The scientific response to TB – the other deadly global health emergency

In 1993, the WHO declared TB, an airborne infectious disease, a global public health emergency and urged coordinated efforts by all nations to avert millions of deaths in the coming years. On January 30, 2020, the WHO declared COVID-19, another airborne infectious disease, a public health emergency of international concern. However, the similarity between the global responses to these two pandemics ends there. What we have witnessed in the past 2 years in terms of the scientific, public health, medical, and pharmaceutical communities to COVID-19 is nothing short of spectacular. Within 2 weeks of declaring COVID-19 a global emergency, the WHO had convened a meeting of experts and issued a research roadmap. National governments, especially that of the United States, rapidly committed vast sums of money into research at all levels, from basic virology and immunology to clinical care and prevention. Pharmaceutical companies launched development programs for new products to diagnose, treat and prevent COVID-19. As a result, diagnostics, therapeutics, and vaccines have been developed at a dizzying pace, delivering an array of tools that provide us with the means to control and end the SARS-CoV-2 pandemic. The effective and equitable deployment of those tools is a challenge of monumental proportions, but no one can claim that science has been found wanting in responding to the global crisis.

Since 1993, TB on the other hand has not been treated as a true emergency. This is perhaps because it was not new, was not escalating at a frightening pace, and had already experienced a golden age of discovery in the 20th century. But even if it was not new and there were tools available to combat it, its worldwide distribution, impact on health, and mortality burden was just as dire, and the need for a rapid, coordinated, and adequately resourced scientific response was just as evident. The recently issued 2021 WHO Global Tuberculosis Report discloses the disturbing news that TB incidence remains plateaued at 10 million cases per year, but that in 2020 case detection fell by almost 20% and mortality rose for the first time in a decade to 1.5 million deaths. The decline in diagnoses and increase in mortality is directly attributable to the COVID-19 pandemic.

The US National Academies of Science, Engineer-
glacial. Funding for TB research is less than half of what the United Nations and WHO estimate is necessary to achieve the End TB targets, and the Treatment Action Group’s report on research funding shows a 16-year average of just $659 million from 2005 to 2020. A comparison of investments in TB research vs. research into COVID-19 is astonishing (Table 2). But even with limited funding, there have been some triumphs in TB research in the past decade: molecular assays make diagnosis possible in less than 2 hours, rather than 2–4 weeks; treatment of drug-susceptible TB has been shortened to 4 months, the first reduction in duration in 40 years; and treatment of multidrug-resistant TB has been shortened from 2 years of noxious, injectable agents to 6 months of an all-oral regimen; treatment of drug-susceptible TB has been shortened to 4 months, the first reduction in duration in 40 years; and treatment of TB infection has been cut from 9 months to as short as 1–3 months with safer and better tolerated regimens. However, one universal truth is that none of these transformative advances occurred as quickly as they should have.

Compared to COVID-19 and HIV grants, NIH funding opportunities for TB biomedical research are limited and reviews of TB applications proceed according to a languid schedule. Following funding awards, the sequential and often redundant regulatory and ethical review processes at each participating institution further delay activation of the research, and therefore the results. For example, a suite of TB preventive studies funded by Unitaid and addressing WHO high-priority areas (such as interactions between TB drugs and antiretroviral drugs in pregnant women and children with HIV infection) continues to be held in a stranglehold by regulatory procedures. The WHO’s Ethics Committee, supported by overtaxed volunteer experts, can take an average of 10–12 months to review a protocol. Approval by national and local IRBs in high-burden countries can then take an additional year. The overall timeline for conducting critically important TB research is scandalously long; clinical trials for TB generally take a very long time because the endpoints are slow to accrue, but most studies are unnecessarily prolonged by painfully long administrative and regulatory review processes.

The broader problem, however, is much larger than the mechanics of individual funding agencies or regulatory bodies. First, nobody is treating TB as an actual emergency! As we have seen with COVID-19, when everyone thinks it is an emergency, people act differently, and things move rapidly. At the NASEM meeting, a South African government researcher reported waiting 6 months for approval of a minor protocol amendment to a study on lifesaving treatment for multidrug-resistant TB. We have experienced long delays with our trials in a number of countries. Second, the clinical and public health research infrastructure is vastly underfunded and under-supported. Much of our focus is on individual researchers who clamor (justifiedly) for more money, but the remainder of the machinery of clinical research is largely neglected. COVID-19 has demonstrated what is possible when researchers, funders, and regulatory agencies unite to confront a crisis. Game-changing trials of therapeutics and vaccines can be conducted in record time without cutting corners and compromising participant safety and scientific integrity, if everyone acts as if it is an emergency. But to do so requires a radical change in our collective mindset in addition to substantially greater human and financial resources.

Operating in crisis mode for COVID-19, TB, or any other health catastrophe is difficult to sustain. But the COVID-19 pandemic has shown us what works to accelerate progress against a global threat. First, substantial funding for priority research multiplies innovation and progress. As a starting point, governments, pharma/biotech companies, and foundations must increase investment in TB research, at least to the levels laid out in the UN High Level Meeting Report and make TB a central element in global pandemic response strategies. Moving forward, the level of ambition must be raised. There is a growing recognition from the COVID-19 experience that the funding targets for TB research are far too low – and the scale-up of newly developed tools is far

### Table 1 Comparison of TB and COVID-19 vaccine development.

|                          | TB*                                                                 | COVID-19†                                                                 |
|--------------------------|---------------------------------------------------------------------|--------------------------------------------------------------------------|
| Year pathogen discovered | 1882                                                               | 2019                                                                    |
| Number of vaccines licensed for use | 1 (bacille Calmette-Guérin)                                     | 25                                                                     |
| Number of vaccines in clinical trials | 15                                                             | 112                                                                    |
| Development timeline of representative vaccine candidates | M72/AS01E (Gates Medical Research Institute, 2020 onward; formerly GlaxoSmithKline) | BNT162b2 (Pfizer/BioNTech)                                               |
| Preadclinical work begins | Early 2000s (Mtb72F, Corixa Corp)                                 | January 2020 (BioNTech)                                                  |
| First Phase I trial starts | 2004 (results published 2009)                                     | May 2020 (combined phase I/II)                                           |
| Pivotal Phase II trial starts | 2014 (starts) 2018 (primary analysis published)                    | July 2020 (combined phase I/II/III)                                     |
| Phase III starts         | 2023 (expected start)                                              | December 2020 (UK)                                                      |

* Information on TB vaccine pipeline and M72/AS01E development timeline from Treatment Action Group Tuberculosis Vaccines Pipeline Report and WHO Report of the high-level consultation on accelerating the development of the M72/AS01E tuberculosis vaccine candidate
† Information on COVID-19 vaccine pipeline and BNT162b2 development timeline from New York Times Coronavirus Vaccine Tracker (cited 13 December 2021).
Table 2 A comparison of government funding for research on TB and two estimates for COVID-19 therapeutics and vaccines, 2020

|                       | TB* US$      | COVID-19† US$ | COVID-19‡ US$ |
|-----------------------|--------------|---------------|--------------|
| **Total research funding** | 915 million (all areas) | 53 billion (vaccines only) | 104 billion (vaccines + therapeutics) |
| **Funding for vaccine research** | 118.6 million | 53.5 billion | 98.9 billion |
| Public funding for vaccine research | 77.5 million (65%) | 51.4 billion (96%) | 98.9 billion (100%) |
| Percentage of public funding committed via advanced purchase agreements | 0% | 88% | 98% |
| **Philanthropic funding for vaccine research** | 38.7 million | 85.4 million | — |
| Private sector funding for vaccine research | 2.4 million | 517.8 million | — |
| Multilateral funding for vaccine research | 0 | 1.4 billion (CEPI) | — |
| Funding for long-term consequences of disease (post-TB lung disease15 and long COVID) | No estimate, but minimal | 1.15 billion (US only) | — |

* TB funding data comes from the Treatment Action Group and Stop TB Partnership report Tuberculosis Research Funding Trends, 2005–20208 which tracks research expenditures (actual disbursements) across six areas of TB research: basic science, diagnostics, drugs, vaccines, operational research/epidemiology, and infrastructure/unspecified projects.
† The Knowledge Network on Innovation and Access to Medicines published estimates of COVID-19 vaccine funding (disbursements and commitments) with data drawn from the Policy Cures Research COVID-19 R&D trackers and ACT-Accelerator Tracker; last updated July 8, 2021.
‡ The kENUP Foundation published estimates of public funding for COVID-19 vaccines and therapeutics (disbursements and commitments) in the first 11 months of the pandemic in January 2021.

too slow. Governments and other funders must commit to more to end TB by 2030. Second, the funding timeline can be greatly reduced; peer review for NIH HIV-related grants serves as a useful model for TB applications, with review occurring within 2–3 months of submission and funding within 6 months. If the rationale for implementing aggressive timelines for reviewing biomedical research in HIV and COVID-19 was the recognition and fear that these infections would rapidly spread and kill, then TB grants should likewise be reviewed rapidly. Third, the regulatory bottleneck must be cleared. Additional investment in regulatory and ethical infrastructure (including training and international coordination) is necessary to ensure that these critical requirements do not suffocate innovative research. Unnecessary regulatory reviews only add delay while providing no protection for study participants and their communities. Finally, governments must treat TB as a central element in global pandemic response strategies. The new focus on pandemic preparedness – most notably the beginning of negotiations by the WHO to create a legally binding pandemic treaty or similar mechanism – must include a commitment to end ongoing pandemics such as TB. If an annual 1.5 million deaths due to TB is not a pandemic, then what is?

The bottom line is that advances in TB diagnostics, treatments, and prevention that can translate into progress in TB elimination need to be pursued and then scaled up with the sense of urgency they deserve. If we do not behave like TB is a global health emergency, we will continue to see agonizingly slow progress in developing tools to End TB, as well as unacceptable suffering from a disease that has killed more than 20 million people in this century alone.

R. E. CHAISSON1
M. FRICK2
P. NAHID3

1Johns Hopkins University School of Medicine, Baltimore, MD, 2Treatment Action Group, New York, NY, 3UCSF Center for Tuberculosis, University of California, San Francisco, San Francisco, CA, USA

Correspondence to: Richard E Chaissom, Johns Hopkins Center for Tuberculosis Research, 1530 Orleans St, IM.08, Baltimore, MD 21287, USA.
email: rchaiss@jhmi.edu

Disclaimer: The viewpoints expressed are those of the authors and do not necessarily reflect the views of our institutions or the National Academies of Science, Engineering and Medicine.
Conflicts of interest: none declared.

References
1 World Health Organization. TB, a global emergency. Geneva, Switzerland: WHO, 1994. https://apps.who.int/iris/bitstream/handle/10665/58749/WHO_TB_94.177.pdf?sequence=1&isAllowed=y Accessed November 2021.
2 World Health Organization. Director-General’s statement on IHR Emergency Committee on Novel Coronavirus (2019-nCoV). Geneva, Switzerland: WHO, 2019. https://www.who.int/director-general/speeches/detail/who-director-general-s-statement-on-ihr-emergency-committee-on-novel-coronavirus-(2019-ncov). Accessed November 2021.
3 World Health Organization. COVID-19 strategic preparedness and response plan. Geneva, Switzerland: WHO, 2021. https://www.who.int/publications/i/item/WHO-WHE-2021.02. Accessed November 2021.
4 World Health Organization, Global tuberculosis report, 2021. Geneva, Switzerland: WHO, 2021.
5 Migliori GB, et al. Global Tuberculosis Network. Gauging the impact of the COVID-19 pandemic on tuberculosis services: a global study. Eur Respir J 2021; 58:5: 2101786.
6 The National Academies of Sciences, Engineering, and Medicine. Innovations for Tackling Tuberculosis in the Time of COVID-19: Current Tools and Challenges: Proceedings of a Workshop, 2021 https://www.nationalacademies.org/our-work/innovations-for-tackling-tuberculosis-in-the-time-of-covid-19-a-two-part-workshop-seriesTAG Funding Report. Accessed November 2021.
7 National Institutes for Health, Rapid acceleration of diagnostics (RADx), 2020, https://www.nih.gov/research-training/medical-research-initiatives/radx Accessed November 2021.
8 Treatment Action Group. Tuberculosis research funding report, 2005–2020. New York, NY, USA: TAG, 2021. https://www.treatmentactiongroup.org/wp-content/uploads/2021/12/tb_funding_2021.pdf Accessed December 2021.
9 Boehme CC, et al. Rapid molecular detection of tuberculosis and rifampin resistance. N Engl J Med 2010; 363: 1005–1015.
10 Conradie F, et al. Nix-TB Trial Team. Treatment of highly drug-resistant pulmonary tuberculosis. N Engl J Med 2020; 382: 893–902.
11 Dorman SE, et al.; AIDS Clinical Trials Group; Tuberculosis Trials Consortium. Four-month rifapentine regimens with or without moxifloxacin for tuberculosis. N Engl J Med 2021; 384: 1705–1718.
12 Sterling TR, et al., TB Trials Consortium PREVENT TB Study Team. Three months of rifapentine and isoniazid for latent tuberculosis infection. N Engl J Med 2011; 365: 2155–2166.
13 Swindells S, et al.; BRIEF TB/A5279 Study Team. One month of rifapentine plus isoniazid to prevent HIV-related tuberculosis. N Engl J Med 2019; 380: 1001–1011.
14 World Health Organization. Geneva, Switzerland: WHO, 2018. https://www.who.int/news-room/events/un-general-assembly-high-level-meeting-on-ending-tb Accessed November 2021.
15 Migliori GB, et al. Clinical standards for the assessment, management and rehabilitation of post-TB lung disease. Int J Tuberc Lung Dis 2021; 25(10): 797–813.