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Drug-resistant tuberculosis control in South Africa: scientific advances and health system strengthening are complementary

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We examine some aspects of the South African health system that have contributed to the current multi- and extensively drug-resistant tuberculosis (M(X)DR-TB) epidemic and identify opportunities for change and improvement. Implementation of several recent major scientific advances have the potential to accelerate the control of M(X)DR-TB, but health systems strengthening will be essential.

Keywords: extensively drug-resistant tuberculosis, health systems strengthening, multi-drug-resistant tuberculosis

Decades of sub-standard, poorly implemented tuberculosis (TB) control programme guidelines and interrupted TB treatment, combined with neglect in drug development, has seen the emergence of both multi- and extensively-drug-resistant TB [M(X)DR-TB] [1] – a public health crisis of particular concern in South Africa where both TB and HIV epidemics have converged. Therapeutic options for drug-resistant TB are limited by efficacy, acceptability, availability, tolerability and financial constraints. Since 2006, the number of M(X)DR-TB cases in South Africa has increased temporally [2] and in 2013 the WHO identified South Africa as having the second-highest number of MDR-TB cases worldwide [3]. Since the prevalence of drug-resistant TB is considered a measure of the effectiveness of existing TB control programmes [4], these statistics emphasise grave inadequacies in current treatment and management strategies for TB in South Africa and highlight an ailing health system. Here, we examine aspects of the South African public health system which have contributed to the current situation and evaluate the role of scientific advances in the control of drug-resistant TB.

The treatment of M(X)DR-TB is protracted, toxic, resource-intensive [5] and plagued by poor patient adherence [6]. South African data for the period 2000 – 2004 show that standardised treatment for MDR-TB resulted in an overall treatment success rate of only 46% [7]. The mortality rate in HIV-infected patients was 35.2%, more than double than in HIV-negative patients [7]. Treatment success rates are considerably lower in XDR-TB, an ominous threat to gains made in HIV-TB programmes. A 24-month study of XDR-TB in South Africa in which 73% of the cohort was HIV co-infected, but only 61% received combined anti-retroviral therapy (cART), reported a treatment success rate of just 22% [8]. Since then there has been compelling scientific evidence demonstrating the role of concurrent second-line anti-TB chemotherapy and cART administration in improving sputum conversion rates and reducing mortality in M(X)DR-TB-HIV co-infected patients [8-10]. Encouragingly, the National Tuberculosis Programme responded and the South African ART guidelines were amended in 2010 to incorporate the...
recommendation that TB-HIV co-infected patients be initiated on cART irrespective of the CD4 cell count. Three years on, data from the third quarter of 2013, at one of the largest TB drug-resistant treatment facilities in South Africa, show that 19% of MDR-TB-HIV co-infected patients were not initiated on cART.

There is some evidence that operational challenges in implementing the current guidelines and the sub-optimal use of existing drugs contribute to low treatment success rates [11]. Such operational challenges occur at various levels of the health system. A recent study describes the treatment journey of a typical patient, and highlights numerous flaws within a health system that ultimately fails its dependants [12]. A 6-week time lag for notification of culture results; a month’s delay in hospital admission due to limited inpatient capacity; incomplete treatment regimens as a result of drug stock-outs and delayed cART initiation (attributable to poor integration of TB and HIV services) have since informally been accepted as ‘standard of care’. Alarmingly, recent data, from another province in South Africa, show that only 63% of newly diagnosed MDR-TB patients initiated treatment in 2011 [13]. Further, a random chart review of 186 patients referred to a centralised MDR-TB facility in KwaZulu-Natal in 2010 show that only 25% were initiated on treatment within 8 weeks, leaving 75% of patients experiencing a mean delay of 12.36 weeks (in addition to the 6–8 weeks required for culture and susceptibility testing) [14]. Such issues, in conjunction with patient-related socioeconomic problems make the Millennium Development Goal of eradicating TB by 2050 elusive in this resource-poor setting.

Novel anti-TB agents such as bedaquiline and delamanid and re-purposed drugs such as linezolid, significantly shortened treatment programmes such as the ‘Bangladesh Regimen’ [11] and innovative point-of-care diagnostics such as Xpert MTB/RIF® personify the scientific advances considered essential in regaining control of drug-resistant TB but the question remains whether they can have the desired impact on their own.

The advent of Xpert MTB/RIF has since substantially reduced the time to MDR-TB treatment initiation by approximately two-thirds at some sites in South Africa (Dr Iqbal Master, personal communication, October 2013); however, its ability to provide a diagnosis within a few hours suggests that this delay could, in theory, be further reduced. Xpert MTB/RIF has also paradoxically further intensified the burden, increasing MDR-TB case detection resulting in an escalation in the number MDR-TB cases initiated on treatment (Figure 1) and an unwavering patient admission waiting list (Figure 2). Figure 2 illustrates the number of patients with M(X)DR-TB awaiting admission at a TB specialist hospital with a maximum capacity of 192 beds, over a 7-year period. Although fluctuating, the numbers remain unacceptably high and reveal a lack of intervention to ensure a sustained decline in waiting time, despite this facility initiating treatment in 65% of patients in outpatient care. The limited inpatient capacity has other important public health implications. Pietersen et al. found that 42% of patients with XDR-TB who were discharged had failed treatment [15]. Further, these patients had a median survival time of 19.84 months from the time of discharge. These data reveal a critical gap in current TB control programmes where potential disease transmission requires urgent interruption.

Unlike scientific advances, TB control programmes appear to have stagnated. These factors demonstrate a health system unable to exploit the potential of available resources. Cumulatively, they are an ominous threat to drug-resistant TB control, exacerbating patient-related socioeconomic problems,
fuelling transmission in vulnerable communities, urgently requiring strategic intervention.

**Expert opinion**

Major developments in drug availability, shortened, better-tolerated treatment regimens and innovative diagnostic tools have the potential to vastly enhance current programmes and accelerate the control of M(X)DR-TB. Although these advances highlight the way forward, they too will inevitably fail in a fragile health system. The burden of the M(X)DR-TB has prompted a culture that strives towards quantity as opposed to quality of healthcare administered, often neglecting to implement even the most basic principles of care without capitalising on available tools. It is imperative that we do not underestimate the role and value of health systems’ strengthening. We need to identify weaknesses and develop strategies that target such limitations. Stakeholder commitment to TB control, improved provider performance and enhanced morale in healthcare workers, combined with the necessary support and infrastructure, will be essential in containing the scourge of this disease. Unless these interventions complement each other, an unfortunate outcome is inevitable.

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**Declaration of interest**

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Bibliography


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