Concurrent infection with *Mycobacterium tuberculosis* confers robust protection against secondary infection in macaques


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**Abstract**

For many pathogens, including most targets of effective vaccines, infection elicits an immune response that confers significant protection against reinfection. There has been significant debate as to whether natural *Mycobacterium tuberculosis* (Mtb) infection confers protection against reinfection. Here we experimentally assessed the protection conferred by concurrent Mtb infection in macaques, a robust experimental model of human tuberculosis (TB), using a combination of serial imaging and Mtb challenge strains differentiated by DNA identifiers. Strikingly, ongoing Mtb infection provided complete protection against establishment of secondary infection in over half of the macaques and allowed near sterilizing bacterial control for those in which a secondary infection was established. By contrast, boosted BCG vaccination reduced granuloma inflammation but had no impact on early granuloma bacterial burden. These findings are evidence of highly effective concomitant mycobacterial immunity in the lung, which may inform TB vaccine design and development.

**Author summary**

Tuberculosis (TB), a lung disease caused by the bacterial pathogen *Mycobacterium tuberculosis*, is endemic in many developing countries. This infection is transmitted from a person with active tuberculosis through coughing, talking, and singing. Exposure to this bacterium can result in a spectrum of infection outcomes, including in the majority of persons asymptomatic infection, known as latent TB. However, re-exposure to those with active disease occurs frequently, particularly in crowded conditions. Here we demonstrate...
Introduction

There has been significant debate as to whether natural Mtb infection confers protection against reinfection. Epidemiologic studies from the pre-antibiotic era suggest that primary *Mycobacterium tuberculosis* (Mtb) infection provides up to 80% protection against TB disease due to secondary exposure[1], although assessing protection against actual reinfection is not possible. However, up to ~20% of patients who complete drug treatment develop TB again, in part due to reinfection[2–6]. Recent studies also suggest that mixed infections (two or more Mtb isolates) are detectable in 2–18% of individuals with active TB[4, 7, 8], although some of these studies are in HIV+ individuals. In mice, ongoing or treated Mtb infections only reduce bacterial burdens in the lung by ~10-fold, roughly equivalent to BCG vaccination[9, 10]. These data are often used to inform the field’s understanding of the extent of protection that a primed immune response can provide against Mtb infection, which is a critical question for vaccine development. However, these conclusions are confounded by uncertainties regarding host immune status and prior exposure in the human studies, outcome measures of disease rather than establishment of a secondary infection, and the relevance of the small animal models to human infection.

Here, we sought to quantitatively assess the effect of a concurrent Mtb infection on rechallenge in the cynomolgus macaque model[11–13], which recapitulates nearly all aspects of human Mtb infection. To probe the dynamics of reinfection, we used serial [18F]-fluorodeoxyglucose (FDG)-PET-CT imaging to track timing of granuloma formation following secondary exposure[14–16], which allowed all granulomas to be retrieved at necropsy. The outcomes of primary and secondary challenges were defined with Mtb libraries marked with unique DNA identifiers that were tracked by sequencing and/or a custom direct hybridization (NanoString) assay[14, 17, 18]. Our findings indicate that primary ongoing infection provides robust protection against infection upon secondary exposure to Mtb.

Results

Primary infection restricts new granuloma formation after re-challenge

Eight cynomolgus macaques received a primary challenge with Mtb Erdman library A (<15 CFU) (Fig 1A). As expected based on our published work[12, 13, 19], this infection resulted in a range of outcomes by 16 weeks as assessed by PET-CT[13], from minimal to progressive disease (Fig 1B and S1 Table). Sixteen weeks after primary infection, animals were rechallenged with Mtb Erdman library B (<15 CFU). Six naïve control animals were challenged with library B in parallel (Fig 1A). Granuloma formation after library B challenge was tracked for ~4 weeks by PET-CT imaging (S1 Table) (Fig 1C). The number of granulomas detectable by PET-CT imaging at 4 weeks post-infection in naïve animals is one correlate of the number of bacteria that successfully establish infection[14, 15]. Individual granulomas and lymph nodes were obtained at necropsy (4 weeks post-library B) for bacterial and immunologic analyses. At this time point, which is just prior to the onset of adaptive immunity in naïve animals, granuloma bacterial burdens are relatively uniform and at their highest levels with minimal bacterial killing in naïve animals[14, 20][21]. Thus, comparing granulomas at 4 weeks post-library B in naïve and reinfected animals allows a direct comparison of early bacterial control.
Using PET-CT to identify formation of new granulomas, there was no aggregate difference between new granuloma formation after library B challenge in the presence or absence of ongoing infection (S1A Fig). However, this metric alone cannot distinguish new granuloma formation due to library B challenge from granulomas formed by ongoing dissemination from sites of primary infection (library A). Indeed, three animals had apparent dissemination just prior to or during secondary infection, based on imaging (monkey IDs: 19415, 19515, 19615). Moreover, imaging alone fails to capture the potential for library B seeding sites of existing infection as described in fish, frogs, and mice[22, 23]. Therefore, we deconvoluted the identity of library tags in granulomas formed after primary and secondary infections. Tags were identifiable both from bacteria cultured from granulomas and, in most cases, from tissue homogenates including the homogenates of sterile lesions.

As expected, DNA tags from sampled granulomas in naïve animals mapped entirely to library B (Fig 1D and S2 Table). In animals challenged during ongoing primary infection, there was no significant difference in total number of granulomas containing library B as compared to naïve animals (p = 0.1525) (S1B Fig). Nine granulomas (of 95 total analyzed) from reinfected monkeys contained both library A and library B DNA (S2 Table), suggesting that
occasionally library B seeded pre-existing sites of infection. However, concurrent infection did reduce the formation of new granulomas attributable only to library B (Fig 1D, p = 0.0083). The number of granulomas attributable only to library B at 4 weeks was also significantly lower than the number of granulomas established by library A granulomas at 4 weeks (as determined by PET CT) in the same animals (p = 0.0156, S1C Fig). Indeed, there was complete protection in one animal with no detectable library B DNA in any tissues (monkey ID: 19815) (Figs 1D and S1A and S1B).

**Primary infection significantly reduced growth and increased killing of the Mtb rechallenge strain**

To assess growth of the secondary challenge strain, we evaluated the bacterial burdens of individual granulomas in both cohorts and used the library tags to attribute these bacterial loads to library A and/or library B (Fig 2A). In naïve animals, the distribution of granuloma bacterial burdens (library B) was consistent with our previous studies[14] with a median bacterial load of 8300 and few sterile granulomas. By contrast, most granulomas from reinfected animals had lower bacterial loads than in naïve animals, with many sterile granulomas (Fig 2A, p<0.0001). Strikingly, in concurrently infected animals, the few library B granulomas that did form had markedly lower bacterial burdens than in naïve macaques where these granulomas were assessed at the same time post-library B infection (Figs 2A and 3C). In addition to the one animal that had no library B DNA (monkey ID: 19815), four animals had no viable library B bacteria; in these animals library B DNA tags were only found in homogenates of sterile granulomas (S3 Table). Only 1 of the 9 granulomas with both library A and B DNA tags grew library B bacteria. Thus, primary infection fully protected 5 out of 8 animals against productive (CFU+) infection with the challenge strain (Fig 2A). Collectively, these data suggest that primary Mtb infection initiates an immune response which leads to rapid neutralization of the challenge strain.

Note that the bacterial loads in granulomas containing library A, delivered 20 weeks prior to necropsy, were also lower than library B granulomas from naïve animals analyzed ~4 weeks post infection. However, this decrease in bacterial load over time is consistent with our previously published studies[14] and data from historical macaques infected with Mtb Erdman (S2 Fig), where CFU/granuloma decreases after the onset of adaptive immunity, and did not support the hypothesis that rechallenge altered the course of the primary infection. To further investigate this question, we assessed inflammation in individual library A granulomas by PET CT prior to and after library B challenge and compared this to similar time points in historical macaques challenged with Mtb Erdman, and again saw no evidence that secondary infection altered the inflammatory dynamics of the preexisting infection (S3 Fig).

We next sought to distinguish whether the lower bacterial burden in granulomas formed after reinfection reflected restriction of bacterial growth and/or true enhancement of bacterial killing. We assessed granuloma bacterial genome counts (CEQ) attributable to library B in naïve and rechallenged animals and then quantified killing of the challenge strain by relating viable library B live bacteria counts (CFU) to library B genome counts (CEQ), as described[14, 24] (Fig 2B). In rechallenged animals, library B granulomas had significantly lower CEQ than in naïve animals, indicating reduced replication of the infecting bacteria. Library B granulomas had concomitantly lower CFU (Fig 2B), which also reflected significantly increased killing (as reflected by CFU/CEQ) of the library B in granulomas from rechallenged animals relative to naïve animals (Fig 2B and 2C median log killing = -1.41 (naïve) vs. -3.19 (reinfected), p = 0.0002; ~1.75 log increase in killing). Interestingly, there was no correlation between bacterial killing of library B Mtb and host lung inflammation at the time of second challenge (Fig 2D, r = 0.03, p = 0.8832), nor was there any association between total thoracic bacterial burden
at necropsy in those macaques who had few library B+ granulomas and those who had none (S5 Fig). Thus, the extent of disease at time of secondary exposure does not appear to affect the protection against reinfection.

**Primary infection provides substantially more protection against Mtb challenge than a fusion protein-boosted BCG vaccine**

To assess the protection provided by concurrent infection in the context of current vaccine strategies, we compared the bacterial loads from granulomas in our naïve and reinfection
monkeys to those from macaques vaccinated with BCG boosted with an adjuvanted fusion protein (H56 in CAF01[25, 26]). We previously showed that BCG+H56 provides protection against reactivation, reduces pathology and improves survival of Mtb-infected macaques[27, 28]. Moreover, the size and FDG avidity of early granulomas in BCG+H56 vaccinated animals were reduced compared to unvaccinated animals, as was found in the granulomas formed in the setting of reinfection (Fig 3A and 3B). However, at the early time points assessed here, the

Fig 3. Primary Mtb infection reduces bacterial burden in secondary challenge granulomas, while boosted BCG has no effect on early granuloma bacterial burden. (a, b) Size (mm, by CT) and inflammation (as measured by $^{18}$F-FDG avidity via standard uptake value, SUV) for library B (for naïve and reinfection macaques) or Mtb (for BCG+H56 and unvaccinated controls) granulomas. (c) CFU of Mtb library B granulomas from naïve (blue) and reinfection (red) macaques and 5–6 week granuloma CFU in boosted BCG (green) and unvaccinated control (purple) macaques challenged with Mtb Erdman. For a, b, and c, each symbol is a granuloma, p < 0.0001, Kruskal-Wallis tests, Dunn’s multiple comparison adjustment was used and adjusted p-values are reported in the figure above corresponding comparisons. All granulomas from N = 6 macaques (naïve); N = 7 macaques (reinfection); N = 6 BCG+H56 macaques; N = 6 unvaccinated macaques are represented. (d) Left panel: Number of thoracic lymph nodes from each macaque which were CFU+ for Mtb library B. Right panel: Number of thoracic lymph nodes from unvaccinated or BCG+H56 vaccinated CFU+ positive for Mtb at 5–6 weeks post-infection. N = 6 (naïve), N = 8 (reinfection), and N = 6 (unvaccinated). Each symbol is a macaque, Mann-Whitney test p-values reported in figure.

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bacterial control engendered by primary Mtb infection was dramatically superior to that found in the vaccinated animals (Fig 3C). In addition, primary infection prevented dissemination of the reinfection strain to thoracic lymph nodes while BCG+H56 did not have a similar effect (Fig 3D); the number of granulomas at 4–5 weeks post-challenge in BCG-vaccinated macaques was similar to contemporaneous control macaques (S1D Fig). By contrast, in aggregate primary infection provided ~10,000 fold protection against Mtb reinfection as assessed by granuloma bacterial burdens (Fig 3C), reflecting the combined effects of restricted establishment of infection, limited bacterial growth, and increased bacterial killing.

**Immunologic landscape in lungs after reinfection**

To provide insight into the local immunologic landscape that may contribute to the protection seen with reinfection, we first assessed uninvolved lung tissue from reinfeected monkeys using Luminex (S4A Fig) compared to responses in an uninfected monkey, where it was only possible to euthanize one animal for analysis of normal lung tissue. These data suggested that even in uninvolved tissue, infection is associated with the expression of innate cytokines and chemokines. Though this analysis was limited by the availability of uninfected animals, the data suggest increased IL-8 in lung tissue from infected monkeys compared to uninfected control monkeys, possibly implicating increased neutrophil migration in infected tissues, and a decrease in IL-1 in infected tissues, with a trend towards increased IL-1RA. CXCL9, a chemokine that binds CXCR3 was also slightly higher in infected lung tissues, suggesting possible increased T cell migration, while CXCL13, which is chemotactic for B cells, was reduced. We also assessed uninvolved lung tissue from a separate set of Mtb-infected monkeys (not reinfeected) at similar time points for Mtb-specific (ESAT6/CFP10) T cell responses (S4B Fig). Mtb-specific T cell responses were higher in infected lung tissue, compared to uninfected lung tissue, indicating the presence of T cells resident within lung tissue that could rapidly activate macrophages upon encountering a new Mtb bacillus. Thus, a reinfecting bacillus encounters innate and adaptive immune responses in lung tissue, which may underlie the more rapid killing of the bacterium that we observed.

We next assessed T cell responses in the few library B granulomas that did establish in the reinfeected monkeys and compared these to library A granuloma in the same animals and library B granulomas that established in naïve animals (Fig 4). In general, T cell responses in library A and B granulomas in reinfeected macaques were similar, and the data reflect the normal variability seen in immune responses in granulomas[20]. There were no significant differences in the frequency of T cells making T\(_H\)1 cytokines among library B granulomas from naïve or reinfeected macaques, or library A granulomas in reinfeected macaques. However, IL-10 was significantly higher in both library A and B granulomas in reinfeected macaques compared to granulomas in naïve macaques. Our prior data suggested that the combination of T cells expressing IL-10 and T cells expressing a pro-inflammatory cytokine in granulomas was associated with the sterilization of Mtb granulomas[20]. The higher levels of IL-10 in granulomas with lower bacterial loads as compared to granulomas in naïve animals is consistent with this finding.

More broadly, production of inflammatory cytokine and chemokines (measured in granuloma supernatants) was lower in library B granulomas in animals with primary infection relative to those formed in naïve animals (Fig 5). TNF, IL-1β, IL-18, IL-1RA, IL-8, MCP-1, and MIG were all present at significantly (\(p<0.05–0.0001\)) lower levels in secondary granulomas, with a trend towards lower IFN-γ. This likely reflects the low bacterial burden in the reinfection granulomas and again may indicate a healing response as those granulomas rapidly kill the bacteria. Further study into the precise kinetics and cellular organization of these granulomas is needed to help inform intervention strategies.
Discussion

In this study, we present evidence for robust concomitant immunity in the cynomolgus macaque model of TB, using a combination of PET CT imaging and molecular analysis of DNA-tagged strains of M. tuberculosis in a macaque model. We found complete protection against productive secondary infection in five macaques with ongoing infection (Figs 1D and 2A), almost no productive dissemination to lymph nodes (Fig 3D), and a ~10,000-fold decrease in live Mtb in library B (reinfection) granulomas compared to age-matched granulomas in naïve animals (Figs 2B, 2C and 3C). One reinfected monkey had no trace of library B DNA despite receiving a secondary challenge dose of 10 CFU. The number of new granulomas
that were seen by PET CT and attributed to library B by Q-tag analysis at 4 weeks post-reinfection was significantly lower in macaques with ongoing primary (library A) infection, compared to the number of library B granulomas at 4 weeks post-infection in naïve (i.e. no primary infection) macaques. The new library B granulomas that were established in the reininfected animals were often sterile or had very low bacterial burden, in stark contrast to 4 week granulomas in naïve macaques, where the bacterial burden is generally quite high. Using CEQ analysis, we determined that the library B Mtb bacilli did not grow to the same extent in reininfected macaques as in naïve macaques, and that bacterial killing was increased. Thus, reduced new granuloma formation, reduced growth and increased killing resulted in a 10,000 fold decrease in live Mtb library B bacilli in reininfected macaques, compared to the bacterial burden of library B Mtb in naïve macaques at the same time point.

Fig 5. Reinfection challenge strain granulomas have reduced levels of pro-inflammatory cytokines and chemokines. Supernatants from library B granulomas from naïve or reinfection macaques were assessed for cytokine and chemokines. N = 27 (naïve) and N = 12 (reinfection). Each dot is a granuloma, p < 0.05–0.0001, Mann-Whitney test.

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The extent of protection in the setting of reinfection as compared to that provided by intradermal BCG followed by a fusion protein boost suggests that there is a fundamental difference between the immune profile elicited by primary Mtb infection and that generated by BCG intradermal vaccination. We suspect that this reflects the difference in immunity expressed by local (i.e. lung) resident T effector cells maintained by ongoing infection versus the systemic response promoted by a distal boosted BCG vaccination. Our data suggest that T cells specific for Mtb proteins are present in uninvolved lung tissue of infected macaques, which supports this hypothesis. In addition, we found modestly increased innate cytokines in uninvolved lung tissue from infected macaques, which may also contribute to reducing initial establishment of infection. Further studies are required to determine which factors are necessary or sufficient for protection against re-infection.

Immunologic assessment of the few library B granulomas that were established following reinfection showed increased IL-10+ T cell responses compared to those from naïve macaques. The IL-10+ T cell responses in reinfection granulomas were similar to those in the primary (library A) infection granulomas in the same macaques. We previously published that IL-10+ T cells in conjunction with T cells producing a proinflammatory cytokine (i.e. IL-17) in individual granulomas was associated with lower bacterial burdens, including sterility[20]. We interpret this to mean that either an immune balance of pro- and anti-inflammatory cytokines is a robust pathway to sterilization in granulomas, or that the IL-10 signal is a sign of granuloma healing once bacteria have been killed. At this point, we cannot distinguish between those possibilities. Nonetheless, the fact that the 20 week library A granulomas (most of which have succeeded in killing many of the bacteria within them[14, 17]) and the 4 week Library B granulomas in the same animals shared this increased IL-10+ T cell profile suggests that the library B granulomas are primed to quickly dispense with the infection, with IL-10 production being one factor associated with that clearance.

The data presented here have implications for TB vaccines. The extent of protection that we find against rechallenge suggests that robust protection via a vaccine is possible. This level of protection would be truly remarkable for a TB vaccine, and suggests that a protective vaccine for TB may have to induce long lived, likely local, immune responses that mimic those induced by primary infection. The recent study showing substantial protection in rhesus macaques by a CMV vector expressing multiple Mtb antigens suggests that sustained high level T cell responses are important for immunity but the correlate of protection analysis also pointed to the potential importance of an activated innate immune response in mediating bacterial clearance [29]. We could not formally define a correlate of protection conferred by concurrent infection since all concurrently infected animals were robustly protected against reinfection; defining a correlate requires failure of protection in some animals. There were no immunologic or disease parameters that segregated animals that fully sterilized the challenge from those that sterilized many but not all sites of infection (S5 Fig). However, our immunologic analysis does provide evidence of an activated innate response and Mtb specific T cells in the uninvolved lung tissue of infected animals that could participate in clearance.

In the current study, the primary infection and the reinfection strains were the same: virulent Mtb Erdman. Whether a primary infection with one strain could provide such robust protection against a heterologous strain remains to be tested. Given the broad immune responses induced by Mtb against multiple antigens, and the absence of data in human-like models indicating that immune responses against a single or a few antigens provide robust protection, we suspect that a heterologous re-challenge would be well-controlled. Nonetheless, such experiments are necessary to address this clinically relevant point.

Concomitant immunity has been described in other systems, e.g. tumor rejection models [30]. In terms of infectious diseases, this is best characterized for leishmaniasis, where the
presence of a live but contained *Leishmania* infection prevents establishment of a new infection. In the Leishmania model, this protection depends on the presence of high levels of T effector cells rather than central memory T cells, and is lost when the original infection is cleared[31]. Our study supports concomitant immunity in Mtb infections, although it remains to be determined whether complete clearance of primary Mtb infection with drug treatment will abrogate the extraordinary protective effects shown here[32]. While the precise immune mechanisms underlying the protection described here are not yet fully characterized, this study provides exciting clues for revised TB vaccine strategies to achieve both sterilizing immunity and protection from infection.

**Methods**

**Ethics statement**

University of Pittsburgh IACUC reviewed and approved the study protocol. The protocol assurance number for our IACUC is A3187-01. Specific Approval number: 15066174. The IACUC adheres to national guidelines established in the Animal Welfare Act (7 U.S.C. Sections 2131–2159) and the Guide for the Care and Use of Laboratory Animals (8th Edition) as mandated by the U.S. Public Health Service Policy.

All macaques used in this study were housed at the University of Pittsburgh in rooms with autonomously controlled temperature, humidity, and lighting. Animals were singly housed in caging at least 2 square meters apart that allowed visual and tactile contact with neighboring conspecifics. The macaques were fed twice daily with biscuits formulated for nonhuman primates, supplemented at least 4 days/week with large pieces of fresh fruits or vegetables. Animals had access to water ad libitum. Because our macaques were singly housed due to the infectious nature of these studies, an enhanced enrichment plan was designed and overseen by our nonhuman primate enrichment specialist. This plan has three components. First, species-specific behaviors are encouraged. All animals have access to toys and other manipulata, some of which will be filled with food treats (e.g. frozen fruit, peanut butter, etc.). These are rotated on a regular basis. Puzzle feeders foraging boards, and cardboard tubes containing small food items also are placed in the cage to stimulate foraging behaviors. Adjustable mirrors accessible to the animals stimulate interaction between animals. Second, routine interaction between humans and macaques are encouraged. These interactions occur daily and consist mainly of small food objects offered as enrichment and adhere to established safety protocols. Animal caretakers are encouraged to interact with the animals (by talking or with facial expressions) while performing tasks in the housing area. Routine procedures (e.g. feeding, cage cleaning, etc) are done on a strict schedule to allow the animals to acclimate to a routine daily schedule. Third, all macaques are provided with a variety of visual and auditory stimulation. Housing areas contain either radios or TV/video equipment that play cartoons or other formats designed for children for at least 3 hours each day. The videos and radios are rotated between animal rooms so that the same enrichment is not played repetitively for the same group of animals.

All animals are checked at least twice daily to assess appetite, attitude, activity level, hydration status, etc. Following *M. tuberculosis* infection, the animals are monitored closely for evidence of disease (e.g., anorexia, weight loss, tachypnea, dyspnea, coughing). Physical exams, including weights, are performed on a regular basis. Animals are sedated prior to all veterinary procedures (e.g. blood draws, etc.) using ketamine or other approved drugs. Regular PET/CT imaging is conducted on most of our macaques following infection and has proved very useful for monitoring disease progression. Our veterinary technicians monitor animals especially closely for any signs of pain or distress. If any are noted, appropriate supportive care (e.g. dietary supplementation, rehydration) and clinical treatments (analgesics) are given. Any animal

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considered to have advanced disease or intractable pain or distress from any cause is sedated with ketamine and then humanely euthanatized using sodium pentobarbital.

Macaque infections, PET-CT imaging, and tissue excision

Fourteen adult cynomolgus macaques (*Macaca fascicularis*) were obtained from Valley Biosystems (Sacramento, California) and screened for Mtb and other comorbidities during a month-long quarantine. Each macaque had a baseline blood count and chemical profile and was housed according to the standards listed in the Animal Welfare Act and the Guide for the Care and Use of Laboratory Animals. All procedures were approved by the Institutional Animal Care and Use Committee at the University of Pittsburgh. The animals were separated into two cohorts: 8 macaques were assigned to reinfection and 6 were assigned to naïve 4-week only controls. All animals were infected with barcoded strain Erdman Mtb via bronchoscopic instillation as previously published[11, 12]. The infection schema is provided in Fig 1A; 8 macaques were infected with Mtb library A, followed 16 weeks later by Mtb library B; 6 naïve macaques were infected with Mtb library B in a series of matched infections. All animals received an inoculum of <15 CFU (determined by plating a sample of the inoculum and counting CFU after 3 weeks) with the details listed in S1 Table. The animals were further subdivided such that 4 reinfection animals were directly paired with 2, naïve animals; 2 additional naïve animals were infected separately (S1 Table). Each macaque was followed with serial 2-deoxy-2-[18F]-fluoro-D-glucose ([18F]-FDG) PET-CT imaging as previously described[14–16] to identify and track lesion formation and progression over time. PET-CT scans were performed monthly during the primary infection, and as noted in S1 Table after reinfection. Total FDG activity in lungs was used to estimate thoracic bacterial burden prior to reinfection, as previously published[13, 33]. Granulomas were individually characterized by their date of establishment (scan date), size (mm), and relative metabolic activity as a proxy for inflammation ([18F]-FDG standard uptake normalized to muscle [SUVR][13, 33]). Granulomas ≥1mm can be discerned by our PET-CT imaging analysis (Fig 1B).

All macaques were necropsied at ~4 weeks post-library B infection. The pre-necropsy PET-CT scan was used to map new and old granulomas. To avoid barcode cross-contamination, individual granulomas were separately excised and processed, along with all thoracic lymph nodes. Each sample was homogenized to a single cell suspension, plated for CFU on 7H11 agar supplemented with oleic albumin dextrose catalase (OADC), and frozen aliquots were stored for DNA extraction.

Isolation and preparation of bacteria genomic DNA from tissue samples

A small portion of granuloma homogenate was frozen for qTag sequencing and chromosomal equivalent (CEQ) analysis. For analysis of bacteria that grew from the granulomas, bacterial colonies from granuloma and lymph node plates were scraped to obtain all colonies, and frozen. Genomic DNA was extracted as previously published[14, 24]. In brief, gDNA was twice extracted with phenol:chloroform:isoamyl alcohol (25:24:1, Invitrogen) with an intermediate bead beating step using 0.1mm zirconia-silica beads (BioSpec Products, Inc.).
Barcode/qtag determination via sequencing and nanostring

To identify DNA tags from each library, genomic DNA was diluted to 10 ng/μL and amplified with Q5 polymerase (New England Biolabs) with two rounds of PCR of 8–15 cycles each, using the primers described in Martin et al[17]. Samples were then sequenced on an Illumina MiSeq using v2 chemistry. Barcodes were identified using custom scripts as described in Martin et al. To identify qTags, genomic DNA was diluted to 100 ng/μL and amplified with 24–36 cycles of PCR, using the primers listed in S3 Table. PCR product was used as input in the NanoString nCounter assay (NanoString Technologies) with custom-designed probes to determine qTag identity. We attributed the bacteria in any mixed lesion to library B, a conservative assumption that would lead us to overestimate the success of library B in the setting of rechallenge.

Luminex on granuloma supernatant and lung tissue

Supernatants from granuloma and uninvolved lung tissue homogenates were frozen at -80˚C until time of assay. After thawing, samples were filtered using a 0.22um syringe filter to remove any infectious bacteria and kept on ice. Thirty cytokines and chemokines were assayed from the granuloma supernatants in duplicate using a ProcartaPlex multiplex immunoassay (Invitrogen) specific for nonhuman primate samples according to manufacturer’s instructions, with an additional dilution of the supplied standard curve to extend sample detection range. Multiplex results were read and analyzed by BioPlex reader (BioRad).

T cell responses in granulomas and uninvolved lung tissue

Flow cytometry was performed on individual granulomas from naive and reinfected macaques and a random sampling of lung tissue without any disease pathology from 6 Mtb infected macaques (with no reinfection) and an uninfected macaque. For granulomas, no restimulation was performed because of the high level of antigens in the granulomas, as previously noted[20]. Single cell suspension of the lung tissue was stimulated with peptide pools of Mtb specific antigens ESAT-6 and CFP-10 (10ug/ml of every peptide) in the presence of Brefeldin A (Golgiplug: BD biosciences) for 3.5 hours at 37˚C with 5% CO2. The cells were then stained for Viability marker (Invitrogen), surface and intracellular cytokine markers. Flow cytometry for cell surface markers for T cells included CD3 (clone SP34-2; BD Pharmingen), CD4 (Clone L200, BD Horizon) and CD8 (clone SK1, BD biosciences). Intracellular cytokine staining panel included pro-inflammatory cytokines: Th1 [IFN-γ (Clone B27), IL-2 (Clone: MQ1-17H12), TNF (Clone: MAB11)] and Th17 [IL-17 (Clone: eBio64CAP17)]. Data acquisition was performed using an LSR II (BD) and analyzed using FlowJo Software v.9.7 (Treestar Inc, Ashland, OR).

PBMC PCA analysis

A dimension reduction technique was performed on data extracted from PBMCs sampled just prior to reinfection (15 weeks post infection). Principal components analysis (PCA) was utilized here to investigate whether protected animals cluster separately from unprotected animals using blood signatures. Unfortunately, with such a small sample size (n = 8 macaques) PCA is not stable, so these results should be interpreted with caution.

Statistical analysis

Graphs were created in Graphpad Prism and JMP and all statistical analysis was performed in Graphpad Prism. In datasets that included zeroes, data were transformed by adding either 1 or 0.01 so that zeroes could be graphed on a log-scale. The D’Agostino and Pearson test was used.
to test for normality. If data were found to be normal, groups were compared using t-tests; otherwise, groups were compared using the non-parametric Mann-Whitney test for two groups or the Kruskal-Wallis test for three groups (with Dunn’s multiple comparison adjustment). All tests were two-sided with significance defined as $p < 0.05$.

**Supporting information**

S1 Fig. **There is no difference in the total numbers of granulomas that contain library B in reinfection.** (a) All new granulomas seen by PET CT in naïve and reinfected macaques at 4 weeks post-Library B infection. (b) All granulomas with Library B DNA in naïve or reinfected macaques. This count includes both newly established and pre-existing granulomas that contained library B in the reinfection cohort. Monkey ID: 19815 had no detectable library B in any tissue. (c) Number of new granulomas seen by PET CT established by library A or library B 4 weeks post-infection in the same animal. $p = 0.0156$, Wilcoxon-matched pairs signed rank test. (d) Number of granulomas seen by PET CT at 4–5 weeks post-Mtb Erdman infection between unvaccinated and BCG+H56 vaccinated macaques. Statistics: Mann-Whitney test.

S2 Fig. **Bacterial burden of granulomas is reduced over time.** CFU/granuloma was compared from Mtb Erdman infected macaques at 4–6, 11–12, and 20–24 weeks post-infection, using historical controls ($N = 17$). This is similar to data published in Lin, Ford, et al, but using a different set of monkeys. On the left side of the graph, CFU/granuloma of library A in our reinfected animals is shown ($N = 8$); these values are similar to the values of Erdman at 20–24 weeks.

S3 Fig. **Size (in mm) and FDG activity (SUVR) of individual library A granulomas in animals pre- and post- Library B reinfection and historical controls from 16–20 weeks post-infection.** (A) Granulomas do not change significantly in size after infection with Library B in the reinfection animals ($p = 0.2851$). Similarly, granulomas from animals infected with the Erdman strain do not change significantly in size from 16 to 20 (or 24) weeks post-infection ($p = 0.1848$). (B) Previously established (library A) granulomas do not significantly increase or decrease in FDG activity (SUVR) after reinfection with Library B ($p = 0.1351$). Likewise, in Erdman-infected historical controls, granulomas do not significantly change in SUVR ($p = 0.7878$) from 16 to 20 weeks post-infection. (C) Change in size (mm) per granuloma by animal. (D) Change in FDG activity (SUVR) per granuloma by animal. Each symbol represents a granuloma. Statistics: Wilcoxon matched-pairs signed rank test.

S4 Fig. **Uninvolved lung tissue from infected macaques has increased levels of chemokines and higher Mtb-specific T cell responses compared to lung tissue from an uninfected macaque.** A. Luminex analysis on supernatant from uninvolved (no granuloma) lung tissue was compared between reinfected macaques and an uninfected macaque. B. Using a separate set of macaque lung tissue (20–24 weeks post-infection but not reinjected), the T cell responses following ESAT-6 and CFP10 stimulation were assessed by flow cytometry, and compared to an uninfected macaque. No statistics were performed, due to the small sample size for the uninfected macaque lung.

S5 Fig. **Immunological and disease parameters do not segregate fully protected macaques from monkeys with new secondary granulomas.** (A) Principal Components Analysis of
PBMC data. Far left is the Eigenvalue Pareto Plot showing the cumulative percentage of variation accounted for in each principal component. Middle is a scatterplot of the first two components color-coded for animals with CFU-positive B granulomas (blue), and those without (red). There is no obvious clustering of these groups. Far right is a loading plot showing the correlations of the original variables to the first two principal components. B. Total thoracic CFU ($\log_{10}$) is similar between animals with and without CFU-positive Library B granulomas ($p = 0.7857$, Mann-Whitney test).

(TIF)

S1 Table. Parameters of macaque infection, serial imaging, bacterial burden and disease pathology.

(PDF)

S2 Table. Number of granulomas recovered with DNA identifiers for Library A or B.

(PDF)

S3 Table. Percentage of Lib. B of total lung bacterial burden.

(PDF)

S4 Table. Parameters of macaque infection, serial imaging, bacterial burden and disease pathology for BCG+H56 study.

(PDF)

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References


