Reducing the Activity and Secretion of Microbial Antioxidants Enhances the Immunogenicity of BCG

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Abstract

Background: In early clinical studies, the live tuberculosis vaccine Mycobacterium bovis BCG exhibited 80% protective efficacy against pulmonary tuberculosis (TB). Although BCG still exhibits reliable protection against TB meningitis and miliary TB in early childhood it has become less reliable in protecting against pulmonary TB. During decades of in vitro cultivation BCG not only lost some genes due to deletions of regions of the chromosome but also underwent gene duplication and other mutations resulting in increased antioxidant production.

Methodology/Principal Findings: To determine whether microbial antioxidants influence vaccine immunogenicity, we eliminated duplicated alleles encoding the oxidative stress sigma factor SigH in BCG Tice and reduced the activity and secretion of iron co-factored superoxide dismutase. We then used assays of gene expression and flow cytometry with intracellular cytokine staining to compare BCG-specific immune responses in mice after vaccination with BCG Tice or the modified BCG vaccine. Compared to BCG, the modified vaccine induced greater IL-12p40, RANTES, and IL-21 mRNA in the spleens of mice at three days post-immunization, more cytokine-producing CD8+ lymphocytes at the peak of the primary immune response, and more IL-2-producing CD4+ lymphocytes during the memory phase. The modified vaccine also induced stronger secondary CD4+ lymphocyte responses and greater clearance of challenge bacilli.

Conclusions/Significance: We conclude that antioxidants produced by BCG suppress host immune responses. These findings challenge the hypothesis that the failure of extensively cultivated BCG vaccines to prevent pulmonary tuberculosis is due to over-attenuation and suggest instead a new model in which BCG evolved to produce more immunity-suppressing antioxidants. By targeting these antioxidants it may be possible to restore BCG’s ability to protect against pulmonary TB.

Introduction

Oxidants fulfill signaling roles during the activation of innate immune responses and promote the development of adaptive immunity [1–3]. Growing evidence implicates microbial antioxidants in the suppression of host immune responses to Mycobacterium tuberculosis. The iron co-factored superoxide dismutase (SodA) and its secretion channel, SecA2, suppress macrophage activation in vitro and inflammatory and immune responses in vivo [4–6]. An extracytoplasmic-function sigma factor, SigH, controls the expression of several antioxidants including thioredoxin and is associated with lung immunopathology [7–9]. Other antioxidants including catalase-peroxidase, alkathydroperoxidase C, and the NADPH quinone reductase Rv3303c also contribute to mycobacterial virulence [10–12].

The current vaccine against tuberculosis (TB), the bacillus of Calmette and Guérin (BCG), also secretes large amounts of antioxidants. BCG is widely administered however it has failed to control the enormous global burden of TB that causes about 9 million new active cases and 1.7 million deaths annually [13–16]. BCG is a live vaccine derived from M. bovis via the loss of RD1, the “region of deletion 1” that encodes the ESX-1 secretion system.
involved in virulence [17]. During the first half of the 20th century BCG was sent to investigators throughout the world and maintained by serial passage, i.e., by transferring a portion of an aging culture into fresh media. Over decades this practice produced multiple BCG daughter strains with additional regions of genomic deletion as well as regions of genomic duplication and other mutations [17–19]. These substrains represent divergent evolution from the original BCG and some have been transferred over 1500 times [17]. Some BCG daughter strains exhibit genomic duplication of sigH, trxC (thioredoxin), trxB2 (thioredoxin reductase), whiB1, whiB7, and lpdA (Rv3303c) as well as increased expression of genes encoding other antioxidants including SodA, thiol peroxidase, alkali-hydroperoxidases C and D, and other members of the whiB family of thioredoxin-like protein disulfide reductases [17,19–21].

Although attenuated from the loss of RD1, some BCG substrains exhibit potent immune suppressive properties. BCG inhibits phagolysosomal fusion in macrophages [22], reduces monocyte apoptosis [23], down-regulates MHC class II molecules [24], and suppresses the maturation of dendritic cells [25]. The immune suppressive capacity of BCG is perhaps most apparent in vivo. Some BCG daughter strains persist indefinitely in small animals [4,26–28], induce lung pathology [4,29], and cause death when administered as a large dose [30]. In man, the rising prevalence of HIV infection and virulence attributes of BCG have raised concerns about disseminated BCG infection after routine neonatal vaccination and it is estimated that more than 2000 serious complications of BCG vaccination occur annually [31].

To test the hypothesis that the duplication of SigH and production of SodA contribute to the immune suppressive capacity of BCG, we modified an extensively cultivated BCG daughter strain to eliminate both copies of sigH and to reduce the activity and secretion of SodA. The modified vaccine induced stronger immune responses than the parent BCG vaccine during primary vaccination as well as enhanced recall immune responses upon subsequent challenge. These observations have important implications for efforts to develop a more effective live vaccine against TB. They further suggest the novel hypothesis that the extensively cultivated BCG daughter strains are immune suppressive as a consequence of increased antioxidant production. This contrasts with the theory that the extensively cultivated BCG daughter strains are over-attenuated although both theories attempt to explain the apparent decline in efficacy of BCG against pulmonary TB during decades of cultivation in vitro [32,33]. Finally, the findings highlight the importance of host-generated oxidants in the development of adaptive immunity.

Results

Construction and in vitro characterization of 3dBCG, a modified BCG vaccine

We used allelic inactivation to eliminate secA2 and sigH in BCG Tice, and dominant-negative (dn) interference techniques to reduce the activity of SodA. To construct BCGΔsecA2ΔsigH (“double-deletion BCG”, “DDBCG”) secA2 and sigH were inactivated in sequence (Fig. 1A). During attempts to inactivate sigH with a hygromycin-resistance (hygR) cassette we found that only about 15% of colonies lacked sigH whereas most colonies contained an intact sigH along with DNA restriction fragments suggesting successful interruption of sigH by the hygR allele. This finding is consistent with reports that most BCG daughter strains have two copies of sigH due to tandem duplication of the region of chromosomal DNA containing sigH [17,19].

After verifying that sigH was absent from the colony selected for further modification, we inserted an allele encoding a dominant-negative mutant of SodA into the chromosome of DDBCG to yield 3dBCG (BCGΔsecA2ΔsigHΔsodA). SodA contains histidines that chelate the active-site iron [34] and we eliminated H28 and H76 to produce an inactive SodA mutant. Enzyme activity assays (Fig. 1B) and immunoblotting (Fig. 1C) demonstrated a dominant-negative effect with reduced enzyme activity in 3dBCG compared to DDBCG despite comparable amounts of SodA protein, presumably because AH28AH76 monomers were incorporated into the tetrameric, functional form of SodA and inhibited enzyme activity. Compared to BCG, the inactivation of secA2 in DDBCG nearly eliminated SodA secretion however enzyme

Figure 1. Construction of modified BCG vaccines. (A) Southern hybridization demonstrating inactivation of secA2 and sigH. Drall-digested DNA from each strain was probed for secA2, sigH and the hygR cassette used to inactivate sigH. Lane 1, BCG Tice; lane 2, BCGΔsecA2; lane 3, BCGΔsigh; lane 4, BCGΔsecA2Δsigh (DDBCG); lane 5, a colony of BCGΔsecA2 with only one copy of sigh inactivated. secA2 contains two internal Drall sites and Drall-digested DNA was predicted to yield fragments of 307 bp, 785 bp, and 10544 bp. sigh was predicted to be on a 2895 bp fragment. The hygR cassette contains an internal Drall site and was predicted to yield 2938 bp and 1612 bp fragments after inactivation of sigh. (B) Differences in the localization and activity of SodA in BCG, DDBCG, and 3dBCG. A representative experiment shows the SOD units per ml of concentrated supernatant or lysate. (C) Immunoblot showing comparable amounts of SodA in lysates of DDBCG (lane 1) and 3dBCG (lane 2), despite the marked difference in SOD activity as shown in (b).

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activity was present in the lysate. Compared to DDBCG, 3dBCG exhibited a 93% reduction in enzyme activity. In summary, we introduced three modifications into BCG to reduce antioxidant production, yielding 3dBCG.

3dBCG induces stronger recall T cell responses

Most vaccines are administered to generate memory lymphocytes that can proliferate rapidly, thereby accelerating the kinetics and magnitude of immune responses to a subsequent infection. To determine whether vaccination with 3dBCG or BCG induce recall immune responses, we immunized C57Bl/6 mice subcutaneously with BCG, 3dBCG, or sterile media (control vaccination group). BCG typically can be recovered months after inoculation of C57Bl/6 mice depending upon the strain, dose, and route of administration [4,26,27], however BCG-induced T cell responses were induced rapidly almost to pre-vaccination levels within a few months of treating vaccinated mice with antibiotics [35]. Thus, to assess recall immunity without interference from persisting vaccine bacilli, we treated the vaccinated groups of mice with isoniazid and rifampin from day 30 to day 60 post-vaccination and this reduced the spleen titer of vaccine bacilli to below the lower limits of detection (50 CFU). After resting the mice for an additional 2 to 9 weeks, they were challenged intravenously with 2×10⁵ CFU of BCG Tice.

Recall immunity was evaluated on days 26 to 30 post-challenge by incubating splenocytes on uninfected and BCG-infected bone marrow-derived macrophages (BMDMs). Antigen-specific cytokine production, quantified by intracellular cytokine staining and flow cytometry (Fig. 2A), was reported as the number of cytokine+ T cells per spleen. The magnitude of the CD4+ IFN-γ T cell response following challenge in BCG-vaccinated mice (median 67,560/spleen) was not significantly greater than in media-vaccinated control mice (median 43,300/spleen) undergoing a primary response to the BCG challenge (Fig. 2B). In contrast, more IFN-γ+ CD4+ T cells were present in 3dBCG-vaccinated mice (median 138,500/spleen). The number of CD4+ lymphocytes producing TNF-α or both IFN-γ and TNF-α was also greater in 3dBCG-vaccinated mice than in BCG-vaccinated or control mice.

The production of IL-2 is associated with multi-functional, high-quality memory T cells [36] and thus as a secondary endpoint we also evaluated the IL-2 recall response. The number of IL-2+ and IFN-γ+IL-2+ (double-positive) CD4+ T cells in BCG-vaccinated mice was not significantly greater than the response of control mice (Fig. 2B). In contrast, more IL-2+ and IFN-γ+IL-2+ CD4+ T cells were present in 3dBCG-vaccinated mice than in control mice. For the comparisons involving 3dBCG- and BCG-vaccinated mice, there was a trend towards statistical significance (P>0.05, <0.1). The numbers of IFN-γ- or TNF-α-producing CD8+ T cells in the spleen were low at day 26 to 30 post-challenge (data not shown) compared to the number of CD4+ T cells producing these cytokines. However, five of nine 3dBCG-vaccinated mice exhibited greater than 10,000 IL-2+ CD8+ T cells in the spleen compared to none of seven media-vaccinated mice (P=0.03, Fisher’s exact, two-tailed) and one of nine BCG-vaccinated mice exhibited greater than 10,000 IL-2+ CD8+ T cells.

After noticing higher cytokine production in re-stimulated splenocytes of 3dBCG-vaccinated mice in the first experiment, we enumerated challenge bacilli in spleins of mice in subsequent experiments and thus determined post-challenge CFU values for six vaccinated mice per group. Fewer bacilli of the BCG challenge dose were found in the spleens of 3dBCG-vaccinated mice than in BCG-vaccinated mice (median Log_{10} CFU of 5.05 and 5.80, respectively, P=0.034) or media-vaccinated mice (median Log_{10} CFU 5.95, P=0.013) (Fig. 3A). An inverse correlation between median CFU and median CD4+ IFN-γ T cell responses from each of the three vaccinated groups was observed (R² = 0.99) (Fig. 3B), indicating that the stronger 3dBCG-induced recall CD4+ IFN-γ responses were associated with better clearance of the challenge bacilli. Therefore, under circumstances in which the live vaccine strain is not allowed to persist, prior vaccination with 3dBCG enhanced BCG-specific immune responses post-challenge and reduced the number of challenge bacilli, whereas prior vaccination with BCG did not confer much advantage over unvaccinated mice.

3dBCG induces stronger CD8+ T cell responses during primary vaccination

Results from the recall immune response experiments above led us to evaluate primary T cell responses in more detail. We used Balb/c mice to track peptide-specific responses to immunodominant epitopes including TB10.474-88 - ST15, a CD4 epitope [37], and TB10.3/10.4_20-28 - GL9, a CD8 epitope [38]. Preliminary experiments showed comparable numbers of cytokine-producing CD4+ T cells in mice vaccinated subcutaneously and intravenously, however the intravenous group demonstrated greater CD8+ T cell responses by flow cytometry (data not shown). Thus, to determine the kinetics and peak of the primary response we inoculated Balb/c mice intravenously with 5×10⁵ CFU of BCG or 3dBCG and analyzed the number of IFN-γ-producing CD8+ and CD4+ T cells at 8, 9, 12, 17 and 30 days post-inoculation. The CD8+ T cell response to peptide GL9 peaked at day 9 post-inoculation (Fig. 4A) and the CD4+ T cell response to peptide ST15 peaked at day 12 (Fig. 4B). The kinetics of the T cell responses to BCG and 3dBCG vaccination were similar however vaccination with 3dBCG induced higher peak IFN-γ+, IFN-γ+TNF-α and IFN-γ+IL-2+ CD8+ T cell responses to GL9 than vaccination with BCG (Fig. 4C).

3dBCG induces more IL-2-producing CD4+ T cells

The presence of IL-2 during the contraction phase of the primary immune response is critical for the development of memory T cells that can subsequently proliferate to provide a recall T cell response [36,39,40]. The kinetics of the immune response after subcutaneous vaccination of Balb/c mice with 5×10⁵ CFU of BCG or 3dBCG demonstrated declines in the number of IFN-γ+ CD4+ T cells and increases in the number of IL-2+ CD4+ T cells in response to peptide ST15 from day 17 to day 30 post-vaccination (Fig. 5A,B). The number of ST15-specific IFN-γ- or TNF-α-producing CD4+ T cells at the peak (day 17) or memory phase (day 30) CD4+ T cell response was similar in the BCG and 3dBCG groups. In contrast, vaccination with 3dBCG induced more ST15-specific IL-2+ CD4+ T cells during the memory phase (median 18,400/spleen) than vaccination with BCG (median 10,760/spleen) (Fig. 5C). The higher ratio of IL-2+ to IFN-γ+ or TNF-α+producing CD4+ T cells in 3dBCG-vaccinated mice compared to BCG-vaccinated mice may have contributed to the greater recall immune response in 3dBCG-vaccinated mice during subsequent challenge.

3dBCG induces greater early expression of IL-12p40, RANTES, and IL-21 mRNA

The cytokine milieu after vaccination influences the differentiation and polarization of naïve CD4+ T cells, IFN-γ, IL-12, and RANTES (CCL5) promote a Th1 response, IL-1α Th2 response, IL-6 a Th17 response, and IL-21 a T follicular helper response [41–44]. To determine whether BCG and 3dBCG induce different patterns of expression of cytokines that may explain some of the differences in adaptive immunity reported above, we inoculated C57Bl/6 mice intravenously with...
1.5 × 10^7 CFU of BCG, 3dBCG, or PBS (control). Spleens were harvested 72 hours later and the mRNA from whole spleens was evaluated by RT-PCR for these Th-polarizing cytokines. BCG- and 3dBCG-vaccinated mice exhibited increased expression of IFN-γ, IL-12p40, IL-6, and IL-21 compared to control mice (Fig. 6) whereas the expression of IL-4 was not significantly different in either vaccinated group or controls. The expression of some genes was significantly greater in 3dBCG- than BCG-vaccinated mice including 33% more IL-12p40 (IL-12b), 34% more IL-21, and 30% more RANTES, which was suppressed in BCG-vaccinated mice compared to the control and 3dBCG groups. In summary, within three days of inoculation, BCG and 3dBCG induce different patterns of expression of immune response genes likely to influence subsequent T cell responses.
Figure 3. Spleen mycobacterial titers post-challenge and correlation between bacilli counts and T cell responses. (A) BCG colony-forming units (CFU) persisting in the spleens of mice on day 26–30 after intravenous challenge with $2 \times 10^7$ CFU of BCG. Data are from six mice per group from two experiments. Individual and median values are displayed. The variance of the medians of the three groups was significant, $P = 0.0186$, Kruskal-Wallis test. Mann-Whitney $P$ values for two-group comparisons are shown in the figure. (B) Plot of median Log$_{10}$ CFU in spleens of vaccinated mice versus the median CD4+ IFN-$\gamma$ responses for the three vaccination arms. Coefficient of correlation, $R^2$ was calculated using median values from each vaccination group.

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Figure 4. Kinetics of the primary immune response and peak CD8+ T cell responses. (A) Numbers of IFN-$\gamma$+ CD8+ T-cells in spleens after re-stimulation with TB10.3/10.4,20-28 peptide (GL9) at 8, 9, 12, 17 and 30 days after intravenous inoculation of media, BCG or 3dBCG. The symbols and error bars represent median and range of responses of 3 to 9 mice per time point. (B) Numbers of IFN-$\gamma$+ CD4+ T cells in spleens after re-stimulation with TB10.4,20-82 peptide (ST15) on days after vaccination. (C) Numbers of IFN-$\gamma$+, and IFN-$\gamma$+TNF-$\alpha$+, and IFN-$\gamma$+IL-2+ CD8+ T cells in spleens after re-stimulation with TB10.3/10.4,20-28 peptide - GL9 at the peak of the primary CD8+ T cell response (day 9 post-inoculation). Each symbol represents the number of cytokine-producing CD8+ T cells from one mouse. The bars represent median values for each group and Mann-Whitney $P$ values for two-group comparisons are displayed in the figure. The variances of the medians of the three groups were significant: IFN-$\gamma$, $P = 0.0009$; IFN-$\gamma$+TNF-$\alpha$, $P = 0.0013$; and IFN-$\gamma$+IL-2, $P = 0.0415$, Kruskal-Wallis test.

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3dBCG grows slower in vitro and persists less well in vivo

We have previously reported that under culture conditions wherein a detergent (e.g., Tween 80) is used to reduce the clumping of bacterial cells, antisense mutants of M. tuberculosis H37Rv with diminished production of SodA grow slower than the parent H37Rv strain [4]. Furthermore, these SodA mutants are attenuated in their ability to cause death and to persist in vivo [4]. Attenuation has also been reported for secA2 deletion mutants of M. tuberculosis [5]. To determine whether the multiple mutations in antioxidant production introduced into BCG similarly affects in vitro growth and in vivo persistence, we monitored the absorbance of broth cultures of BCG and 3dBCG over time and compared the number of viable bacilli in the spleens of Balb/C at four weeks after intravenous inoculation. Relative to BCG, 3dBCG exhibited about 30% less of an increase in absorbance over ten days of cultivation (Fig. 7A), indicating a modest reduction in its rate of growth. Furthermore, the number of 3dBCG recovered from the spleens of mice was reduced 32-fold compared to BCG (Fig. 7B). The recovered bacilli represented 1.8 percent and 38.6 percent, respectively, of the number of bacilli inoculated (P = .027, two-sample T test). Thus, the stronger immune responses induced by 3dBCG compared to BCG as demonstrated above cannot be attributed to an effect of higher numbers of persisting vaccine bacilli. Instead the reduced persistence of 3dBCG in vivo is a consequence of the stronger immune responses induced by 3dBCG along with its slower rate of growth.

Discussion

The capacity of M. tuberculosis and M. bovis to secrete multiple antioxidants may have evolved to enable growth under aerobic conditions by reducing oxidative damage to cell wall lipids and proteins, which comprise nearly 40 per cent of the dry weight of mycobacteria. However, beyond their role in microbial physiology, the production and secretion of large amounts of antioxidants gives the bacterium obvious advantages within mammalian hosts. Although best known for their microbicidal properties, oxidants produced by immune cells also help to coordinate the host response to infection. Oxidants have important signaling roles in the activation and apoptosis of macrophages [1], the maturation of dendritic cells [2], and the proliferation of T cells [3]. Chronic granulomatous disease (CGD), a genetic disease in which the phagocyte oxidase fails to assemble and produce superoxide, causes defects in signaling cascades that impair phagocyte activation [45], antigen presentation [46], and the development of memory lymphocytes [47]. By analogy, M. tuberculosis secretion of SodA, an enzyme that inactivates superoxide, may induce a similar albeit localized deficiency of superoxide that inhibits the development of adaptive immunity, resulting in chronic infection rather than eradication of the pathogen. Granulomatous pathology is a hallmark of both CGD and TB and the pathophysiology of inflammation in CGD involves impairment of a superoxide-dependent step in tryptophan metabolism [48]. In summary, the inactivation of host-generated oxidants by mycobacterial antioxidants may disrupt redox signaling during early infection and...
Figure 6. Expression of genes encoding Th polarizing cytokines. Mice were inoculated by the retro-orbital route and spleens harvested 72 hours later. Each data point represents the level of gene expression by individual mice in each group as determined by RT-PCR and normalized to the mean value of the PBS-vaccinated group. The bars represent median values for each group and Mann-Whitney P values for two-group comparisons are displayed in the figure. Statistical comparisons of the variances of the medians of the three groups were: IL-12b, P = 0.0006; IFN-γ, P = 0.0033; IL-4, P = 0.0702; IL-21, P = 0.0011; IL-6, P = 0.0033; and RANTES, P = 0.0112, Kruskal-Wallis test.
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Figure 7. In vitro growth and in vivo persistence of BCG and 3dBCG. (A) Plot of the absorbance of cultures of BCG and 3dBCG during ten days of cultivation. A representative experiment shows A₆₀₀ values every two days after Middlebrook 7H9 media containing Tween 80 in roller bottles was seeded with BCG or 3dBCG. The initial inoculum was adjusted to an A₆₀₀ value of 0.2. (B) Spleen CFU at four weeks after intravenous inoculation of Balb/C mice with BCG or 3dBCG. Inocula were prepared based on absorbance values and then titered to determine the number of viable bacilli administered to each group of three mice – 6.6 x 10⁶ CFU for BCG; 7.7 x 10⁶ CFU for 3dBCG. The numbers represent the weights of the spleens in grams.
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promote TB pathogenesis by weakening host immune responses and promoting tissue-damaging immunopathology.

By reducing antioxidant activity and secretion in BCG to yield 3dBCG, we unmasked immune responses during vaccination with 3dBCG that were suppressed by the parent BCG vaccine. Enhanced cytokine responses were detected within 3 days of vaccination. These differences may have clinical relevance as IL-12 and RANTES contribute to protection against TB in man [49,50]. The early differences in cytokine responses were followed by differences in T cell responses. IL-12 and RANTES augment Th1 responses [41,42] and IL-21 and IL-12 promote the expansion of CD8+ T cells [51,52]. Thus the greater expression of IL-12p40, RANTES and IL-21 observed on day 3 post-vaccination in the 3dBCG group may have contributed to stronger antigen-specific IL-2+ production by CD4+ T cells as well as stronger CD8+ T cell responses. The enhanced CD8+ responses in 3dBCG-vaccinated mice may also have clinical relevance as CD8+ T cells contribute to protection against pulmonary TB in man [53,54]. In effect, the modifications to BCG correct a deficiency in vaccine immunogenicity that may partly be responsible for the suboptimal efficacy of BCG against pulmonary TB.

Another deficiency of some current BCG vaccines is the poor induction of immune memory. The duration of protection conferred by vaccination with BCG in mice appears to be limited to the time that BCG persists in vivo [55]. However, persisting live vaccine bacilli represent a risk for disseminated BCG infection, especially in persons infected with HIV [31]. Thus, modifications to BCG that promote the development of immune memory and confer protection independent of persisting vaccine bacilli should improve the safety of BCG. In summary, the differences in the immune responses of 3dBCG- and BCG-vaccinated mice suggest that reducing the activity and secretion of microbial antioxidants in BCG leads to greater activation of innate immunity, stronger antigen-specific T cell responses, and better memory immunity. We suspect there are also other differences in response to these vaccines and this is a focus of ongoing investigation.

The enhanced immunogenicity of 3dBCG compared to BCG challenges the hypothesis that extensively cultivated BCG daughter strains are over-attenuated [32,33]. To briefly review the background of the over-attenuation hypothesis and its influence on BCG vaccination practices, as early as the late 1940s concerns were raised about the possible over-attenuation of BCG [55]. The context was the observation of highly variable protection, from 0% to 80%, against pulmonary TB in human studies of BCG vaccination [13,14]. Furthermore, investigations in mice indicated that BCG daughter strains differed in their ability to persist in vivo and to confer protection against challenge by virulent M. tuberculosis [27,56,57]. Indeed, protection in mice correlated directly with the invasiveness and persistence of the BCG daughter strain [57]. Such observations, along with the assumption that virulence would be lost rather than gained during cultivation in vitro, led to the conclusion that virulent BCG substrains should be preferred as they were the most like the original protective vaccine [30] that had exhibited 80% protective efficacy against pulmonary TB in uncontrolled studies of nursing and medical students in the 1920s [30], and also in an early controlled clinical trial in the 1930s [39].

The opinion that virulent BCG daughter strains were the most like the original vaccine influenced the selection of the relatively virulent BCG Danish 1331 and BCG Pasteur 1173P2 vaccines for the large controlled trial of vaccination with BCG in the Chingleput region of South India between 1968 and 1971 [13,60,61]. However neither vaccine was effective in persons over 14 years of age. Although this failure is often attributed to a confounding effect by environmental mycobacteria found in tropical climates [14], this explanation ignores the fact that in an earlier trial in the Madanapalle region of South India, only 105 kilometers from Chingleput, BCG exhibited 60% protective efficacy with an annual rate (per 100,000) of 190 and 79 bacillary cases, respectively, among controls and BCG vaccinees between 25 and 45 years of age [62]. The protective vaccine used in the Madanapalle study came from a 1948 subculture of BCG Danish before it was freeze-dried in 1960 as BCG Danish 1331 [63]. Another clinical trial in the United Kingdom also used BCG Danish vaccine prepared between 1950 and 1952 to immunize 14 to 15 year-olds, finding a 77% reduction in pulmonary TB [64]. Thus, the clinical data indicate that at one time some BCG vaccines conferred protection against pulmonary TB in adolescents and adults, including tropical countries with a high prevalence of TB and environmental mycobacteria. However, as evidenced by the experience with BCG Danish, something happened during years of in vitro cultivation to make BCG become less effective against pulmonary TB. Yet as BCG Pasteur 1173P2 and Danish 1331 are relatively virulent BCG substrains [27,30,56,61], their failure to protect against pulmonary TB at Chingleput and in subsequent case-control trials where other less virulent BCG daughter strains exhibited about 55% protective efficacy against pulmonary TB [65–67] should not be attributed to over-attenuation.

Apart from its failure to explain BCG’s unreliable protection against pulmonary TB, the over-attenuation hypothesis is inconsistent with the observation that the “late” BCG vaccines Pasteur 1173P2 and Danish 1331 cause more adverse reactions in man and exhibit greater virulence in animal models than the “early” BCG Tokyo 172 vaccine strain [14,28,61,68]. Although the greater virulence of late BCG daughter strains that were passaged more than 1000 times compared to an early BCG vaccine passed only 172 times after it was brought to Japan by Kiyoshi Shiga in 1924 could be a coincidence of divergent evolution, observations involving vaccines representing “linear” evolution in the BCG Danish family similarly suggest that over time BCG became more virulent instead of more attenuated. In 1954, a few years after BCG Danish was the source of vaccine for the trials in Madanapalle and the United Kingdom in which protection against pulmonary TB was demonstrated, a freeze-dried seed lot from the 1077th transfer of BCG Danish was preserved as BCG London/Glaxo [63]. Six years later BCG Danish 1331 was preserved [63]. Yet despite its later derivation, BCG Danish 1331 is more virulent than BCG London/Glaxo in animal models. In one study the intraperitoneal injection of a 5 mg dose of Danish 1331 killed male golden hamsters within 100 days whereas more than 50% of hamsters infected with the same amount of BCG London/Glaxo survived for more than 400 days [30]. Experiments to determine the minimum “sensitizing” dose, i.e., the lowest number of bacilli capable of inducing delayed type hypersensitivity in guinea pigs or bank voles show that it takes only 10 cfu of Danish 1331 compared to about 200 cfu of BCG London/Glaxo [61,69]. Thus between transfer 1077 and transfer 1331, the BCG Danish family of vaccines became more invasive and lethal.

When combined with recent reports of the duplication and increased expression of genes encoding antioxidants in BCG [17,19], our results suggest a new model to explain how BCG became more virulent in small animals and less effective in controlling pulmonary TB in man. Whereas reference isolates of M. bovis and BCG Tokyo 172 have a single copy of sigH, BCG Pasteur 1173P2 and Danish 1331 contain bacilli with duplicated or triplicated sigH [17]. Indeed, BCG Danish appears to be
both early and late BCG daughter strains remain highly efficacious against TB meningitis and miliary TB in young children, and it is pulmonary TB that causes the greatest burden of disease globally [16]. Our results suggest it might be possible to make improvements in BCG that enhance protection against pulmonary disease in man. Molecular genetic techniques make it possible to reduce the activity of antioxidants below the level found in any BCG vaccine, past or present, and this may enable the construction of highly immunogenic BCG-derived vaccines that match or surpass the high protective efficacy against pulmonary TB observed in early clinical trials. Our findings demonstrate the value of targeting SigH and SodA in BCG. Other antioxidants that are more highly expressed in BCG daughter strains than M. bovis include thiol peroxidase, Rs3303c, and members of the WhiB family of protein disulfide reductases [17]. If these antioxidants similarly suppress host immune responses they may also be promising targets for enhancing BCG immunogenicity.

Beyond the implications of these findings for single dose vaccination with BCG to prevent pulmonary TB, a more immunogenic BCG should be a better priming vaccine for subsequent boosting with a heterologous vaccine in accordance with new TB vaccination strategies [75]. A more immunogenic BCG also has implications for cancer as increasing host production of multiple cytokines with anti-tumor activity (i.e., IL-12p40, RANTES, IL-21, and IL-2) has the potential to overcome some of the limitations of current BCG vaccines as cancer immunotherapy [76,77]. Finally, BCG occupies a unique place in the vaccination schedule of infants throughout much of the world. Thus, a more immunogenic BCG may become a useful vector for expressing antigens of other pathogens including HIV and Plasmodium species to produce strong cell-mediated responses, including CD4+ helper responses, as part of vaccination regimens against AIDS, malaria, and other infectious diseases.

Materials and Methods

Bacterial isolates, plasmids, media, and growth conditions

Genetic tools and bacterial isolates are listed in Table S1. Genetic modifications were made to BCG Tice (Organon Teknika Corp., Durham, NC). Of note, although historically BCG Tice was considered to be a weak BCG vaccine that exhibited poor ability to persist in mice or confer protection against challenge with M. tuberculosis in mice [27,57], it was reformulated with a subculture of BCG obtained from the Institut Pasteur in 1951 and subsequently BCG Tice has exhibited in vivo characteristics similar to BCG Pasteur 1173P2 [68,78,79].

E. coli strains were grown in LB media. To prepare vaccine inocula and to compare rates of growth, BCG Tice and modified strains were grown in roller bottles containing Middlebrook 7H9 media with 10% oleic acid-dextrose-catalase (OADC) enrichment supplemented with 0.2% glycerol, and 0.05% Tween80. Kanamycin (50 µg/ml or 25 µg/ml), apramycin (50 µg/ml), and hygromycin B (100 µg/ml or 50 µg/ml) were used to select colonies after genetic manipulations in E. coli or BCG, with higher concentrations for E. coli and lower concentrations for BCG.

Molecular genetic manipulations

Plasmid and chromosomal integration vectors for allelic inactivation or expression of dominant-negative monomers of SodA were electroporated into BCG using standard methods [80]. After electroporation, the strains were incubated in Middlebrook 7H9 media with 5% CO2 at 37°C for 24 hrs before the suspension was selected on Middlebrook 7H11 agar containing antibiotics as...
needed. Successful transformation was confirmed by PCR of DNA unique to the vector.

The inactivation of secA was performed using a two-step allelic exchange strategy with plasmid pMB179, as previously described in *M. tuberculosis* [5]. The inactivation of sigH was performed by using the cosmids-plasmid pYUB83-4-phAE87 system, containing the higR cassette, to construct a specialized transducing phage [80]. Homologous recombination in BCG was predicted to eliminate the terminal 316 nucleotides of sigH including the region encoding a putative DNA binding motif. Southern hybridization of *DraIII*-digested DNA was used to verify allelic inactivation.

To construct dominant-negative (dn) enzyme monomers, *sodA* was PCR-amplified from DNA of *M. tuberculosis* strain H37Rv, ligated into pcR2.1-TOPO, and propagated in *E. coli* TOP 10. Site-directed mutagenesis was performed using primer overlap extension methods [81] and codon deletion was verified by DNA sequencing. The AH28AH76 mutant was verified to lack superoxide dismutase (SOD) activity by its inability to complement the growth of the SOD double-negative (*sodA-* , *sodB-* ) *E. coli* strain CK9C1891 [82]. The dn-*sodA* open reading frame was then ligated behind DNA from a 341-bp region upstream of isocitrate lyase (*icl*) from H37Rv, a macrophage-inducible promoter [83], in an effort to not severely impair the growth of SodA-diminished strains in vitro while reducing enzyme activity in vivo. Finally, alleles encoding dominant-negative enzyme monomers were ligated into shuttle plasmid or chromosomal integration vectors.

**Assays of SodA quantity and activity**

Immunoblotting was used to compare SodA enzyme quantity. Bacterial lysates were adjusted to a standard A280 value, applied to a SDS-12% PAGE gel for separation of proteins by electrophoresis and transferred to nitrocellulose membranes and hybridized with a 1:1000 dilution of rabbit polyclonal antisera against recombinant SodA [84]. To measure superoxide dismutase activity in lysates and supernatants of vaccine cultures, we monitored the % inhibition of a superoxide anion (O$_2^{-}$) - induced reduction of a water-soluble tetrazolium (WST-1) salt to a formazan dye by colorimetric methods (SOD Assay Kit-WST, Dojindo Molecular Technologies, Inc., Gaithersburg, MD). One unit was defined as the amount of SodA that inhibited color development at 440 nm by 50%.

**Mice and Infections**

Experiments in mice were approved by the Vanderbilt Institutional Care and Use Committee and conform to regulatory standards for research involving vertebrate animals. Female C57Bl/6 mice and Balb/c mice aged 5-6 weeks were purchased from Jackson Laboratories (Bar Harbor, ME). To assess persistence of the vaccine strain in vivo, mice were inoculated retro-orbitally with approximately 5×10$^7$ CFU. To assess immune responses, mice were vaccinated subcutaneously or retro-orbitally with sterile broth media or with preparations of BCG or 3dBCG (BCGΔsecA2ΔsigHΔhigR) in broth media at 5×10$^7$ CFU. In order to eliminate the vaccine strain before a mycobacterial challenge, mice were treated with isoniazid and rifampin by adding 100 mg/liter of each antibiotic to drinking water [85]. After 30 days on this regimen, mice were then rested for at least two weeks before intravenous challenge (retro-orbital or tail-vein) with 2×10$^7$ CFU of BCG Tice.

**Gene expression analysis**

C57Bl/6 mice, six per group, were injected retro-orbitally with 1.5×10$^7$ CFU of BCG, 3dBCG, or phosphate-buffered saline (control). Mice were euthanized 72 hours later and the spleens were harvested and immersed in liquid nitrogen. To recover total cellular RNA, each spleen was homogenized with a mortar and pestle in the presence of liquid nitrogen. The homogenized tissue was extracted with Trizol (Gibco/Invitrogen, Carlsbad, CA) followed by a chloroform extraction and isopropanol precipitation. The pellet was resuspended in DNase/RNase free water (Gibco/Invitrogen, Carlsbad, CA) and the sample was DNase-treated (Promega, Madison, WI) for 30 minutes at 37°C. Total cellular RNA was further purified with a Qiagen RNeasy column purification system (Qiagen, Valencia, CA), quantitated with a spectrophotometer, and normalized. cDNA was prepared from each RNA sample with the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA) and analyzed with Taqman primer-probe pairs (Applied Biosystems) on an Applied Biosystems 7900HT Real-Time PCR system.

**Measurement of Immune Responses**

To obtain lymphocytes for measurement of immune responses after vaccination or challenge, mice were euthanized and splenocytes were isolated and suspended in RPMI, enumerated, and three million cells were used for each stimulation condition. Synthetic peptides TB10.474-488 - ST15, a CD4 epitope [29] and TB10.3/10.420-28 - GL9, a CD8 epitope [30] (Genemed Synthesis Inc, San Antonio, TX) and BCG-infected bone-marrow derived macrophages (BMDMs) were used as stimulants. BMDMs were derived from femur bone marrow stem cells which were cultured for 5 days in petri-dishes in DMEM medium containing 10% fetal calf serum and 20% L-929 conditioned medium. Three days before harvesting splenocytes, the BMDMs were transferred to 24-well plates at 1 million cells per well, and infected the next day with 5:1 MOI BCG Tice in the presence of 20 ng/ml IFN-γ for 4 hours. Paired wells of BMDMs were similarly treated with IFN-γ but left uninfected. Forty-eight hours after infection of BMDMs, 3 million harvested splenocytes were re-stimulated on uninfected and BCG-infected BMDMs for 20 hours, the last 18 hours in the presence of brefeldin A. For measurement of cytokine responses to peptide, harvested splenocytes were stimulated with or without peptides for 6 hours, the last 4 hours in the presence of brefeldin A. Cytokine production was assessed by intracellular cytokine staining and flow cytometry.

**Intracellular cytokine staining and flow cytometry**

Stimulated splenocytes were stained sequentially with live/dead fixable cell stain kit - violet fluorescent reactive dye (Invitrogen Molecular Probes, Carlsbad, CA) and a cocktail of surface antibodies - anti -CD3 APC-Cy7 , -CD4 PE-Texas Red, and -CD8 Pacific Orange (Invitrogen), fixed and permeabilized with cytofix/cytperm solution and stained with anti-IFN-γ PE-Cy7, -TNF-α FITC, and -IL-2 PE (all from BD biosciences, San Jose, CA) and a cocktail of surface molecules - anti-CD3 APC-Cy7, -CD4 PE-Cy7, and -CD8 PE-Cy7, normalized to a cocktail of surface markers. The total number of cytokine+ CD4+ and CD8+ T cells per whole spleen was calculated from the products of the total number of splenocytes multiplied by the proportion of cytokine+ CD4+ and CD8+ T cells under each stimulation condition. Antigen-specific cytokine+ T cells were calculated as the difference between cytokine+ cells from the unstimulated and stimulated conditions.

**Statistics**

The planned comparison in each experiment was between the immune response to BCG and the immune response to 3dBCG. Unless otherwise indicated, these comparisons were performed using the Mann-Whitney determination for a non-parametric
**Supporting Information**

**Table S1** Tools for genetic manipulations and bacterial strains

| References |
|------------|
| 1. Forman HJ, Torres M (2001) Signaling by the respiratory burst in macrophages. JUIMBB Life 51: 365–371. |
| 2. Kantergwa S, Jornot L, Deneogues C, Nicod LP (2003) Superoxide anions induce the maturation of human dendritic cells. Am J Respir Crit Care Med 167: 431–437. |
| 3. van der Veen RC, Dotin TA, Karapetian A, Holland SM, Hofman FM (2004) Extra-cellular superoxide promotes T cell expansion through inactivation of nitric oxide. J Neuroimmunol 133: 103–109. |
| 4. Edwards KJ, Cynamon MH, Voladri RK, Hager CC, DeStefano MS, et al. (2001) Iron-cofacted superoxide dismutase inhibits host responses to Mycobacterium tuberculosis. Am J Respir Crit Care Med 164: 2211–2219. |
| 5. Braunstein M, Espinosa BJ, Chan J, Belisle JT, Jacobs WR Jr (2003) SecA2 functions in the secretion of superoxide dismutase A and in the virulence of Mycobacterium tuberculosis. Mol Microbiol 48: 453–464. |
| 6. Hinchee J, Lee S, Leon BY, Basaraba RJ, Venkataswamy MM, et al. (2007) Enhanced priming of adaptive immunity by a protopootic mutant of Mycobacterium tuberculosis. J Clin Invest 117: 2979–2989. |
| 7. Raman S, Song T, Puyang X, Bardarov S, Jacobs WR Jr, Husson RN (2001) The alternative sigma factor SigH regulates major components of oxidative and heat stress responses in Mycobacterium tuberculosis. J Bacteriol 183: 6119–6125. |
| 8. Manganelli R, Voskuil MI, Schoonh GL, Duhnae E, Gomez M, Smith J (2002) Role of the extracytoplasmic function factor σE in Mycobacterium tuberculosis global gene expression. Mol Microbiol 45: 365–374. |
| 9. Kaushik D, Schneider BG, Tyagi S, Yoshimatsu T, Scott C, et al. (2002) Reduced immunopathology and mortality despite tissue persistence in a Mycobacterium tuberculosis mutant lacking alternative σ factor, SigH. Proc Natl Acad Sci U S A 99: 8330–8335. |
| 10. Ng VH, Coe JS, Suzana AC, MacMicking JD, Kinney JD (2004) Role of KatG catalase-peroxidase in mycobacterial pathogenesis: counteracting the phagocyte oxidative burst. Mol Microbiol 52: 1291–1302. |
| 11. Wilson T, de Lisle GW, Mencinikevica JA, Blanchard JS, Collins DM (1998) Antimicrobial RNA to ahpC, an oxidative stress defense gene involved in nonaizal resistance, indicates that AhpC of Mycobacterium buccae has virulence properties. Microbiology 144: 2687–2695. |
| 12. Akhtar P, Srivastava S, Srivastava A, Srivastava M, Srivastava BS, Srivastava R (2000) Rv3303c of Mycobacterium tuberculosis protects tubercle bacilli against oxidative stress in vivo and contributes to virulence in mice. Microbes Infect 2: 2853–2862. |
| 13. Ten Dam HG (1993) BCG vaccination. In: Reichman LB, Hershfield ES, eds. Tuberculosis: a comprehensive international approach. New York: Marcel Dekker. pp 251–274. |
| 14. Fine PEM, Carneiro IAM, Milsten JB, Clements JD (1999) Issues relating to the use of BCG in immunization programs. A discussion document. Document WHO/VKB/99.23. |
| 15. Anderssen P, Doherty TM (2005) The success and failure of BCG - implications for a novel tuberculosis vaccine. Nat Rev Microbiol 3: 656–662. |
| 16. (2008) Global tuberculosis control: surveillance, planning, financing. World Health Organization. http://www.who.int/tb/publications/global_report_2008/en/index.html. |
| 17. Borch R, Gordon SV, Garnier T, Eigenmier K, Frigui W, et al. (2007) Genome plasticity of BCG and impact on vaccine efficacy. Proc Natl Acad Sci U S A 104: 5096–5101. |
| 18. Behr MA, Wilson MA, Gill WP, Salamon H, Schoonh GK, et al. (1999) Comparative genomics of BCG vaccines by whole-genome DNA microarray. Science 284: 1520–1523. |
| 19. Brosch R, Gordon SV, Buchriere C, Pyn AS, Garnier T, Cole ST (2000) Comparative genomics uncovers large tandem chromosomal duplications in Mycobacterium bovis BCG Pasteur. Year 17: 111–123. |
| 20. Alam MS, Garg SK, Agrawal P (2008) Studies on structural and functional divergence among seven WhiB proteins of Mycobacterium tuberculosis H37Rv. FEBS J. |
| 21. Jaeger T, Bulle H, Eohe L, Menge U, Singh M, et al. (2004) Multiple thiorhodocin-mediated routes to detoxify hydroperoxides in Mycobacterium tuberculosis. Arch Biochem Biophys 425: 182–191. |
| 22. Via LE, Deretic D, Ullner R, Hilder NS, Huber LA, Deretic V (1997) Arrest of mycobacterial phagosome maturation is caused by a block in vesicle fusion between stages controlled by rab5 and rab7. J Biol Chem 272: 13526–13531. |
| 23. Kremar L, Estague J, Brandt E, Ammend J, Loich C (1997) Mycobacterium bovis Bacillus Calmette-Guérin infection prevents apoptosis of resting human monocytes. Eur J Immunol 27: 2450–2456. |
| 24. Wojciechowski W, DeSanctis J, Skamene E, Bardzie D (1999) Attenuation of MHC class II expression in macrophages infected with Mycobacterium bovis bacillus Calmette-Guérin involves class II transactivator and depends on the Nurr1 gene. J Immunol 163: 2688–2696. |
| 25. Gagliardi MC, Teltori R, Giromoni F, Battioni M, Sargentini V, et al. (2005) Mycobacterium bovis Bacillus Calmette-Guérin infects DC-SIGN- dendritic cells and causes the inhibition of IL-12 and the enhancement of IL-10 production. J Leukoc Biol 78: 106–113. |
| 26. Kamath AB, Behar SM (2005) Anamnestic responses of mice following Mycobacterium bovis BCG vaccination. Infect Immun 73: 6130–6138. |
| 27. Dubos RJ, Pierce CH, Schaefer WB (1956) Differential characteristics in vitro and in vivo of several substrains of BCG. III. Multiplication and survival in vivo. Am Rev Tuberc. 74: 683–698. |
| 28. Lagranderie MR, Balazuc AM, Derich A, Cedric CD, Ghoergia M (1996) Comparison of immune responses of mice immunized with five different Mycobacterium bovis BCG vaccine strains. Infect Immun 64: 1–9. |
| 29. Tree JA, Williams A, Clark S, Hall G, Marsh PD, Iwany J (2004) Intranasal bacille Calmette-Guérin (BCG) vaccine dosage needs balancing between protection and lung pathology. Clin Exp Immunol 138: 405–409. |
| 30. Bunch-Christensen K, Ladefoged A, Guld J (1978) The virulence of some strains of BCG for golden hamsters. Further studies. Bull World Health Organ 45: 63–70. |
| 31. Mak TK, Hesseling AC, Hussey GD, Cotton MF (2008) Making BCG vaccination programmes safer in the HIV era. Lancet 372: 786–787. |
| 32. Behr MA, Small PM (1997) Has BCG attenuated to impotence? Nature 389: 133–134. |
| 33. Mostowy S, Tsolaki AG, Small PM, Behr MA (2003) The in vitro evolution of BCG vaccines. Vaccine 21: 4270–4274. |
| 34. Cooper JB, McIntyre K, Badasso MO, Wood SP, Zhang Y, et al. (1995) X-ray structure analysis of the iron-dependent superoxide dismutase from Mycobacterium tuberculosis at 2.0 Angstroms resolution reveals novel dimer-dimer interactions. J Mol Biol 246: 531–544. |
| 35. Olsen AW, Brandt L, Agger EM, van Pintxeteren LA, Anderssen P (2004) The influence of remaining live BCG organisms in vaccinated mice on the maintenance of immunity to tuberculosis. Scand J Immunol 60: 273–277. |
| 36. Seder RA, Darragh PA, Roederer M (2000) T-cell quality in memory and protection: implications for vaccine design. Nat Rev Immunol 3: 656–662. |
| 37. Hervas-Stubbs S, Majlessi L, Simsova M, Morova J, Rojas MJ, et al. (2006) High frequency of CD4+ T cells specific for the TB10.4 protein correlates with protection against Mycobacterium tuberculosis infection. Infect Immun 74: 3396–3407. |
38. Kamath A, Woolworth JS, Behar SM (2006) Antigen-specific CD8+ T cells and the development of central memory during Mycobacterium tuberculosis infection. J Immunol 177: 6361–6369.

39. Blattman JN, Grayson JM, Wherry EJ, Kaech SM, Smith KA, Ahmed R (2003) The therapeutic use of IL-2 to enhance antiviral T-cell responses in vivo. Nat Med 9: 540–547.

40. Williams MA, Tynzik AJ, Bevan MJ (2006) Interleukin-2 signals during priming are required for secondary expansion of CD8+ memory T cells. Nature 441: 890–893.

41. Corhay A (2006) A three-cell model for activation of naive T helper cells. Scand J Immunol 64: 93–96.

42. Luther SA, Cyster JG (2001) Chemokines as regulators of T cell differentiation. J Immunol 167: 551–559.

43. Zhou L, Ivanov II, Spolski R, Min R, Shenderov K, et al. (2007) IL-6 programs airway epithelial cells to increases CD4+ memory T cell accumulation. J Immunol 178: 6361–6369.

44. Vogelzang A, McGuire HM, Yu D, Sprent J, Mackay CR, King C (2008) A fundamental role for interleukin-2 in the generation of T follicular helper cells. Immunity 29: 127–137.

45. Faúndez L, Chan CK, Grinstein S, Downey GP (1993) Regulation of tyrosine phosphorylation in neutrophils by the NADPH oxidase. Role of reactive oxygen intermediates. J Biol Chem 268: 17131–17137.

46. Heijnen CJ, van der Meer JW, Zegers BJ (1986) Altered antigen-presentation in T lymphocytes of rheumatoid arthritis patients. Arthritis Rheum 29: 111–117.

47. Vogt G, de Beaucoudrey L, et al. (2008) Decreased CD4+CD25+ (memory) T cells in patients with chronic granulomatous disease. J Infect Dis 196: 983–985.

48. Romani L, Fallarino F, De Luca A, Montagnoli C, D’Angelo C, et al. (2008) Defective tryptophan catabolism underlies inflammation in mouse chronic granulomatous disease. Nature 451: 211–215.

49. Filipe-Santos O, Bustamante J, Chapgier A, et al. (2008) Decreased CD4+CD25+ (memory) T cells in patients with chronic granulomatous disease. J Infect Dis 196: 983–985.

50. Chu SF, Tam CM, Wong HS, Kam KM, Lau YL, Chiang AK (2007) Decreased CD4+CD25+ (memory) T cells in patients with chronic granulomatous disease. J Infect Dis 196: 983–985.

51. Trinchieri G (2003) Interleukin-12 and the regulation of innate resistance and adaptive immunity. Nat Rev Immunol 3: 133–146.

52. Pittman RE, Taylor HC (1957) The problem of standardization of BCG. J Infect Dis 96: 1155–1159.

53. Wilkinson RJ, Zhu X, Wilkinson KA, Lalvani A, Ivanyi J, et al. (1998) 38 000 tuberculosis prevention trial (1979) Trial of BCG vaccines in south India for tuberculosis prevention: first report. Bull World Health Organ 76: 314–320.

54. Aronson JD, Aronson CF, Taylor HC (1957) The problem of standardization of BCG. J Infect Dis 96: 1155–1159.

55. Irvine KN (1949) B.C.G. Vaccination in Theory and Practice. London and Oxford: Blackwell Scientific Publications, Ltd. 120 p.

56. Suter WE, Dubos RJ (1951) Variability of BCG strains (Bacillus Calmette-Guerin). J Exp Med 93: 559–572.

57. Dubos RJ, Pierce CH (1956) Differential characteristics in vitro and in vivo of Mycobacterium bovis BCG strains. Am Rev Tuberc 75: 692–693.

58. Aronson JD, Widmar H, Heron RD, Jackson EK, Guerin. J Exp Med 93: 559–572.

59. Aronson JD, Widmar H, Heron RD, Jackson EK, Guerin. J Exp Med 93: 559–572.

60. Tuberculosis Prevention Trial (1979) Trial of BCG vaccines in south India for tuberculosis prevention: first report. Bull World Health Organ 57: 819–827.

61. Ladefoged A, Bunch-Christensen K, Guld J (1976) Tuberculin sensitivity in guinea-pigs after vaccination with varying doses of BCG of 12 different strains. J Infect Dis 133: 623–630.

62. Fricch-Moller J, Acharyulu GS, Parthasarathy R (1968) Observations on the protective effect of BCG vaccination in a South Indian rural population: Third Report. Indian Journal of Tuberculosis 13: 40–46.

63. Oettinger T, Jorgensen M, Ladefoged A, Haslov K, Andersen P (1999) Development of the Mycobacterium bovis BCG vaccine: review of the historical and biochemical evidence for a genealogical tree. Tuberc Lung Dis 79: 243–250.

64. Hart PD, Sutherland I (1977) BCG and vole bacillus vaccines in the prevention of tuberculosis in adolescence and early adult life. Br Med J 2: 293–295.

65. Comstock GW (1960) Identification of an effective vaccine against tuberculosis. Am Rev Respir Dis 138: 479–480.

66. Comstock GW (1994) Evaluating vaccination effectiveness and vaccine efficacy by means of case-control studies. Epidemiol Rev 16: 77–89.

67. Comstock GW (2000) Simple, practical ways to assess the protective efficacy of a new tuberculosis vaccine. Clin Infect Dis 30 Suppl 3: S250–S253.

68. Shor NA, Chaparas SD, Pearson J, Charies M (1973) Virulence of six strains of Mycobacterium bovis (BCG) in mice. Infect Immun 8: 736–742.

69. Ladefoged A, Bunch-Christensen K, Guld J (1970) The protective effect in bank voles of some strains of BCG. Bull World Health Organ 43: 71–90.

70. Aronson JD, Schneider P (1950) The problem of standardization of BCG vaccine. Am J Public Health 40: 333–344.

71. Saunders BM, Frank AA, Orme IM, Cooper AM (2002) CD4 is required for the development of a protective granulomatous response to pulmonary tuberculosis. Cell Immunol 216: 65–72.

72. Mahomed H, Kibel M, Hawkridge T, Schaal HS, Hanekom WA, et al. (2006) The impact of a change in bacille Calmette-Guerin vaccine policy on tuberculosis incidence in children in Cape Town, South Africa. Pedaet Infect Dis J 25: 1167–1172.

73. Davies V, Hanekom WA, Mianoor N, Gamieldien H, Gelderbloem SJ, et al. (2006) The effect of bacille Calmette-Guerin vaccine strain and route of administration on induced immune responses in vaccinated infants. J Infect Dis 193: 531–536.

74. Whitmire JK, Eam B, Whitton JL (2008) Tentative T cells: memory cells are quick to respond, but slow to divide. PLoS Pathog 4: e1000041.

75. Skelley YA, Sadowski JC (2006) Advances in tuberculosis vaccine strategies. Nat Rev Microbiol 4: 469–476.

76. Herr HW, Morales A (2008) History of bacillus Calmette-Guerin and bladder cancer: an immunotherapy success story. J Urol 179: 53–56.

77. Brandau S, Suttman H (2007) Thirty years of BCG immunotherapy for non-muscle invasive bladder cancer: a success story with room for improvement. Biomed Pharmacother 61: 299–305.

78. Dubos RJ, Pierse CH (1957) Tice strain of BCG. Am Rev Tuberc 75: 692–693.

79. Horwitz MA, Harth G, Dillon BJ, Maslesa-Galic S, et al. (2009) Commonly administered BCG strains including an evolutionarily early strain and evolutionarily late strains of disparate genealogy induce comparable protective immunity against tuberculosis. Vaccine 27: 441–445.

80. Braunstein M, Bardarov SS, Jacobs WR Jr (2002) Genetic methods for deciphering virulence determinants of Mycobacterium tuberculosis. Methods Enzymol 358: 67–99.

81. Ho SN, Hunt HD, Horton RM, Pollen JK, Pease LR (1989) Site-directed mutagenesis by overlap extension using the polymerase chain reaction. Gene 77: 51–59.

82. Carlloz A, Touati D (1968) Isolation of superoxide dismutase mutants in Escherichia coli: is superoxide dismutase necessary for aerobic life? EMBO J 5: 623–630.

83. Graham JE, Clark-Curtiss JE (1999) Identification of Mycobacterium tuberculosis RNAs synthesized in response to phase transition by human macrophages by selective capture of transcribed sequences (SCOTS). Proc Natl Acad Sci U S A 96: 11554–11559.

84. Lakey DL, Vlodari RK, Edwards KM, Hager C, Samten B, et al. (2000) Enhanced production of recombiant Mycobacterium tuberculosis antigens in Escherichia coli by replacement of low-usage codons. Infect Immun 68: 233–236.

85. Collins FM, Watson SR (1980) Effect of chemotherapy on suppressor T cells in BCG-infected mice. Immunology 40: 529–537.