With tuberculosis (TB) having plagued mankind for centuries, there can be no doubt that Mycobacterium tuberculosis, the causative agent of human TB, has been successful in adapting for human infection. M. tuberculosis belongs to the Mycobacterium tuberculosis complex (MTC), itself comprised of bacterial agents responsible for TB or TB-like disease. Members of the MTC are known to infect mammalian hosts, and the extent and consequence of this infection is gaining greater recognition in part because of the availability of diagnostic tools to classify specific isolates appropriately. This article introduces the tools and terminology used for this classification and illustrates their utility by discussing work from independent laboratories that have established a genome-based phylogeny for the MTC [1–5]. Next, it considers the use of these markers to distinguish atypical isolates not conforming to attributes of traditional MTC members [6,7]. Finally, it discusses the current genomic evidence regarding the origin and evolution of M. tuberculosis in the context of its relevance for TB control in humans and other mammalian hosts.

Characteristics of the Mycobacterium tuberculosis complex

The MTC consists of bacteria that genetically share identical 16S rRNA sequence and greater than 99.9% nucleotide identity. M. tuberculosis, M. africanum, M. microti, and M. bovis have been regarded as the four traditional species of the MTC, although the extent of MTC speciation is not yet resolved. In this article, MTC organisms are referred to as members, and the nomenclature provided in the most recent literature is used.

Members characteristically differ in their host range, epidemiology, clinical presentation in humans, and laboratory phenotype, although little is known about these differences or why these differences have evolved. The human form (M. tuberculosis sensu stricto) and the bovine form (M. bovis) have been nominally distinct for more than a century; other members have been identified more recently (Table 1). The members classically were described by their biochemical properties or by targeting their specific genetic regions. Genomic insights now show a new approach to MTC speciation outside the scope of these more traditional tools [8].

Genetic resources to study the Mycobacterium tuberculosis complex

With the availability of complete sequence information, several methodologies have developed to understand the MTC genetically. These methods can
be categorized as genetically fast or slow and as having phenotypic consequences or not. Each methodology has advantages and disadvantages. Whereas each methodology has proven useful, a tool is only as informative as the question toward which it is applied. Responsible contributions ideally should draw information from all available typing methods to conclude with the most parsimonious scenario.

**Fingerprinting patterns**

The use of DNA fingerprinting patterns, in which samples are genotyped by restriction-fragment-length polymorphisms using genetic attributes specific to the MTC as markers, has proven valuable for tracking MTC disease [9]. Molecular epidemiologic markers used include the MTC-specific insertion sequence IS6110 [10], polymorphic glycine- and cytosine-rich sequences [11], the direct-repeat region [12], spacer-oligonucleotide typing (spoligotyping) [13,14], and variable-number tandem repeats of genetic elements termed mycobacterial interspersed repetitive units [15,16]. Although these genetic markers are known to mutate at rates suitable for tracing a chain of disease transmission, their patterns of change are potentially too common to act as reliable markers over longer periods of evolutionary time. Therefore, they do not seem to be reliable for phylogenetic studies and speciation of clinical isolates.

**Table 1**

**Myobacterium tuberculosis** complex members

<table>
<thead>
<tr>
<th>MTC member</th>
<th>Natural host</th>
<th>Mouse</th>
<th>Guinea pig</th>
<th>Rabbit</th>
<th>Unique attribute</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>M. canettii</em></td>
<td>Human?</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>Most ancestral recognized MTC member, anecdotal isolation</td>
</tr>
<tr>
<td><em>M. tuberculosis</em></td>
<td>Human</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>Predominant cause of human TB</td>
</tr>
<tr>
<td><em>M. africanum</em> subtype II</td>
<td>Human</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>Reclassified as atypical <em>M. tuberculosis</em></td>
</tr>
<tr>
<td><em>M. africanum</em> subtype I (a)</td>
<td>Human?</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>Rarely isolated</td>
</tr>
<tr>
<td><em>M. africanum</em> subtype I (b)</td>
<td>Human?</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>Phenotypically heterogeneous</td>
</tr>
<tr>
<td><em>M. pinnipedi</em></td>
<td>Pinnipeds</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Closely related to <em>M. microti</em></td>
</tr>
<tr>
<td><em>M. microti</em></td>
<td>Vole</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>Attenuated, used as live vaccine in humans</td>
</tr>
<tr>
<td><em>M. caprae</em></td>
<td>Goat</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Only described in Europe?</td>
</tr>
<tr>
<td><em>M. bovis</em></td>
<td>Cow</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Dynamic pathogen with wildlife reservoirs?</td>
</tr>
<tr>
<td><em>M. bovis</em> BCG</td>
<td>None</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>Family of laboratory adapted strains of <em>M. bovis</em> used as live vaccine</td>
</tr>
</tbody>
</table>

**Abbreviations**: +, animal typically succumbs to infection; –, animal typically survives infection.

a Although previously suggested as a unique member of the MTC, *M. africanum* subtype II isolates cannot be genomically distinguished from *M. tuberculosis* [7]; and throughout this article, *M. africanum* subtype II is included within the *M. tuberculosis* lineage.

b Refers to the genotype ‘(a)’ of *M. africanum* subtype I having deleted RD9, but not RD7, RD8, and RD10.

c Refers to the genotype ‘(b)’ of *M. africanum* subtype I having deleted RD9, RD7, RD8, and RD10.

Data from Refs. [38,39,42,43,62,63].

**Sequenced genomes**

A wealth of genomic insight for the Mycobacterium genus is available through whole-genome sequence information for several species (Table 2), including six entire MTC genomic sequences completed or in progress. These are *M. tuberculosis* H37Rv [17], *M. tuberculosis* CDC1551 [18], *M. tuberculosis* 210 [18a], *M. microti* OV254 [19], *M. bovis* 2122 [20], and *M. bovis* bacille Calmette-Guerin (BCG) Pasteur [20a]. Mycobacteria sequenced or being sequenced outside the MTC include *M. leprae* [21], *M. ulcerans* [22], *M. avium* 104 [51], *M. paratuberculosis* K10 [22a], *M. marinum* [22b], and the relatively fast-growing *M. smegmatis* MC2 155 [18a]. Even the most distant of these sequenced mycobacterial genomes are minimally related by 60% DNA/DNA homology, and comparative genomic analysis has shown that gene loss is a significant part of the ongoing evolution of the slow-growing mycobacterial pathogens [23].

**Single-nucleotide polymorphisms**

Single-nucleotide polymorphisms (SNPs) can result in a silent amino acid substitution in which the protein coding sequence remains unchanged (synonymous) or can alter the protein-coding sequence (nonsynonymous) and hence act as a substrate for
### Table 2
Overview of mycobacterial genome sequencing projects

<table>
<thead>
<tr>
<th>Species</th>
<th>First author/date [reference]</th>
<th>Genome size (base pairs)</th>
<th>No. of protein-coding genes</th>
<th>G + C nucleotide content (%)</th>
<th>Insight from sequencing project</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>M. tuberculosis</em> CDC1551</td>
<td>Fleischmann, 2002 [18]</td>
<td>4,403,836</td>
<td>4249</td>
<td>65.6</td>
<td>Polymorphisms among <em>M. tuberculosis</em> strains more extensive than initially anticipated</td>
</tr>
<tr>
<td><em>M. tuberculosis</em> 210</td>
<td>The Institute for Genomic Research [18a]</td>
<td>4,400,000a</td>
<td>NA</td>
<td>NA</td>
<td>Describes hyper-virulence of ‘Beijing’ strain family?</td>
</tr>
<tr>
<td><em>M. bovis</em> 2122</td>
<td>Garnier, 2003 [20]</td>
<td>4,345,492</td>
<td>3951</td>
<td>65.6</td>
<td><em>M. bovis</em> is derivative compared to <em>M. tuberculosis</em></td>
</tr>
<tr>
<td><em>M. microti</em></td>
<td>Brodin, 2002 [19]</td>
<td>4,400,000a</td>
<td>NA</td>
<td>64.0a</td>
<td>Loss of RD1 contributed to attenuation of <em>M. microti</em></td>
</tr>
<tr>
<td><em>M. bovis</em> BCG Pasteur</td>
<td>Sanger Institute [20a]</td>
<td>4,083,000a</td>
<td>NA</td>
<td>NA</td>
<td>Describes live vaccine administered to humans</td>
</tr>
<tr>
<td><em>M. marinum</em></td>
<td>Sanger Institute [22b]</td>
<td>6,636,827</td>
<td>NA</td>
<td>65.73%</td>
<td>Describes causative agent of TB-like disease in fish and ‘fish tank granuloma’ of humans</td>
</tr>
<tr>
<td><em>M. ulcerans</em></td>
<td>Stinear, 2004 [22]</td>
<td>6,032,000a</td>
<td>NA</td>
<td>65.0a</td>
<td>Plasmid-encoded toxin responsible for Buruli ulcer</td>
</tr>
<tr>
<td><em>M. leprae</em></td>
<td>Cole, 2001 [21]</td>
<td>3,268,203</td>
<td>1604</td>
<td>57.8</td>
<td>Massive gene decay in the leprosy bacillus</td>
</tr>
<tr>
<td><em>M. avium avium</em> 104</td>
<td>Semret, 2004 [51]</td>
<td>5,475,491</td>
<td>4480</td>
<td>69</td>
<td>Extensive genomic polymorphism among <em>M. avium</em> sub-species</td>
</tr>
<tr>
<td><em>M. avium paratuberculosis</em> K10</td>
<td>GenBank [22a]</td>
<td>4,829,781</td>
<td>4350</td>
<td>69.3</td>
<td>Describes causative agent of Johne’s disease in cattle</td>
</tr>
<tr>
<td><em>M. smegmatis</em> MC2 155</td>
<td>The Institute for Genomic Research [18a]</td>
<td>7,000,000a</td>
<td>NA</td>
<td>NA</td>
<td>Describes fast growing, model organism for mycobacteria</td>
</tr>
</tbody>
</table>

Mycobacteria are listed in the order of 16S rRNA sequence relatedness to *M. tuberculosis* [64].

Abbreviations: G + C, guanine plus cytosine; NA, not available.

*a* Parameter estimates.
evolutionary selection. Both types of mutations have been applied toward differentiation and diagnostics of MTC members [24–26]. In a landmark study, a first sequence analysis of MTC isolates revealed that allelic polymorphism is impressively rare, occurring on the order of 1 in 10,000 base pairs (bp), suggesting that the complex could be dated to about 15,000 to 20,000 years of age [24].

Genomic comparison of multiple sequenced MTC strains has made possible the identification of SNP markers for studies of evolution, pathogenesis, and epidemiology in clinical M. tuberculosis [27] and M. bovis [20], supporting a clonal evolution of the MTC without detectable lateral gene exchange. The ratio of SNP types within a genome can act as a molecular clock [28] in which the high ratio of non-synonymous to synonymous mutations across coding sequences within MTC genomes suggests a recent divergence of M. bovis and M. tuberculosis [20].

Large-sequence polymorphisms

Unlike other mycobacterial species in which horizontal gene transfer has been demonstrated [22], this mode of generating genomic diversity has not been observed for the obligately intracellular MTC. Genomic comparisons for the MTC reveal a prominent role of genomic deletions relative to the sequenced strains of M. tuberculosis. For example, the complete genome sequence of M. bovis 2122 contains 66,037 bp less than M. tuberculosis H37Rv, and no genomic region exclusive to M. bovis but consistently absent from M. tuberculosis has been detected [20].

To uncover deletions in nonsequenced strains efficiently, one can hybridize whole genomic DNA of a MTC member against a spotted array [29] or an Affymetrix GeneChip (Santa Clara, California) [30] representative of the entire M. tuberculosis H37Rv genome [17]. Regions of the prototype strain that seem to be absent from the test strain are then confirmed by performing polymerase chain reaction (PCR) with primers approximating the deleted region, to amplify across the deletion. This amplicon is then sequenced to define the deletion point precisely. Isolates are said to share a genomic deletion when sequencing shows the deletion occurs in different isolates at exactly the same cut point [2]. Because independently arisen chromosomal rearrangements sometimes involve the same strategically located elements, only upon exact description of the specific genomic event (ie, genomic location within a reference strain) can one determine with confidence whether genomic deletions behave as unidirectional event polymorphisms (UEPs). These UEPs, which represent one-time events in the evolution of the organism, can serve as robust markers of clonal organisms, useful for determining phylogenetic classification.

A valuable use of these genomic deletions pertains to their application in defining specific MTC members and accurately assessing their prevalence in clinical specimens [8,26]. Unlike biochemical testing, which for individual results had imperfect sensitivity and specificity, the use of genomic events in these studies provided unambiguous classification, thereby simplifying the process considerably. To explore the basis for the previously observed biochemical attributes used for MTC speciation, an association was sought between deleted sequences and phenotypic results for isolates assigned as M. africanum. Results indicate that convergent biochemical profiles can be independently obtained in different MTC members. For instance, organisms presenting the distinct deletion profile of M. africanum and M. bovis can manifest the same biochemically based profile [7]. These results confirm the limitations of biochemically derived speciation and, by extension, challenge the taxonomic divisions currently in place for classifying members of the MTC.

Beyond diagnostics, different studies have all supported the potential value of most MTC genomic deletions (with the exception of mycobacteriophage DNA) as evolutionary markers. In separate genomic studies of M. bovis BCG vaccine strains, it has been documented that BCG-specific deletions superimpose perfectly on the historical record [3,29]. In studies of genomic deletions within clinical isolates of M. tuberculosis [31,32], mycobacterial clones shared the same genomic deletions, again suggesting that deletions can be used to reconstruct phylogenetic trees. Finally, a recent analysis of 100 M. tuberculosis clones from San Francisco has again confirmed that these deletions are UEPs [5], and therefore genomic deletions can effectively brand a particular clone [33]. A practical use of this approach will be to provide a secure genomic definition for prominent strains, such as the Beijing [34] and Manila [35] strains of M. tuberculosis, and to assess their prevalence through space and time.

Genomic deletions and the origin and evolution of the Mycobacterium tuberculosis complex

The long-recognized presence of a human TB bacillus and a closely related bovine form has given rise to speculation that TB originally came to humans as a zoonotic infection from cattle [36]. In
retrospect, this view was probably influenced by the types of *M. tuberculosis* isolates available for study, biased toward hosts (namely cattle and humans) for which a diagnosis of TB would lead to microbiologic investigation. To explore the evolutionary relationship of members of the MTC, the presence or absence of deletions was tested within complex isolates derived from different hosts and from isolates in various geographic locales [1,2]. Analysis revealed a stepwise accumulation of genomic deletions among isolates interrogated, but the distribution of genomic deletions argued against present-day *M. bovis* as the evolutionary precursor of *M. tuberculosis*, making it improbable that human TB originated with the domestication of cattle. Instead, a number of MTC organisms, both long established and more recently described, present genomic profiles that seem to be intermediate between the ancestor of modern *M. tuberculosis* and that of present-day *M. bovis*.

The availability of improved laboratory tools has facilitated the description of a number of novel variants of the MTC, including *M. canettii* [37,38], *M. caprae* [39,40], *M. pinnipedii* [41,42], and the dassie bacillus [2,43] (Table 1).

Before these tools were available, MTC members had presented a well-established host range, presumably biased by expectations: *M. tuberculosis* (and sometimes *M. africanum*) is classically isolated from humans, *M. microti* from voles, and *M. bovis* from a broad range of hosts including (but not limited to) cows. More careful study, however, has revealed a wider range of host animals. A practical issue arising from these studies involves the generally held belief that *M. bovis* infects an extensive range of animal species, including the badger, opossum, elk, cougar, and buffalo. Until recently, *M. caprae* and *M. pinnipedii* were considered to be forms of *M. bovis* [40,42]. Although *M. bovis* might be versatile enough to accommodate such a dynamic host range, the inclusion of such organisms probably overestimates the true host range of *M. bovis*. Detailed genomic analysis of isolates from unusual hosts is underway, with the expectation that results will continue to challenge accepted notions of MTC speciation and taxonomy [2].

Just as genomic deletions have proven unique to isolates of *M. tuberculosis* affecting only human hosts [5,30], deletions unique to these other MTC members permit resolution of their phylogetic situation (Fig. 1) [1,2]. A first observation from this distribution of organisms is that the MTC affects a number of undomesticated and domesticated mammals, both terrestrial and aquatic. *M. marinum* can be considered a piscine/amphibian mycobacterium [44] and *M. avium* an avian mycobacterium [45]. The MTC is a relatively broad-ranging mammalian mycobacterium. Organisms of the four most ancestral lineages (*M. canettii*, *M. tuberculosis*, and both genotypes of *M. africanum* subtype I) have been cultured predominantly from humans. Because isolation of *M. canettii* has been extremely rare [37,38], an unrecognized nonhuman reservoir might exist, with humans representing an accidental or circumstantial host. The next three MTC lineages (*M. microti*, *M. pinnipedii*, and the dassie bacillus) affect undomesticated mammals irrespective of their geographic location. *M. microti*, first identified in Europe, infects the field vole [19], the dassie bacillus infects the dassie and the surikat from Africa [6,43], and *M. pinnipedii* globally infects a variety of seals and sea lions from Oceania to South America [4,42]. Finally, more derivative forms of the MTC are seen in goats (*M. caprae*) and subsequently cattle (classic *M. bovis*), suggesting that the organism was introduced into livestock in the order of their domestication. More recently, spillover of *M. bovis* from farms has been seen in the case of badgers in the United Kingdom [46] and the brushtail opossum in New Zealand [46a]. Far from suggesting that human TB originated with livestock, the genomic record suggests that, directly or indirectly, humans were responsible for bringing MTC to the farm, with secondary foci of spread now observed in animals associated with this setting.

**Geographic, chronologic, and ecologic origins of the *Mycobacterium tuberculosis* complex**

An absolute chronology of the TB epidemic is difficult to discern by genomic deletions, because they do not evolve on a predictable time scale. The genetic record can potentially point to the geographic origins, however, because the ancestral form *M. canettii* [1,4,25] has been isolated only in persons living in Africa [37,38]. If the origins of human TB are situated in the same the continent as the origins of man, it is conceivable that the organism spread with humans during the paleomigration, explaining the presence of MTC DNA in 5000-year-old samples from Egypt [47] and pre-Columbian mummies from Ecuador [48]. Because more derivative organisms are found in hosts domesticated 10,000 to 12,000 years ago, these clues suggest that the organism accessed humans before that era and subsequently spread to other hosts, either from man directly or through an unrecognized vector.
Using deletions to assign directionality to the MTC phylogeny, one can employ sequence-based analysis to estimate the chronology of this scenario and refine the previous nucleotide-based analysis that suggested a 20,000-year divergence between *M. tuberculosis* and *M. bovis* [24]. Another approach to date these events uses testing for genomic regions directly on paleo-DNA samples [49] (Mostowy et al, unpublished data). Because these samples can be carbon dated independently, it is possible to provide genomic signatures for samples of human or non-human provenance and to derive minimal estimates for the ages of genomic events portrayed in Fig. 1.

Turning to the ecologic origins of the MTC, a livestock source seems to be unlikely, because human forms diverged before the modern caprine and bovine forms. Although it is attractive to consider another mammalian host as the ancestral niche, observations for other mycobacteria suggest that nonmammalian reservoirs such as plant or insects deserve consideration [50]. With the ability to test rapidly for genomic deletions by PCR, one can test putative wildlife reservoirs for variants of the MTC to find the natural host of relatively ancestral forms such as *M. canettii*.

**What is being deleted from the *Mycobacterium tuberculosis* complex?**

When compared with other bacterial species, members of the MTC present relatively little genomic diversity. Estimates of large-sequence polymorphism diversity among MTC members [32], in agreement with similar conclusions drawn from estimates of SNPs [25], have been consistently described as low in comparison with other microbes. Nonetheless,
Genomic flexibility does seem to exist within the MTC for specific host adaptation, and a similar potential is beginning to reveal itself among other mycobacterial complexes. Although the amount of diversity revealed within the Mycobacterium avium complex is 10-fold more than that of the MTC [51], host-specific genomic contents are being observed there as well (M. Semret and M. Behr, unpublished data). Taken together, these data highlight a comparative genomics approach to understanding an evolutionary potential of mycobacterium pathogenesis, in which genomic content can suggest DNA features for host-specific adaptation.

Genomic deletions and virulence

With host-specific MTC extending beyond a domesticated setting, how TB spreads from host to host is difficult to ascertain. From what is known about humans and cows with TB, it is reasonable to expect that transmission would occur through aerosols from a diseased animal to a contact animal. If so, a requisite of host adaptation is a certain degree of virulence in that host. Although greater virulence might facilitate transmission, too much virulence could be detrimental if host mortality is excessive or if the organism causes an invasive form of TB that is generally nontransmissible (such as TB meningitis). Thus, optimal transmissibility requires some degree of virulence (ie, pulmonary pathology) but a sufficiently contained disease process to generate the agents required for spread (ie, aerosols).

Support for this notion comes from studying the content of the genomic regions that have been deleted in different MTC members. A general observation is that although each deletion noted in Fig. 1 is unique to the bp, both genomic regions and the predicted function of implicated genes are nonrandom. Several regions of difference (RD) seem to be prone to genomic deletion, with different specific deletions having occurred near the same locus [6,7]. Most prominent among these is RD1, a series of nine genes implicated in the attenuation of M. bovis BCG strains, that has suffered three distinct genomic deletions. Although this confluence of deletions might point to genetic instability at this locus, a study of 100 circulating M. tuberculosis clones documenting 176 deletions failed to detect a single deletion in this region, arguing against an inherently elevated mutation rate [32]. The absence of RD1 was first observed for BCG vaccines [51a]; subsequently, targeted disruption of RD1 from M. tuberculosis was shown to decrease bacterial replication and educe pulmonary pathology in a mouse model [52,53]. More recently, M. microti and the dassie bacillus also were shown to have deletions in the RD1 region; notably, both have been characterized as having low virulence in animal models [19,43]. More detailed analysis of the RD1 region revealed it contains genes encoding a novel secretion system of two important secreted antigens (CFP-10, ESAT-6) [53,54]. Presumably a metabolically expensive process, the loss of this region in BCG was probably advantageous with no selective pressure in favor of synthesizing and secreting antigenic proteins in vitro. Although the independent loss of CFP-10 and ESAT-6 in M. microti and the dassie bacillus explains their attenuated phenotype, the selective pressures for their deletion in vivo are more speculative.

Given the documented impact of the RD1 region on virulence, the observation of its deletion in both the vole and dassie hosts is provocative. Nothing evident points to why the genetically distant vole (a rodent) and dassie (closely related to the elephant) would share some unique immunologic susceptibility. A more likely explanation might involve social conditions and transmissibility [55], given that voles and dassies congregate in high-density underground communities, unlike other MTC hosts that predominantly live aboveground in open-air conditions. Such congregate living settings would be extremely favorable for TB transmission, and an organism of lesser virulence might be successful in such burrowing hosts so long as host populations remain sufficiently abundant [56]. Conversely, conditions for transmission aboveground between goats and seals are less ideal and would probably require an organism of relatively high virulence to optimize transmissibility.

Summary of catalogued deletions

From Fig. 1, deletions represented along the vertical line of the phylogeny preceded spread of the bacillus into new hosts; those along the horizontal axes arose during coevolution of the organism with new hosts. Evidence supporting this scenario is that organisms lacking RD7, RD8, RD9, and RD10 have been recovered from the entire MTC host range, whereas the precise deletions seen along the horizontal lineages are observed in only restricted, one-host settings. To derive a scenario for the loss of genomic regions in vivo, genes lost on the vertical axis and those lost along horizontal lineages can be directly compared, pointing to nonrandom distinction between the functional classification of these two sets of deleted genes. Although such studies generate hypotheses regarding the evolution of MTC members in different hosts, these studies are naturally biased
to successful pathogenic strains, as opposed to those that caused infection but not disease. In this light, an examination of BCG vaccines provides telling insights into MTC evolution when the selective pressures for virulence were absent in the host and limited to the culture media employed. Remarkably, the sheer volume of genomic disturbance incurred by BCG vaccines during a half-century of laboratory evolution has favored the elimination of regulatory elements and antigens. These results reinforce the notion that the capacity to engage the host immune system is selected for during in vivo conditions, consistent with MTC members being professional pathogens [57]. More practically, the absence of numerous antigens from BCG vaccines may in part explain its limitations as an immunizing agent [58].

Summary and concluding thoughts: lessons for tuberculosis control

*M. tuberculosis* has probably been with humans for millennia and thus probably became adapted to humans during times of low population density and predominantly outdoor living. Unlike diseases such as HIV that rapidly spread in epidemic form soon after introduction into humans, the TB epidemic peaked in Western Europe during the nineteenth century and has arguably yet to peak in certain parts of the world. Although it is possible that the organism has evolved toward greater virulence in recent centuries, it seems more probable that that social changes brought about by industrialization were paramount in altering the transmission dynamics. This argument would also apply to *M. bovis*, in which an organism that evolved to persist in free-ranging cattle would predictably wreak havoc in the environment provided by modern-day factory farming. Finally, whereas tuberculous animals in the wild might normally succumb to predation, the increasing protection of these hosts in wildlife refuges and zoos should provide a greater chance for progression to TB, as attested to by reports of TB in farmed deer [59], zoo tigers [60], and seal-trainers [61]. Although these cases are generally rare, these anecdotes do serve notice that MTC has the capacity to adapt to the immunologic environment it engages and suggest that nonhuman reservoirs may become pertinent to human TB control.

References


