior treatment history), microbiological susceptibility (e.g., drug-resistant forms of TB), comorbidities (e.g., HIV, diabetes), or other social factors (e.g., living in migrant, urban slum, or indigenous communities).

Second, **why are patients falling out of the cascade?** Understanding barriers to engaging in TB care that contribute to poor outcomes are important to inform intervention development. Such barriers may occur at the level of the health system (e.g., poor quality of care or user experience), the patient (e.g., substance use, depression), the patient's family and community (e.g., TB-related stigma), or society (e.g., structural barriers).

Third, **what interventions are needed to reduce gaps in the care cascade?** Beneficial interventions might involve using novel technologies to address health system or patient barriers, social and behavioral interventions to address psychosocial barriers, social protection schemes for patients, or incentives to change healthcare provider (HCP) behavior, including in the private health sector. Intervention development would ideally be informed by research on the first two questions described above.

2.3. Pathway for generating evidence

Diverse research approaches will be required to understand which patients are being lost, identify reasons for these losses, develop interventions, and implement these interventions in routine clinical practice (Fig. 2). Systematic reviews help to aggregate evidence about reasons for patient dropout across care cascade stages and the effectiveness of interventions to reduce these gaps. For example, systematic reviews have synthesized evidence on barriers to TB medication adherence from qualitative studies [4] and assessed the effectiveness (or lack thereof) of directly observed therapy (DOT) [5–9] and other interventions for improving adherence, including digital adherence technologies (DATs) [8,10]. Systematic reviews have not evaluated reasons or interventions for other care cascade gaps, such as pretreatment loss to follow-up (PTLFU) (Gap 3) and post-treatment relapse or death (Gap 5). Studies of HIV care delivery provide helpful examples to guide similar systematic reviews for TB [11–15].

By identifying research gaps, systematic reviews may guide further qualitative and quantitative observational research to identify novel risk factors for patient dropout. Findings of observational studies may in turn guide theory-informed intervention development to address risk factors, using iterative implementation and refinement. Implementation research frameworks—including the Unified Theory of Acceptance and Use of Technology (for technology-based interventions) [16], the RE-AIM framework, and the Consolidated Framework for Implementation Research—may guide approaches to designing, evaluating, and
implementing interventions.

Intervention testing can take a variety of approaches. Since interventions to retain patients often require health system changes, cluster-randomized trials may facilitate rigorous evaluations of such interventions. However, such resource-intensive research approaches may not always be practical. Due to poor quality of care at later care cascade stages, interventions that address one gap may improve surrogate endpoints without translating into benefits in long-term outcomes, such as TB cure or recurrence-free survival, which does not necessarily mean that the intervention is not beneficial [17]. In addition, multi-component interventions are more likely to improve long-term outcomes, but development and assessment of such interventions may be more amenable to quality improvement cycles (e.g., plan-do-study-act) and observational studies embedded in routine clinical practice, rather than randomized trials, to enable real-time iterative improvements [18]. Ideally, such quality improvement initiatives would be aligned to the TB care cascade—as an organizing framework and outcome measure—and be informed by theories of change aimed at strengthening health systems. Such initiatives, if well implemented, have the potential to continuously generate ideas and interventions for health system change while allowing assessment of the feasibility of those ideas.

3. Research to address key gaps in the care cascade

In the following sections, we describe specific questions that may be relevant to each gap in the cascade (Table 1).

3.1. Gap 1. Case-finding

Addressing the case-finding gap is contingent on understanding who is missed by current case-finding efforts and how to decrease delays faced by those who are eventually diagnosed. There are a few reasons why individuals with TB in the community may not get evaluated and access a TB test. First, they may not have access to TB services, due to distance or other barriers. Second, they may not seek care for their symptoms, even if services are available. Finally, even if they do seek care, HCPs might not recognize their symptoms as being concerning for TB and initiate appropriate evaluation.
Table 1
Research questions relevant to each gap in the TB care cascade.

| Research questions                                                                 | Potential research approaches                                                                 | Relevance                                                                 |
|-----------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|--------------------------------------------------------------------------|
| **Gap 1: Case-finding**                                                            |                                                                                               |                                                                          |
| Which populations do not have access to TB services?                               | • Analysis of data from national demographic and health surveys                                | • May help to identify locations where TB services need to be expanded to ensure access to high-risk populations |
|                                                                                  | • Local exercises mapping the geographic distribution of notified patients in relation to the availability of TB services | • May identify populations that would benefit from novel community-based strategies such as use of health extension workers for TB screening |
| Why do some individuals with active TB in the population not seek care or delay seeking care? | • Interviews with individuals diagnosed with TB in prevalence surveys who have not sought care | • May help guide targeting of public education strategies via radio, television, or social media |
|                                                                                  | • Interviews with individuals with symptoms concerning for TB in the community who have not sought care | • May help identify the types of individuals who should be prioritized in community active case-finding activities |
|                                                                                  | • Interviews with TB patients who had substantial delay in seeking care                         |                                                                          |
| Why do some healthcare providers (HCPs) not refer individuals for TB testing?       | • Questionnaires using clinical vignettes to assess HCP knowledge                               | • May help identify types of HCPs who lack necessary knowledge or provide suboptimal care with regard to TB evaluation and testing |
|                                                                                  | • Standardized patient studies to assess actual HCP behavior                                    | • Standardized patient and knowledge assessments provide approaches for testing the benefits of interventions aimed at modifying behavior, including education of HCPs, use of incentives, and provision of support through public-private initiatives |
|                                                                                  | • Qualitative research to understand HCP’s clinical decision-making                             | • Understanding HCPs clinical decision-making may facilitate educational strategies targeted at shifting their behavior |
| How can case detection rates of active case-finding (ACF) initiatives be increased? | • ACF trials focusing on high-risk groups, such as household contacts, people living with HIV (PLHIV), or individuals with silica exposure | • May help identify the most efficient approaches for focusing ACF initiatives to increase the case detection and therefore the number of individuals entering the TB care cascade |
|                                                                                  | • ACF trials using identification of geographic TB hotspots to facilitate spatial targeting of case-finding approaches |                                                                          |
| **Gap 2: Diagnosis**                                                              |                                                                                               |                                                                          |
| Which patients disproportionately do not get diagnosed with TB?                    | • Cross-sectional studies using exit interviews with structured or qualitative data collection to identify patients presenting to different health system levels who have not been tested for TB despite having symptoms | • May help to identify whether certain groups are being disproportionately missed |
|                                                                                  | • Cohort studies to understand which patients are not being appropriately tested                |                                                                          |
| Why do some patients not get appropriately diagnosed with TB, despite getting evaluated and tested? | • Patient pathways analyses to understand where TB tests are available in relation to patient care-seeking | • May help identify types of health facilities where World Health Organization (WHO)-approved TB tests are not accessible or feasible to implement, requiring a triage and referral mechanism |
|                                                                                  | • Cohort studies to understand risk factors for patient attrition during the TB diagnostic workup | • May help to identify patient characteristics that predict attrition during TB evaluation to facilitate development of targeted interventions |
|                                                                                  | • Qualitative research to understand barriers in the TB evaluation process                      | • May help to identify health system barriers that need to be addressed to facilitate completion of the TB diagnostic process or whether the appropriate diagnostic algorithms are being used |
| How do we improve diagnosis of TB test-negative (i.e., smear-negative, Xpert-negative) TB patients? | • Cohort studies to understand patient attrition during the TB diagnostic workup, with a specific focus on TB diagnostic test-negative patients | • May facilitate approaches for simplifying algorithms for the diagnostic workup of test-negative TB to reduce patient attrition |
|                                                                                  | • Qualitative research to understand barriers in the TB evaluation process                      |                                                                          |
| **Gap 3: Linkage to care**                                                         |                                                                                               |                                                                          |
| Why do some diagnosed TB patients experience pretreatment loss to follow-up (PTLFU)? | • Cohort studies to understand patient attrition during linkage to care                          | • May help to identify patient characteristics that predict PTLFU |
|                                                                                  | • Qualitative research to understand challenges in the process of linkage to care               | • May help to identify health system barriers contributing to PTLFU |
|                                                                                  |                                                                                               | • May inform development of technology- and human-resource-based interventions to improve linkage to care |
| **Gap 4: Retention on therapy and medication adherence**                           |                                                                                               |                                                                          |
| Why do some patients experience suboptimal TB treatment outcomes or medication non-adherence? | • Cohort studies to understand patient attrition during TB treatment or non-adherence to medications | • May help to identify patient characteristics that predict suboptimal treatment outcomes or medication non-adherence |
|                                                                                  |                                                                                               | • May help to identify health system barriers contributing to               |

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Identifying high-risk populations that have poor access to TB services is an initial step to reducing Gap 1. For example, historically marginalized populations—such as indigenous people living in the Brazilian and Peruvian Amazon, rural Canada, and rural India—have particularly poor access to TB services [19–22]. At the national level, demographic and health surveys may provide insights into populations that have poor access to TB services [23]; however, addressing this problem at a local level may require mapping exercises to understand health service availability in an area relative to the geographical distribution of notified TB patients. Such exercises would also need to account for biases that may result from notifications being higher in areas with better access to health facilities [24].

In settings where TB services are relatively accessible, it is critical to understand why some individuals with TB in the population may not seek care. TB prevalence surveys provide an opportunity to study this problem. Individuals diagnosed with TB during prevalence surveys can be interviewed to understand whether they have sought care, and, if they have not, what prevented them from seeking further care [3]. Prevalence surveys also provide quantitative information that may shed light on disparities in care-seeking behavior. For example, findings from prevalence surveys show that a meaningful proportion of individuals with TB may not seek care because they are asymptomatic, suggesting that the only way to identify such individuals early may be by using chest X-ray or novel biomarker-based screening as part of active case-finding [25]. In addition, a recent systematic review found discrepancies in prevalence and notification data that suggest men may be less likely to seek or access care in many settings [26]. In situations where care-seeking data are unavailable from prevalence surveys, similar data may be available for individuals in the population with TB-related symptoms [1]. Studies examining factors associated with delays in TB care seeking may also provide valuable information [27], since patients experiencing long delays may serve as a surrogate for understanding those who do not seek care at all.

Understanding why HCPs do not refer individuals with symptoms for TB testing requires research into HCP knowledge and behavior. Recent studies using standardized patients in India, China, Kenya, and South Africa have provided insights into HCP behavior when evaluating individuals with TB symptoms [28]. In addition to revealing universally low TB testing rates by HCPs, these studies show that patient characteristics—including gender, age, and biometric characteristics (e.g., body mass index)—have little association with HCPs’ decisions to test for TB [28,29], although male standardized patients reported significantly shorter interactions with providers and felt providers were less likely to take their worries seriously [29]. In contrast, HCP characteristics did influence rates of TB testing and correct management. HCPs with MBBS degrees perform better than non-MBBS providers in India [30]. Public sector HCPs perform better than private sector HCPs in Kenya [31]. Qualitative studies also provide unique insights into HCP behavior with regard to TB evaluation [32–34]. For example, in India, HCPs often defer or delay bacteriological TB testing in favor of empirical treatment [34]. Patient-pathway analyses (PMA) may help identify not only where patients seek care but also gaps in diagnostic capacity in public versus private or lower- versus higher-level health-care facilities [35,36].

Each of these problems in Gap 1 has different solutions that warrant evaluation. Increasing availability of TB services may be possible in geographic areas that are unconnected to health facilities using novel approaches, such as health extension workers [37]. Care-seeking behavior at the population level may be modified by public education strategies disseminated by radio, television, or social media. HCP knowledge of appropriate TB evaluation is low in many contexts [38,39]; however, even when HCPs have adequate knowledge, they often still do not appropriately evaluate for TB (the “know-do” gap) [40]. As such, increasing TB testing rates may require supporting HCPs, including ancillary providers such as community pharmacists, through public-private collaborations or provision of incentives [41,42].

Active or enhanced case-finding (ACF) strategies can circumvent the challenges of these other interventions by bringing TB screening to the doorstep of high-risk individuals; however, the optimal ACF approach remains elusive and will likely vary across settings. For example, a community-randomized trial of household-level enhanced case-finding in Zambia and South Africa did not demonstrate a decrease in TB incidence [43], while a trial in Vietnam demonstrated that conducting ACF on household contacts of TB patients was more effective in detecting TB than passive case finding alone [44]. With the advent of digital radiography with automated computer evaluation, there is renewed interest in community-based mass chest radiography screening campaigns, which were used with relative success in high-income countries in the 1930s-1960s [45].

There is also increasing recognition that people who have previously had TB are an important risk group, such that longitudinal follow-up of these individuals may increase case detection [46]. Refining ACF strategies in high-risk groups—such as household contacts, people living with HIV (PLHIV), or people exposed to silicosis—is a critical area for implementation research. Research is also needed to understand the benefits of spatial targeting of ACF by focusing on geographical hotspots with high TB incidence, which may also increase case detection [47].

### 3.2. Gap 2: Diagnosis

In the Indian and South African care cascades, Gap 2 revealed that...
many TB patients did not get successfully diagnosed, despite reaching health facilities and accessing TB diagnostic tests [1,2]. Certain groups are known to be at higher risk of missed diagnoses, often due to the imperfect sensitivity of existing diagnostic tests. These groups include PLHIV or those who are immunosuppressed for other reasons and children. Of note, these groups are more likely to have extra-pulmonary TB, which is more challenging to diagnose due to the need for biopsies and lower sensitivity of diagnostic tests on non-sputum specimens [48]. Studies that have used exit interviews with patients who present to healthcare settings in high-incidence settings identify missed opportunities for TB screening [49]. Further research is needed to identify whether certain groups, for example, women versus men [50], or patients with substance use are less likely to undergo recommended diagnostic evaluation.

The diagnostic gap may occur for several reasons. Sputum microscopy, which has relatively poor sensitivity, remains the dominant diagnostic modality in many high TB burden countries. Diagnosis of smear-negative pulmonary TB often relies on patients finishing multiple-step diagnostic algorithms associated with high rates of patient attrition [51,52]. Few high TB incidence countries have made higher-sensitivity WHO-approved TB tests (e.g., Xpert MTB/RIF) available at the most decentralized level (L0), which consists of care provided at health posts or by community health workers [53]. As such, patient pathways analyses suggest that TB patients are likely not accessing the best WHO-approved tests [36]. There has also been wide variability in the way in which Xpert MTB/RIF has been implemented in terms of indications for testing as well as geographic availability (e.g., urban versus rural settings [54]). In high incidence countries, such as India, where patients are more likely to initially seek private sector care, a modeling study suggests that rolling out Xpert MTB/RIF with restricted testing indications in the public sector alone might have limited impact on TB incidence [55]. Further research will help to understand test- and location-specific differences in diagnostic gaps in different contexts.

In order to close Gap 2, it is essential to understand that a diagnostic test in isolation cannot improve patient outcomes without efforts to strengthen the entire care cascade [17]. Several high-profile randomized trials of the implementation of diagnostic tests such as Xpert and urine LAM have not demonstrated mortality benefit [56-58]. Research is critical to understand the limitations, unrelated to a diagnostic test’s accuracy, which may result when a new test is implemented in real world settings. For example, the benefits of TB diagnostic tests have been undermined in South Africa by high rates of empirical treatment [59], centralized laboratory testing, and challenges in obtaining sputum samples [60]. Qualitative research can provide insights into how patients navigate diagnostic ecosystems, including understanding why tests may not function as intended in real world settings [61].

While improving access to existing WHO-endorsed diagnostic tests is critical for closing the diagnostic gap, there is also need for new TB diagnostic tests that could help close Gaps 1 and 2 by allowing for more rapid TB diagnosis, facilitating identification of drug-resistant TB via rapid susceptibility testing, and facilitating triage and disease rule out in the community [62,63]. Research should also investigate how diagnostic algorithms for bacteriological test-negative TB can be simplified—for example, by earlier use of radiological studies—to ensure patients get diagnosed before being lost to follow-up.

### 3.3. Gap 3: Linkage to care

Systematic reviews suggest that patient losses from PTLFU (Gap 4) may be more substantial than those during the entire TB treatment course in some high TB burden settings [1,2,64]. The reasons that patients diagnosed with TB do not start treatment are diverse and include patient and health system factors [64]. For example, some studies suggest that particular patient characteristics predict higher risk of PTLFU, including having previously been treated for TB [65], older age [65,66], male sex [66], and weakness due to advanced TB [66–68]. Of these, having a prior TB treatment history is of particular concern, since these patients often also have poorer treatment outcomes and are at higher risk for having drug-resistant TB [1,65]. Health system factors found to contribute to PTLFU include: site of diagnosis (e.g., hospitals [69] or tertiary and TB specialty centers [65]), failure to communicate sputum test results to patients [64,68], challenges in navigating between health facilities [68], and dissatisfaction with waiting times [64]. In Indian studies, missing patient contact information in health records was a major barrier to being able to track these “lost” patients [65,66,70,71]. There is a notable paucity of qualitative research evaluating PTLFU, highlighting an area where further studies are needed. The high-quality qualitative studies that have been conducted emphasize the role of health system barriers in contributing to PTLFU [68,72].

The literature suggests that interventions should at least partly focus on addressing health system barriers. Technology may have a role in improving efficiency of care delivery after diagnosis. For example, electronic medical records may improve recording of patient contact information, so that patients can more easily be tracked, and automated SMS texts may help notify patients of their TB diagnoses [73]. Human resource-based solutions are perhaps even more critical. For example, patient navigators (i.e., individuals tasked with helping patients reach next steps in care) and patient tracking interventions may help prevent loss to follow-up, especially from high-volume tertiary hospitals [74]. The literature on interventions to reduce PTLFU is sparse, highlighting a need for high-quality implementation studies. Few studies have looked at PTLFU in higher-risk patients, such as those with drug-resistant TB.

### 3.4. Gap 4. Retention on therapy and medication adherence

Gap 4 comprises poor outcomes during TB therapy, due to treatment failure, loss to follow-up or death [3]. This has historically been the only gap routinely reported by national TB programs. As a result, Gap 4 has been a central focus of TB care delivery research in recent decades. Systematic reviews have evaluated studies on TB treatment outcomes to understand reasons for mortality [75], medication non-adherence [4], and interventions to improve adherence and reduce loss to follow-up [8].

One systematic review showed that, in high TB burden settings, individuals with drug-resistant TB, HIV co-infection (especially with advanced immunosuppression), older age, and undernutrition have higher TB case-fatality rates [75]; other studies from high burden settings have highlighted strong associations between tobacco or alcohol use and poor TB treatment outcomes [76,77]. In lower burden settings, non-infectious comorbidities (e.g., diabetes, chronic lung disease, renal disease, malignancy) and injection drug use were additional factors associated with increased TB mortality [75].

Research has helped define approaches for addressing some of these risk factors to improve TB patient outcomes. For example, randomized trials have shown that early initiation of antiretroviral therapy in PLHIV with active TB and advanced immunosuppression is associated with improved survival [78–80], and rigorous evidence affirms the benefits of drug-susceptibility testing and treatment with individualized drug regimens for patients with multidrug-resistant (MDR) TB [81].

In spite of the research already conducted to understand and address poor treatment outcomes, we argue that there is need for new research—particularly on risk factors that have been less actively studied—to inform development of novel interventions to reduce Gap 4. For example, undernutrition is a major predictor of poor treatment outcomes; however, trials to assess benefits of macro- and micro-nutrient supplementation are relatively sparse and inconclusive [82,83]. A recent study from Ethiopia found that 54% of TB patients had probable depression at treatment initiation [84]. Untreated depression was associated with three times increased relative risk of death and nine times increased risk of loss to follow-up [84]. And yet, few studies have assessed the impact of treating depression on TB outcomes. Studies
showing promising benefits of treating alcohol use disorder [85], social protection schemes (e.g., cash transfer) for TB patients [86–88], and using community-based strategies (e.g., psychosocial support groups [89]) merit broader evaluation.

Approaches to monitoring medication adherence, particularly DOT, have been a major focus of research, under the assumption that such monitoring is critical for ensuring optimal treatment outcomes. However, systematic reviews have found conflicting results regarding whether DOT yields better outcomes than self-administered therapy, although most suggest little benefit of most DOT approaches [5–9,90,91]. More recently, research has focused on using DATs to "electronically observe" pill-taking [10,92]. Findings regarding the accuracy and impact of DATs on treatment outcomes remain mixed [10,92]. Studies suggest that some DATs have poor accuracy for measuring TB medication adherence [93,94] and show no benefit for improving treatment outcomes [95,96], while others have found higher accuracy or improvements in adherence or treatment outcomes with use of these technologies [97–99]. These mixed findings regarding both DOT and DATs suggest that benefits of these interventions are dependent on the local context, technology used, design of the monitoring strategy, approach to intervening upon non-adherence, and quality of implementation. Using DATs to facilitate human interaction (e.g., provider-patient communication, early identification of medication adverse effects)—rather than simply using the technology to observe patients—may be associated with improvements in outcomes [98,99].

In general, future research on Gap 4 should focus on interventions that move beyond simply monitoring TB patients and towards actually providing support to address their needs.

3.5. Gap 5. Post-treatment TB recurrence-free survival

Gap 5 comprises TB recurrence or death after completing treatment. Assessing Gap 5 is most important during the initial year after a patient completes TB treatment, because most disease recurrence occurs within 12 months of finishing therapy [100]. Post-treatment deaths are also relevant to capture as part of this gap, because TB patients who achieve cure or treatment completion continue to have elevated mortality, part of which may be due to undiagnosed disease recurrence or TB related sequelae [101].

Patient characteristics that may predict TB recurrence include male sex [102] and prior treatment history [103], highlighting groups who may benefit from additional support during treatment. For all TB patients, the risk of disease recurrence partly reflects quality of care received during therapy. For example, undiagnosed drug resistance [104,105], suboptimal medication adherence [103,104,106], and smoking [76,104] are independently associated with increased risk of disease recurrence. These findings suggest that improvements in diagnostics (to facilitate early identification of drug resistance), support for patient adherence, and treatment of comorbidities could potentially reduce TB recurrence.

Gap 5 has implications not only for patient management during therapy but also for post-treatment care. One potential implication of high TB recurrence rates in some contexts [105,107] is that ensuring regular post-treatment follow-up with ongoing screening for TB symptoms may facilitate early identification of disease recurrence, which could effectively serve as a form of ACF. For example, a recent modeling study suggests that ACF among previously treated TB patients, as well as secondary prophylaxis with isoniazid therapy for some patients, could accelerate reduction in TB incidence in South Africa [46]. Routine post-treatment follow-up could also facilitate management of emerging chronic conditions, including post-TB lung disease [108–110] and increased cardiovascular risk seen in individuals with recent TB [101]. Involving affected communities to provide insights regarding wellbeing after TB is critical to inform the post-TB research agenda, given the breadth of TB-related complications, which range from psychological ill-health to disabilities (e.g., hearing loss) to catastrophic socioeconomic consequences [111].

3.6. Risk factors that contribute to multiple gaps in the care cascade

Some patient characteristics are associated with poor outcomes across multiple care cascade gaps. For example, a recent systematic review suggests that men with TB in the community are less likely to reach care and get notified (i.e., started on treatment) by national programs [26]. Studies from a variety of settings also suggest that men may be at higher risk of death while on TB treatment [75] and for experiencing post-treatment disease recurrence [102]. In some settings, patients with a prior treatment history may be more likely to suffer from PTLFU [65], suboptimal treatment outcomes [1,112], and post-treatment disease recurrence [103]. Individuals with drug-resistant TB in particular suffer from disproportionately poor outcomes at every care cascade gap [1,2,113]. Given that many TB patients lack social support to engage in care, community-based care and strategies for facilitating social support may be beneficial to TB patients at multiple care cascade stages, based on evidence from the HIV and maternal health literature [114–116]. Patients with these various characteristics may benefit from dedicated interventions to address their needs at every stage of care.

4. Applying this research agenda in different geographic scales and populations

The multifaceted agenda described above is meant to highlight key gaps in knowledge that may be addressed through studies conducted at different levels of geographic scale and in diverse populations (Table 2). For example, we have previously advocated that national TB programs could use multisite prospective cohort studies, with representative sampling of health facilities, to achieve nationally-representative estimates of patient losses at key care cascade stages—from diagnosis to recurrence-free survival [3]. Rigorous measurement of clinical, psychosocial, and health system factors for patients included in such studies might simultaneously identify characteristics that predict patient attrition to inform national-level interventions and policies.

However, nationally-representative studies may provide suboptimal information regarding barriers to engagement in the TB care cascade for key high-risk populations—such as PLHIV (particularly in low HIV prevalence settings) [117], people who live in slums [118], people who inject drugs [119,120], prisoners [121], migrants [122], miners [123,124], individuals with silicosis [125], and healthcare workers [126] to name a few. Unique sampling methods may be required for these sub-populations. For example, finding people who inject drugs with TB—to understand care-seeking behavior and care cascade dropout—may require screening and follow-up of individuals recruited by respondent-driven sampling or from opioid agonist therapy centers [127]. Finally, cohort and qualitative studies to understand care cascade outcomes at the local city, district, hospital, or clinic level may help to directly inform local interventions and quality improvement initiatives.

5. Conclusion: Need for bold and innovative research on the care cascade

In the last few years, there has been substantial progress in developing approaches for measuring care cascades for active TB disease in high TB burden countries [1–3,128]. However, while the TB community has gained a better understanding of the scale of patient losses throughout the cascade, we still have major shortcomings in our knowledge of how to reduce these gaps in care [129]. For some gaps, such as PTLFU and post-treatment disease recurrence, there has been a paucity of research given the scale of these problems. Even for interventions aimed at addressing gaps that have historically been dynamic areas of research—such as ACF approaches or strategies for promoting TB medication adherence—there are important limitations in our
knowledge regarding their efficacy or optimal approaches to implementation. As such, research needs to be expanded across all levels of the evidence generation pathway (Fig. 2). TB researchers can take inspiration from the extensive research on the care cascade that has been conducted by the HIV community. Closing gaps in the care cascade has the potential to more rapidly accelerate reduction in TB incidence [128,130]; however, achieving this goal will require urgent investments in a bold and innovative action-oriented research agenda.

**Ethics statement**

Note that this is a review article for which no ethical clearances are required.

**Declaration of Competing Interest**

None.

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**Table 2**

Potential geographic scales or population focuses of interest for the care cascade research agenda.

| Geographic scale or population of interest | Potential research approaches | Limitations of the research approaches when applied in a given geographic scale or population |
|-------------------------------------------|------------------------------|------------------------------------------------------------------------------------------|
| National TB programs / country-level studies | - Nationally-representative cohort studies to identify predictors of poor care cascade outcomes  
- Systematic mapping of populations without access to TB services  
- Assessing reasons individuals with active TB have not sought care in large-scale TB prevalence surveys  
- National service mapping may identify major service gaps but miss barriers to health facility accessibility for local subpopulations  
- Reasons for not seeking care or dropping out of care by may vary for subpopulations in local contexts | |
| Key high-risk populations (e.g., people living with HIV, people who inject drugs, slum residents, tribal populations, migrants, refugees, miners, individuals with silicosis, healthcare workers) | - Cohort studies to identify predictors of poor care cascade outcomes by screening and follow-up of affected individuals at specific sites (e.g., HIV clinics, opioid agonist therapy centers, etc.) or using unique sampling methods (e.g., respondent-driven sampling)  
- Qualitative studies may provide rich information that is generalizable to others in the affected sub-population  
- Findings in a given high-risk population may have limited generalizability outside of that sub-population | |
| Local city or district TB programs, hospitals, or clinics | - Cohort and qualitative studies to understand reasons for poor care cascade outcomes  
- Findings may directly inform local changes in care delivery but may have limited generalizability | |
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