Confronting the scientific obstacles to global control of tuberculosis

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Tuberculosis (TB) is a major threat to global health, recently exacerbated by the emergence of highly drug-resistant forms of the disease-causing pathogen and synergy with HIV/AIDS. In 2006, the Stop TB Partnership published “The global plan to stop TB: 2006–2015,” which set out a vision of halving the prevalence of and mortality caused by the disease by 2015, followed by eliminating the disease as a public health problem by 2050. This vision depends on the development of improved diagnostics, simpler treatment, and more effective vaccination. Recently, active translational research pipelines directed toward each of these goals have been established, but improved understanding of the fundamental biology of this complex disease will prove to be the key to radical advances in TB control.

Introduction

Tuberculosis (TB) is today the second highest cause of death from an infectious disease worldwide, after HIV/AIDS (1), and is the biggest killer of people infected with HIV (reviewed in ref. 2). Much of the burden of this disease falls on those living in the developing world. A recent report on the TB pandemic (3) revealed that in 2005, there were almost 9 million new cases and 1.6 million deaths. The latest data indicate that the incidence rate has probably stabilized, but the challenge is to reverse the trend and reduce global mortality and, in the long term, to eliminate the disease.

Most humans infected with the pathogen that causes TB, Mycobacterium tuberculosis, remain asymptomatic, and only a small proportion develop active TB disease immediately. TB is usually a chronic, slowly progressing disease that often remains undiagnosed in patients for many years. The disease has many clinical manifestations and can affect many organs, but the most common form in adults is a chronic pulmonary disease (Figure 1). In most people, the bacterium establishes a latent infection, and the lifetime risk of developing disease is only 10% unless an individual becomes immunocompromised, at which time the risk increases to about 10% per year (4–6). Unfortunately, it has been estimated that a third of the world’s population is latently infected with M. tuberculosis, providing an enormous reservoir for future disease.

“The global plan to stop TB: 2006–2015” (7, 8) was developed and endorsed in 2005 by the Stop TB Partnership, a network of international organizations, countries, donors from the public and private sectors, governmental and nongovernmental organizations, and individuals who expressed an interest in working together to achieve the goal of eliminating TB as a public health problem. It established a number of specific targets. The target for 2005 was to detect 70% of new sputum smear–positive cases (i.e., cases in which a positive diagnosis is obtained after spreading a sputum sample on a microscope slide, fixing, applying a stain, decolorizing with acid, counterstaining, and visualizing the acid-fast tubercle bacilli; Figure 1) and cure at least 85% of these cases (although this goal was not met, substantial progress toward it was made); for 2015, to sustain or exceed those indicators and reduce the prevalence and death rates of TB by 50% relative to the 1990 level; and for 2050, to eliminate TB as a public health problem. However, the current tools for diagnosing, treating, and preventing TB are inadequate for this task: latent and active TB are most commonly diagnosed using, respectively, a skin test that dates back to 1891 (9) and sputum-smear microscopy from the same era (10). The current “short-course” drug therapy consists of a cocktail of drugs taken over a period of at least six months, and despite its widespread use, the Mycobacterium bovis bacillus Calmette-Guérin (BCG) vaccine is largely ineffective, at least in preventing adult pulmonary disease. A growing problem that has had an impact on the success of treatment programs is the emergence of strains of M. tuberculosis resistant to the drugs used as first-line treatment, and around 400,000 cases of multidrug-resistant TB (MDR-TB), which is defined as resistance to isoniazid and rifampicin, occur per year (11). Furthermore, essentially untreatable outbreaks of extensively drug-resistant TB (XDR-TB), which is defined as MDR-TB plus resistance to a quinolone and one of the second-line anti-TB injectable drugs (Amikacin, Kanamycin, and Capreomycin), have begun to appear. In HIV-positive individuals, XDR-TB has been associated with very high mortality (12).

Achieving the goals in the Stop TB plan will require the development of new tools that are substantially better than those available today. This, in turn, will require an advance in our knowledge of the disease and the biology of M. tuberculosis as well as the application of innovative new approaches and technologies. Much progress has been made in the past decade, and a number of new tools are under development (13–22). However, much more needs to be done. In this Review, we look ahead to how the major challenges that must be overcome in the three areas of diagnosis, treatment, and prevention can be addressed to make a quantum leap forward (Table 1). Progress in each of these areas would also be accelerated by the identification and validation of biomarkers, biological markers that correlate with the disease status of the host or the response to intervention.
is often localized to the lung, and by the time patients present to a
clinic, is an infrequent cause of positive sputum smear tests in coun-
tries where TB is endemic (14). There has been a great deal of pro-
gress recently in developing better tools to detect latent TB (23–26),
and understanding the role and relevance of these assays is an area
of active research. However, we have chosen to focus here on detec-
tion of active disease, as these new tools will have the greatest impact
initially on the countries where the burden of disease is highest.

One would think that these aspects of TB would make it possible
to achieve rapid and substantial gains in controlling TB globally
with a standardized application of existing diagnostic methods.
This indeed was one of the assumptions underpinning the DOTS
(directly observed therapy, short course) TB control strategy
launched by the WHO as a highly cost-effective health intervention
in 1994 (27). DOTS has conventionally relied on passive detection
of microscopy smear–positive samples from patients presenting to
health clinics to detect cases and initiate therapy. In the absence of
an effective vaccine, this has been the cornerstone of TB control.

Although DOTS has been very successful at standardizing care
practices and increasing cure rates, case-detection targets have been
more difficult to achieve (1). Furthermore, overall TB control, as
measured by a decline in incidence, especially in settings where
drug resistance or HIV coinfection are prevalent, has been limited
(1). The core problem is that existing microscopy methods to diag-
nose TB are both technically and practically inadequate for use in
high-burden countries. Microscopy is too complex to implement
and too slow to perform as to be used as a point-of-care test. The pri-
cy clinics where most patients with symptoms of TB first seek
care rarely have AFB microscopy available, and patients must com-
monly wait until they are referred to a specialty clinic or microscopy
center. Even then, most TB cases will be missed, both because of
the inherent limits of the sensitivity of the test, which detects only
60% of culture-positive pulmonary TB patients even in clinical trial
settings, and because of the logistical difficulties of making quality-
assured microscopy widely and freely available. Moreover, micros-
copy misses exactly the patients, those with early disease, that one
would like to detect and put on treatment to block transmission
before it has begun. Delays in diagnosis of 3–6 months are common
because of the lack of sensitivity of microscopy (28, 29), during
which time the disease can progress and cause severe destruction
of the airways (Figure 1) and transmission continues.

Mycobacterial culture is much more sensitive than microscopy
but requires equipment and training beyond what is available at
peripheral public health laboratories. Most countries with a high
burden of TB have no more than a handful of laboratories with
culture capability, so most patients never have access to this tech-
nology. When they do, results are often so slow as to be of little
clinical relevance. This has proven especially true in patients with
advanced HIV infection, where undetected or improperly treated
TB can have a very high and early mortality, with survival mea-
sured in days or weeks (30–32). Conventional indirect drug-sus-
cceptibility testing (DST) on solid media is slower still and requires
laboratory infrastructure that is so rarely available that only 3% of
the 6.5 million TB cases arising annually in high-burden countries
of Asia and Africa have access to DST (33).

A broad effort to develop and make available better diagnostics
for TB is underway, and substantial gains have already been made.
The feasibility and cost effectiveness of using rapid liquid culture
systems for case detection and DST has recently been demonstrated
in a number of high-burden countries. Their broader implementa-
tion has been facilitated by price negotiation with manufacturers
and by the recent development of a simple and inexpensive method
for species identification of culture isolates by immunocapture of the TB-specific protein MTB64 on lateral flow (34). Inexpensive and robust fluorescence microscopy platforms based on LED excitation have also been developed to speed TB detection in microscopy laboratories (35, 36). Although gains will be made with improved diagnostics, access to primary health care by patients is also a critical factor in early detection of TB, and further improvements in this are also required to maximize the impact on the epidemic.

Beyond conventional microbiologic methods, there are three molecular assays that are expected to be available to the public sector in the coming 6 to 18 months that can dramatically speed case detection and MDR screening without the need for biosafety precautions associated with phenotypic methods. One is a reference-level test employing manual PCR followed by line-probe hybridization to detect resistance-associated mutations (37). The other two molecular assays use automation (38, 39) that should allow them to be used in microscopy laboratories or similar settings.

Progress in developing true point-of-care tests, however, has been slower. Our ability to develop the types of tests that are most needed and in general to improve the control of TB through better diagnosis, is hampered by knowledge gaps related to the epidemiology and natural history of tuberculosis disease, host and pathogen biology, and biodetection technologies. Some examples of these knowledge gaps follow.

Epidemiology and natural history of TB. Although careful studies of transmission of the TB-causing pathogen in defined settings, such as during long-haul airplane flights (40), in school classrooms (41), and on ships (42), have been used to make assumptions about the duration and intensity of exposure to M. tuberculosis needed to generate secondary cases, little is known about the kinetics of transmission in the community. Specifically, the fraction of transmission that occurs (in settings of both high and low HIV prevalence) before index cases first report to the formal health system and what fraction occurs during the diagnostic process is unknown. This information would be useful in determining the importance of speed of testing and in understanding the degree to which active case-finding efforts in the community should be considered in different settings.

Although detection of individuals who have TB but are not smear positive has not been prioritized by national TB control programs in most countries, our assumptions related to the importance of such cases in disease transmission have not been well tested. Although there is an extensive literature describing the relatively low level of transmission from patients who are smear negative compared with their smear-positive counterparts (43, 44), little is known about the transmission impact of smear-negative TB cases in settings where prevalent HIV infection increases susceptibility to infection with M. tuberculosis and disease. Furthermore, it is not known what fraction of patients with smear-negative TB go on to develop smear-positive disease nor in what time frame.

Biology. M. tuberculosis causes a range of pathology in humans and has a complex and highly evolved interaction with the immune system. Unfortunately, our understanding of the biology of the pathogen and of host-pathogen interactions in different disease states is incomplete, despite an extensive amount of work in this area. This biological uncertainty has largely frustrated attempts to exploit the host response as an effective diagnostic tool. We do not currently have any biomarkers, for example, that can be used to definitively distinguish individuals with prior, cured infection from those infected with dormant bacilli or from those with bacterial replication or incipient disease.

Laboratory studies have demonstrated that specific proteins can be upregulated in specific in vivo growth conditions, for example, within phagosomes (45). However, we know little about gene expression levels in M. tuberculosis as it is found in pulmonary cavities, inside granulomas, or even circulating in the blood of an infected individual (46). More fundamental, from a diagnostic point of view, is our lack of information about the concentration of potentially detectable M. tuberculosis antigens or other moieties in clinical matrices such as urine, serum, sputum, or saliva. Published studies describing the potential for detecting mycobacterial antigens in clinical samples have, in large part, not been systematic, and most have focused on individual proteins or glycolipids abundant in culture filtrate such as MPT32 (47), the antigen 85 complex (48–50), the 38 kDa protein (51), and lipoarabinomannan (LAM) (52–56). Detection of mycobacterial antigens is an attractive approach to TB diagnosis, with the theoretical benefits of high specificity, correlation to mycobacterial burden, and independence from immune function. A number of proteomic approaches to the discovery of novel diagnostic target projects have been carried out (57, 58), but none has yet identified a readily detectable marker with demonstrated clinical utility.

As mentioned earlier, the development of simple molecular approaches to the detection of drug-sensitive and drug-resistant M. tuberculosis is a promising development. The development of more complete molecular methods for drug-resistance testing, which is both faster and safer than phenotypic methods, is hampered by incomplete knowledge of the molecular determinants of resistance for many of the important second-line agents used in treating MDR disease.

Technology. At present, there are only three methods with proven clinical utility for the diagnosis of TB: microscopy, culture, and nucleic acid amplification. None of these approaches is simple enough to
use in peripheral health care settings such as primary care clinics in the developing world. In fact, the only diagnostic technologies that have been widely used in lower levels of the health system for any purpose are enzymatic test strips for glucose monitoring in diabetes and immunochromatographic tests for pregnancy and various infectious diseases. Thus, developing simple point-of-care tests for TB requires either the dramatic simplification of one of the three current TB diagnostic methods or the successful development of tests based on the detection of diagnostic antibodies or TB-specific antigens. Both of these options present important technical challenges.

Lateral flow tests, such as commonly used for pregnancy testing, and other similar formats, suffer from limitations of sensitivity and are unlikely to detect mycobacterial proteins in the concentrations in which they are likely to be found in serum. Fortunately, a number of diagnostic companies are competing to develop point-of-care technologies that can detect low-abundance molecules in the pg/ml range and below.

TB moieties are likely to be much more abundant in sputum than in urine or serum. Sputum, however, is exceedingly difficult to work with as a diagnostic matrix, being viscous, inhibitory to many types of reactions, and highly variable in both nature and quantity. A range of processing protocols have been developed to liquefy or otherwise prepare sputum for use in diagnostic tests (59), but unfortunately, most of the protocols require laboratory procedures and equipment and are more complex than the detection methods they are meant to support. Although specimen-processing research is not attractive to research funding institutes, simple sample-processing methods that allow detection of DNA, RNA, or unaltered antigens are critically needed to accompany novel product development. Such translational research across all aspects of diagnostic research is now in full swing, and there is real hope for rapid point-of-care tests to diagnose both infection state and drug susceptibility.

Drugs

The current approach to treatment of TB comprises a cocktail of drugs taken daily over a period of at least six months, initially in a two-month “intensive” phase with four drugs — isoniazid, rifampicin, pyrazinamide, and ethambutol — followed by a four-month “continuation” phase with isoniazid and rifampicin. The last decade has seen a revival of drug development activity, and a number of drug candidates have been identified and are undergoing clinical evaluation (18) (Table 2).

Toward an ultra-short treatment regimen. *M. tuberculosis* bacilli are AFB that adapt well to growth in vitro in defined media with relatively simple requirements: a stable pH, an adequate carbon source, abundant oxygen, a surfactant to counteract the natural tendency of the organism to aggregate, a buffer for toxic lipids (typically serum albumin), and sufficient micronutrients to support essential enzymatic functions. The past two decades have seen the sequencing of the *M. tuberculosis* genome (60) and the development of ample genetic tools to manipulate the tubercle bacillus almost as casually as *Escherichia coli* (61). These tools have been employed to map essential genes (62–64) and define the biochemical function of many potential enzyme targets in the tubercle bacillus (65). In theory, the development of new drugs against *M. tuberculosis* would seem to simply matter a matter of recapitulating the formalisms of conventional antibacterial programs — identify an essential target, preferably informed by a genomic analysis, develop potent inhibitors of the selected target, and chemically modify such inhibitors to confer drug-like qualities such as good oral bioavailability and low toxicity for the human host.

Unfortunately, this simple model of drug discovery has numerous pitfalls even when applied to more typical bacterial pathogens (66). Identifying hits can be surprisingly difficult for unusual bacterial enzymes due to a lack of biologically relevant chemical diversity, and translating successful enzyme inhibitors into compounds with activity against whole cells often fails because of poorly understood rules for bacterial cell penetration. Even when potent enzyme inhibitors penetrate cells effectively, failure to understand the biological “system” in sufficient detail to identify not only essential, but also rate-limiting, biochemical transformations often translates into failure. Worse, when applied to *M. tuberculosis*, even this imperfect model of drug discovery, and at times it seems any rational model of drug discovery, completely collapses beneath the complexities and peculiarities of the disease.

By the time an accurate diagnosis of TB is made, there is a substantial amount of tissue destruction spread among multiple lesions — each of which operates as an independent infection. Some lesions, such as those on the surface of cavities (often spherical lesions with air-liquid cores open to the bronchi and leaking mycobacteria to the pulmonary surface), contain actively replicating bacilli, whereas others (nodules or consolidations of various sizes) are still successfully containing their mycobacterial invaders in highly structured collections of immune cells known as granulomas (67). This heterogeneity of mycobacterial populations means either that a particular target must be vulnerable across all of the physiological states occupied by the bacilli or that the agents in a

<table>
<thead>
<tr>
<th>Stage of development</th>
<th>Drug candidate</th>
<th>Sponsor</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phase I clinical trials</td>
<td>LL-3858</td>
<td>Lupin Ltd.</td>
<td>(115, 116)</td>
</tr>
<tr>
<td>Phase I clinical trials</td>
<td>SQ-109</td>
<td>Sequella Inc.</td>
<td>(117–119)</td>
</tr>
<tr>
<td>Phase II clinical trials</td>
<td>OPC-67883</td>
<td>Otsuka Frankfurt Research Institute GmbH</td>
<td>(75, 120)</td>
</tr>
<tr>
<td>Phase II clinical trials</td>
<td>PA-824</td>
<td>Global Alliance for TB Drug Development</td>
<td>(121, 122)</td>
</tr>
<tr>
<td>Phase II clinical trials</td>
<td>TMC207</td>
<td>Tibotec</td>
<td>(84, 123)</td>
</tr>
<tr>
<td>Phase III clinical trials</td>
<td>Moxifloxacin</td>
<td>Bayer</td>
<td>(124, 125, 126)</td>
</tr>
<tr>
<td>Phase III clinical trials</td>
<td>Gatifloxacin</td>
<td>Global Alliance for TB Drug Development CDC University College of London Johns Hopkins University</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>OFLOTUB European Commission WHO-TDR Lupin Ltd.</td>
<td>(127, 128)</td>
</tr>
</tbody>
</table>

CDC, Centers for Disease Control and Prevention; WHO-TDR, WHO Special Programme for Research and Training in Tropical Diseases.
cocktail used must individually have targets that collectively cover all such states (68). Lesions containing mycobacteria in nonreplicating states are particularly problematic, as such mycobacteria have an intrinsically low basal metabolism that renders them much less susceptible to conventional mycobactericidal agents. In addition, the very structures created by the immune system to isolate and contain the mycobacteria (i.e., the granulomas) are poorly vascularized, dense structures, often with necrotic, acellular cores that may represent a substantial barrier to drug penetration from the bloodstream (see the extensive consolidation in the patient in Figure 1 for example). Finally, the cell envelope of M. tuberculosis represents a barrier for drug penetration more formidable than virtually any other bacterium in nature (69).

The conventional whole-cell mode of antibacterial discovery offers one path forward for the discovery of new drugs that target M. tuberculosis. With screens targeting replicating cells, however, such a path is likely to uncover only compounds with properties similar to those we have today and is unlikely to result in the quantum leap forward necessary to achieve rapid sterilization, shorten treatment times, and therefore improve global health. This is not to say that such strategies should be abandoned; MDR- and XDR-TB are important public health threats that deserve considerable effort, but the root cause of the global problem with drug-resistant TB-causing pathogens is the armamentarium of slowly acting agents currently used to treat drug-susceptible disease. Various strategies, which are discussed in detail below, are being engaged to combine the best of conventional methodology with the latest conceptual advances to transform TB drug discovery from lottery to logic.

Targeting nonreplicators — the “hypoxia hypothesis” and beyond. Researchers in the TB field have learned some important lessons from lung cancer, another chronic disease with many similar pathological features. For example, anoxic lesions are a hallmark of some tumors, and there are many reasons to think that such anoxia is also a feature of some tuberculous lesions (70). M. tuberculosis is exquisitely adapted to respond to hypoxia and has a dedicated transcriptional program to realign metabolism for survival under hypoxic conditions (71). There are also in vitro models for generating hypoxia-induced nonreplicating cells, and these cells are refractory to killing by conventional antibiotics yet become susceptible to killing by anaerobe-specific agents like metronidazole (72). That observation has spurred an ongoing human clinical trial of the activity of metronidazole in MDR-TB patients (73). In addition, the hypoxia hypothesis underlies the excitement surrounding the development of nitroimidazoles such as PA-824 and OPC-67683, which are currently in clinical development for the treatment of TB and which have activity against both replicating and nonreplicating anaerobic cells (74–76). It is perhaps more telling that virtually any new agent developed for the treatment of TB today is screened for activity against anaerobic nonreplicating cells. Primary screens to identify lead compounds against such cells are part of several TB drug development programs.

Hypoxia, of course, is just one feature of the microenvironment in which the tubercle bacillus is embedded during disease. The lesional pH, the source of carbon, and the abundance of iron and various vitamins and cofactors are all measurable variables in different niches occupied by M. tuberculosis (77). Understanding the other essential features of the microenvironments present in the spectrum of lesions within patients with both active and latent TB infection would both identify novel targets and allow the reconstruction of in vitro conditions that reproduce these environments exactly — in other words, predictive models of drug efficacy. Current efforts to understand the biology of disease in increasingly realistic higher animal models are also essential to ground experimental hypotheses and test the effect of interrupting specific drug targets during established disease (78).

My kingdom for a target — the chemical genomics revolution. There is deep uncertainty regarding how relevant a target identified as essential for M. tuberculosis growth under aerobic conditions on glucose as a primary carbon source (63) is to growth under conditions encountered by the organism in the host. It is therefore difficult to identify which targets, when interrupted by a small molecule during human disease, will result in a therapeutically beneficial effect. Perhaps even knowing which pathways are engaged by the bacillus during infection does not necessarily identify the step in the pathway that would make a suitable target for small molecule inhibition. In part, this is because some steps in a given pathway are naturally not rate limiting; thus, even a small amount of enzymatic activity may be sufficient to allow enough metabolic flux through a pathway in which even 99.9% inhibition would be insufficient to restrict mycobacterial growth or affect viability.

An attractive combination of conventional approaches using whole-cell screening and more modern genomics techniques is “chemical genomics,” which has been used to explore other areas of biology (79–82). In the case of TB, this technique aims to identify metabolic vulnerabilities of the mycobacteria under defined conditions. The technique at first glance looks like a normal whole-cell compound-based screen for activity, but by manipulating the in vitro assay conditions, one can select hits that are selective for one set of conditions or that are general for many sets. These hits are then coupled with genome-scale biology tools such as transcriptional profiles and/or genome resequencing technologies to identify the specific target being affected by the particular hit. The output from this strategy is a pharmacologically validated target that can then be the starting point for a conventional drug development program starting from high-throughput enzyme-based screening and extending through medicinal chemistry and lead optimization.

What will it take to get to a “Z-pack” for TB? A patient with a bacterial upper respiratory tract infection caused by Streptococcus pyogenes often receives 5 days worth of curative therapy with azithromycin in a single package of 6 pills known as a Z-pack. Two weeks of therapy to cure a TB patient has an almost magical, transformative feeling to it. Scientifically, is this even a reasonable goal? The heart of the issue is really the fundamental killing kinetics of both replicating and nonreplicating cells. A culture containing more than 10^6 replicating cells/ml can be killed by a mixture of isoniazid and gatifloxacin in less than three hours in vitro (83), and although the killing kinetics are slower for stationary-phase cells, it still takes only four days to achieve similar kill with isoniazid and rifampicin. It is unclear why it then takes six months of therapy to cure a TB patient if these rates of kill apply in the patient. In the past, an issue like kill-rate would be a dissociated observation, made long after the commitment to a particular target and or molecular series was firm. This is an inevitable consequence of the old way of screening that puts the compound first and thinks about the biology later. Although this can be occasionally successful, as the development of the ATP synthase inhibitor R207910 clearly demonstrates (84), it doesn’t offer a sustainable way forward since the only real target validation, utility in human disease, must await the results of clinical trials of this agent.
At this point in time, there is much scientific uncertainty in understanding the fundamental difference between the in vitro and the in vivo kill rate, although there is considerable attention and effort directed at this phenomenon. The problem can at least be restated productively as understanding the effect of perturbing one step in a dynamic system encompassing both host and pathogen. From the vantage point of “systems biology,” at least the shape of the solution can be seen. Systems biology is roughly defined as the intersection between experimental and computational biology, and although in its infancy, this emerging alliance of biologists, programmers, and mathematicians holds considerable future promise for revolutionizing the search for TB drugs. Currently it is possible to describe discrete biochemical pathways as “systems” and integrate genetic and biochemical information to explore the effect of perturbing a step in such a pathway and predict which interventions would disrupt the pathway (85). Robust and predictive in silico models of even a single bacterial cell are still elusive but will undoubtedly emerge in the near term. In the future, what we need to do is integrate these models and pathways and reconstruct the physiology of both the organism and the host. Such models will allow us to predict which pathways would provide therapeutically useful responses.

Another equally important strategy is the development and validation of predictive animal models that more faithfully reflect human disease and an understanding of the effect of current agents in detail in these models. Nearly all of the current drugs in use have never been examined in anything other than mice, which develop little of the characteristic pathology of human disease. Without understanding the strengths and weaknesses of our current agents, we cannot begin to understand where they fall short. Likewise, most of the major TB drug trials were done in an era where there were few tools for looking in detail at the response to therapy both as reflected in the human immune response and even physically in terms of high-resolution imaging techniques. The cycle of a major expensive development program followed by a lengthy phase III clinical trial with only relapse as an end point must be broken. We must learn more from fewer patients and we must learn these things much faster if the results are to feed back into the quality of our model systems (whether animal or in silico) and inform the next generation of targets taken forward.

Vaccines

**How close are we?** Considering the success of the natural immune response in controlling most TB infections, it is attractive to propose that a vaccine that can be delivered in advance of exposure to the pathogen and that mimics the natural immune response will further reduce the incidence of disease. Running counter to this “classical” vaccine approach is the fact that a major portion of cases of TB — particularly the sputum smear–positive cases that contribute most to the further spread of infection — arise from a population of individuals who successfully controlled an initial infection but then failed to mount a protective response to a subsequent reactivation. In contrast to a typical vaccine-preventable disease, individuals whose immune system has been stimulated by a prior bout of active TB remain highly susceptible to subsequent reactivation (86). The conventional vaccination model, in which the adaptive immune response learns by repeated stimulation, seems to run backwards in the case of TB.

These two lines of argument offer radically differing prognoses for TB vaccine development. According to the first, we should learn the characteristics of the natural response and reproduce them using subunit and attenuated microbe tools of conventional vaccinology. From the second viewpoint, we should search for deficiencies in the natural response and repair these using some alternative form of immune manipulation. The classical vaccinology approach offers the potential for immediate practical application and is clearly a major priority for global health research and development. In parallel, it is, however, prudent to explore the fundamental biology required to underpin development of alternative vaccine approaches.

**Front-line candidates and the IFN-γ hypothesis.** Albert Calmette and Camille Guérin followed Pasteur’s strategy of mimicking the natural response to infection using an attenuated pathogen to produce the BCG vaccine (87). An extensive literature attests to the fact that prior exposure to BCG vaccine accelerates development of the adaptive immune response to *M. tuberculosis* in animal models, conferring advantages in terms of reduced mycobacterial load and increased time to death (88–91). The same pattern can be seen in humans in the context of severe forms of childhood TB (92). However, BCG has variable efficacy against adult forms of TB, showing least effect where it is needed most (93). This is generally ascribed to an inherently weak immunogenicity, to adverse immune-mediated interactions with environmental mycobacteria, and to waning of immune memory (94). A logical approach to overcoming these problems is to boost BCG by priming an additional pool of mycobacteria-reactive T cells. This can be achieved by delivering *M. tuberculosis* antigens as purified proteins in adjuvant or in the form of recombinant viral vaccine vectors. Four such vaccines are lead candidates in current human clinical trials (22) (Table 3).

**Antigens.** In spite of intensive efforts in experimental models, there are few guidelines for rational selection of vaccine antigens. An ability to elicit a recall response in healthy infected individuals provides evidence that a particular antigen is available as an immune target and is used as a common criterion. Proteins secreted from live mycobacteria become available for antigen presentation before those released by mycobacterial lysis and may have an advantage in triggering an earlier immune response. While this seems logical, rigorous proof for preferential recognition of secreted proteins is lacking and both secreted and nonsecreted antigens are included in current candidates (22) (Table 3). In general, collective effort has favored a family of mycolyl transferases found in the growth medium of mycobacterial cultures (Ag85A and Ag85B) and a family of low molecular weight proteins transported by specialized secretion systems (ESAT6, TB10.4) (95). With increasing use of ESAT6 in diagnostic tests, it would seem preferable to avoid its use in vaccines, as microbial antigens that are under selective pressure as targets of a protective immune response often undergo genetic variation, although there is no evidence of this in the case of the commonly studied *M. tuberculosis* antigens (96). It might be that protective immune responses in TB target a very broad repertoire of antigens, or it might be that we have yet to identify the key protective antigens.

**Cytokines.** Protection against TB in mice is critically dependent on the Th1 immune response, an interplay among DCs, T cells, and macrophages that is promoted by IL-12 and IFN-γ (97, 98). Similarly, in humans, rare genetic defects that impair the IL-12–IFN-γ axis are associated with hypersusceptibility to mycobacterial disease (99). The central role of IFN-γ in stimulating antimicrobial functions of macrophages has encouraged its use as a marker for the beneficial immune response induced by vaccination (100). Although it is clear that some level of IFN-γ is essential for protec
tion, the hypothesis that more IFN-γ equates with better protection remains to be proven. Clinical trials of the current front-line vaccine candidates will test this “more-better” hypothesis.

A multiplicity of mechanisms. Although the IFN-γ hypothesis provides a simple marker for monotheistic vaccine development, many other immune parameters are affected by vaccination and might make an important contribution to efficacy (Figure 2). Multicolor flow cytometry provides a powerful tool for analysis of cytokine production and, by analogy with other infections, polyfunctional T cells that produce a combination of IFN-γ and IL-2 might be relevant to protection (101). It will be important to maximize the information gathered from current vaccine trials by including measurement of a broad range of immune parameters.

The particular cell type responsible for production of IFN-γ might be important. CD4+ T cells produce IFN-γ in response to antigens presented by MHC class II molecules. By responding to antigens presented by MHC class I molecules, CD8+ T cells might trigger responses to different cells at different stages of the infection process. BCG is less effective than M. tuberculosis in promoting presentation of antigens by the MHC class I pathway, due to reduced antigen transfer to adjacent cells (“cross-priming”) and perhaps to differences in lysis of phagosomal membranes (102). Recombinant BCG engineered to express a membrane-damaging hemolysin might have enhanced vaccine properties as a result of increased MHC class I expression (103). Similarly, CD1-restricted and γδ T cells that are specialized in recognizing nonprotein antigens from mycobacteria might offer further support to the overall protective response (104), although their potential for priming by vaccination is uncertain.

In addition to IFN-γ stimulation of antimicrobial functions, CD8+ T cells have the ability to kill infected cells by injection of cytotoxic granules. Depending on coordination with other immune cells, this could have a detrimental effect on promoting dissemination of the M. tuberculosis infection or a beneficial effect on exposing bacteria to freshly activated macrophages. Cytotoxic granules might also include antimicrobial peptides that have a direct effect on the intracellular mycobacteria. This could provide an important killing mechanism that is relevant for neutrophils and NKT cells, in addition to CD8+ T cells, that is complementary to IFN-γ activation, and that involves alternative signaling molecules such as vitamin D metabolites (105).

The range of immune mechanisms that might be involved in protection is increasing as our knowledge of immunology expands. A subset of T cell expressing the cytokine IL-17 is activated during mycobacterial infection. However, it remains to be determined whether this response plays a beneficial role — for example, by stimulating early recruitment of neutrophils — or a detrimental role in promoting pathology (106, 107).

Regulation and collateral damage. The toxic effector molecules of the immune response are released within localized environments to maximize their chances of causing damage to the pathogen rather than to the host. Prolonged activation inevitably results in damage to surrounding tissues, however, and the ability to turn off an aggressive response is as fundamental to survival as the ability to turn it on. The importance of regulation of the immune response to TB is seen in the immune reconstitution inflammatory syndrome (IRIS), which causes acute pathology when an unregulated immune response is restored by antiretroviral therapy in individuals coinfected with HIV and M. tuberculosis (108). Dissection of regulatory mechanisms has proven a major challenge to immunology. Current interest focuses on a subset of CD4+CD25+ T cells that express the Foxp3 transcription factor, with cytokines IL-10 and TGF-β contributing to regulatory control (109). Perhaps rather than building a stronger Th1 response, a TB vaccine should aim to enhance the effectiveness of the preexisting response by resetting regulatory circuits.

Presumably, the limitation in the natural immune response — the reason that only 90% of the population is protected and that
individuals who contain an initial infection remain at risk of secondary disease — is that it must balance the benefit of attacking the pathogen against the risk of unacceptable self damage. This equation might vary with the individual’s age. The first few years of life are associated with high susceptibility to TB, but very little disease is seen in older children (110). Susceptibility reemerges after adolescence and might be further enhanced in old age. From an evolutionary perspective, it can be argued that the natural immune response to TB has been selected to maximize the chance of reaching child-bearing age, leaving the researcher with potential scope for artificial enhancement in later life.

Immune subversion. The lungs of a patient with pulmonary TB are transformed into an aerosol generator, and this is central to effective transmission. This transformation is dependent on an active Th1 immune response, comprising the same cells and cytokines that are required for protection. A successful disease cycle for <i>M. tuberculosis</i> involves an ability to withstand the initial impact of the immune response to establish an infection together with the ability to elicit a subsequent immune response that is sufficiently aggressive to promote transmission. Does <i>M. tuberculosis</i> actively program the host immune response in its own favor? It is attractive to imagine delivery of some inhibitor of the inflammatory response that is effective when mycobacterial numbers are low together with an inhibitor of the regulatory response that is effective at high population densities. Hints that <i>M. tuberculosis</i> might engage in constructive dialogue with the immune system are beginning to emerge from studies of the diversity among clinical isolates of <i>M. tuberculosis</i> (111). At least in animal models, subtle changes in the surface components expressed by <i>M. tuberculosis</i> can have radical effects on subsequent pathogenesis. For example, phenolic glycolipid produced by a subset of <i>M. tuberculosis</i> isolates belonging to the W-Beijing family show “hyperlethality” in murine disease models. Disruption of phenolic glycolipid synthesis results in loss of this hypervirulent phenotype without significantly affecting bacterial load during disease (112).

One form of immune subversion that has been proposed for <i>M. tuberculosis</i> is the induction of a Th2 response — measured in terms of production of IL-4 or its splice variants — analogous to that mounted in response to infection with intestinal parasites and during allergy (113). Th2 cytokines are certainly detectable in TB patients and probably contribute to disease-associated immunopathology. It remains to be determined, however, whether Th2 responses (triggered either by <i>M. tuberculosis</i> or by coinfecting parasites) influence earlier stages of infection.

The importance of immune subversion in the context of vaccine development is that, rather than trying to reproduce all of the details of the natural response to <i>M. tuberculosis</i>, there might be certain elements of the response that we should try and eliminate. Perhaps BCG is suboptimal because it is too close to the natural pathogen? An ideal vaccine strain might be one in which attenuation is mediated, not by loss of some physiological function necessary for survival, but as a consequence of induction of a stronger immune response. The phenotype of such a strain might include rapid clearance during infection of immunocompetent animals.

**Figure 2**
Mycobacterial infection activates a broad spectrum of immune responses. Evidence from experimental models, from rare human genetic conditions, and from HIV-associated TB demonstrate that the Th1-mediated IL-12–IFN-γ axis is essential for prevention of progressive disease. Priming this response provides the major focus for the vaccine candidates currently being assessed in clinical trials. A broader view of the immune response suggests multiple alternative strategies that may influence the outcome of infection, including delivery of IFN-γ by alternative lymphocyte subsets, release of bacteria by cytolysis of poorly microbicidal cells, and activation of IFN-γ–independent pathways of mycobacterial killing. It is probable that all of these activities are subject to regulatory control mechanisms in vivo, and successful vaccination may ultimately depend on establishing or resetting multicellular immune networks rather than overactivation of a single pathway.
**Concluding remarks**

There are formidable obstacles to development of point-of-care TB diagnostic tests, drugs that will make it possible to treat a patient with TB in weeks instead of months, and a vaccine that will provide long-term protection from infection. As the scientific enterprise struggles toward an ever-expanding knowledge base of both the physiology and the host-pathogen dynamic, the disease marches steadily on, killing millions each year. Fortunately, never in history has more brain power been focused on the scientific issues surrounding TB. Never has there been a higher chance that a real breakthrough in detection, treatment, and prevention tools will transform global health.

**Acknowledgments**

The authors wish to thank Adriana Alejandro-Osorio for her help in preparing the manuscript.

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1264