Mechanisms of reactivation of latent tuberculosis infection due to SIV coinfection

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Mechanisms of reactivation of latent tuberculosis infection due to SIV coinfection

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**Graphical abstract**

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HIV is a major driver of tuberculosis (TB) reactivation. Depletion of CD4 T cells is assumed to be the basis behind TB reactivation in individuals with latent tuberculosis infection (LTBI) coinfected with HIV. Nonhuman primates (NHPs) coinfected with a mutant simian immunodeficiency virus (SIVAGY) that does not cause depletion of tissue CD4 T cells during infection failed to reactivate TB. To investigate the contribution of CD4 T cell depletion relative to other mechanisms of SIV-induced reactivation of LTBI, we used CD4RI antibody to deplete CD4 T cells in animals with LTBI without lentiviral infection. The mere depletion of CD4 T cells during LTBI was insufficient in generating reactivation of LTBI. Instead, direct cytotoxic effects of SIV resulting in chronic immune activation, along with the altered effector T cell phenotypes and dysregulated T cell homeostasis, were likely mediators of reactivation of LTBI. These results revealed important implications for TB control in HIV-coinfected individuals.

Introduction
Tuberculosis (TB) and HIV coinfection are a global epidemic (1). Approximately 90% of HIV-negative Mycobacterium tuberculosis-infected (Mtbb-infected) individuals lack clinical signs of TB and are considered to have latent TB infection (LTBI) (1, 2). Although the immune systems of exposed individuals successfully contain Mtbb infection in most cases, bacteria can persist within lung granulomas for long periods before reactivating to TB disease (3, 4). We seek to understand the mechanisms by which HIV coinfection reactivates TB using the Mtbb/SIV coinfection model in rhesus macaques.

CD4 T cell depletion is a hallmark of HIV and SIV infection, and is widely regarded as the primary cause of immunodeficiency. Previously, we identified Mtbb/SIV-coinfected primates that did not progress to TB despite productive coinfection, raising the possibility of CD4 T cell-independent mechanisms of LTBI control (5). To investigate the contribution of CD4 T cell depletion relative to other mechanisms of SIV-induced reactivation of LTBI, we used antibodies to deplete CD4 T cells and compared the outcomes to coinfection with pathogenic SIVAGY and the mutant virus, SIVAGY. Coinfection of Mtbb with pathogenic SIV, but without mutant or antibody-mediated CD4 T cell depletion, resulted in reactivation.

Results and Discussion
To assess the role of lung CD4 T cells in protecting against reactivation of LTBI, 39 Indian rhesus macaques were exposed to low-dose aerosol Mycobacterium tuberculosis (Mtbb) infection to establish TB latency (Figure 1A and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI125810DS1). At week 9 after TB infection, a subset of 16 macaques experiencing asymptomatic LTBI were coinfected with pathogenic SIVmac239. These animals exhibited 2 distinct outcomes: 9 animals developed signs of reactivated TB (reactivators) while 7 retained Mtbb latency despite productive SIV infection and peripheral blood viremia (nonreactivators) (5). To investigate the role of CD4 T cells in our low-dose aerosol model, we coinfected 6 macaques with a novel variant of pathogenic SIVmac29 molecular clone, SIVmac239AGYG (SIVAGY) (6), in which a deletion of 2 amino acids from a trafficking motif in the envelope gp41 cytoplasmic domain leads to viral replication, but does not deplete CD4 T cells in the periphery or in the lamina propria (ref. 7 and Supplemental Table 1). In addition, we used antibody-mediated depletion of CD4 T cells in 8 macaques with LTBI using CD4RI, which was administered every 2 weeks starting at week 9 after Mtbb infection (Figure 1A and Supplemental Table 1).

Importantly, SIVAGY-coinfected and CD4RI-administered macaques retained control of TB similar to nonreactivators. Specifically, only 1 of 8 CD4RI-administered nonhuman primates (NHPs) displayed symptomatology consistent with reactivated TB that necessitated a humane necropsy (Figure 1). SIVAGY-coinfected and CD4RI-administered macaques showed normal serum C-reactive protein (CRP) levels over time (Supplemen-
Figure 1. Comparison of CD4+ T cell–sparing SIVmac239AGY and antibody-mediated CD4+ T cell depletion using CD4R1 in Mtb-infected NHPs. (A) Study outline. (B) Serum CRP levels at necropsy. (C) Percentage of temperature increase (°C) and (D) percentage of weight loss (kg) compared with baseline. (E) Ratio of neutrophils to lymphocytes in clinical blood. Bacterial burden at necropsy measured in (F) BAL, (G) lung, and (H) bronchial lymph nodes. (I) CXR scores established by veterinary clinicians for pneumonia. (J) Percentage of lung involvement determined by board-certified veterinary pathologists. (K) Peak plasma viral loads (PVLs) measured at peak. *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001, 1-way ANOVA with Tukey’s multiple testing correction. C–E represent mean, and B and F–J represent mean ± SEM.

tal Figure 1A) and at endpoint (Figure 1B), comparable to LTBI and nonreactivators and statistically different from reactivators. These animals maintained relatively normal body temperatures (Figure 1C) and weights (Figure 1D). Reactivators, unlike all other groups, had a lower ratio of neutrophils/lymphocytes after SIV coinfecion at week 9 (Figure 1E). SIVAGY-coinfected and CD4R1–administered NHPs had lower numbers of viable Mtb in their bronchoalveolar lavage (BAL) fluid throughout infection (Supplemental Figure 1B), and significantly lower viable Mtb in their BAL at endpoint (Figure 1F). Similarly, both experimental groups harbored low lung (Figure 1G), bronchial lymph node (Figure 1H), spleen (Supplemental Figure 1C), liver (Supplemental Figure 1D), and kidney (Supplemental Figure 1E) bacterial burdens, comparable to the LTBI and nonreactivators. Both experimental groups possessed significantly lower viable Mtb in all tissues at necropsy compared with reactivators. Finally, virtually no tuberculous lung pathology was observed in SIVAGY-coinfected NHPs, demonstrating that coinfection with this virus failed to reactivate LTBI (Figure 1, I and J). One of 8 CD4R1–administered NHPs with LTBI did reactivate, displaying an elevated CRP at necropsy (Figure 1B) and chest x-ray (CXR) score (Figure 1I). Measurement of peripheral viremia in coinfeconed animals suggested that
SIVAGY replicated to comparable levels in the acute phase and established similar set points (Figure 1K). Although significantly lower peripheral viremia was observed at peak in our SIVAGY-coinfected NHPs compared with SIVmac239-coinfected reactivators and nonreactivators, this is not unexpected as rhesus macaques infected with SIVAGY often have variable viremia (8, 9).

NHPs with LTBI/SIVAGY coinfection did not exhibit a significant decline in CD4⁺ T cell levels in peripheral blood (Figure 2A and Supplemental Figure 2, A and C) or BAL (Figure 2B and Supplemental Figure 2, B and D). This was in stark contrast to animals infected with pathogenic SIV (Figure 2, A and B), consistent with previous results (5, 10). Although a significant reduction in CD4⁺ T cells was observed in the lungs (Figure 2C) of SIVAGY-coinfected NHPs, an insignificant reduction was observed in the total CD4⁺ T cell compartment (Supplemental Figure 2E). Previously, SIVAGY had been shown to replicate in the plasma and lymphoid tissues, but to spare gut mucosal tissues (7, 8).

To our knowledge, this was a first-time evaluation of lung CD4⁺ T cell populations in SIVAGY-infected NHPs, so this was a novel finding. There may be sufficient lymphoid tissue in the lungs that SIVAGY was able to replicate nearby, perhaps in inducible bronchus-associated lymphoid tissue (IBALT) (5), which led to CD4⁺ T cell depletion during SIVAGY coinfection. Significant depletion of CD4⁺ T cells was observed in the peripheral blood (Figure 2D), BAL (Figure 2E), and lungs (Figure 2F) of CD4R1-administered NHPs, relative to LTBI NHPs. The levels of CD4⁺ T cell depletion were comparable to SIVmac239-infected animals and the differences in CD4⁺ T cell populations between NHPs administered CD4R1 and NHPs coinfected with SIVmac239 were not significant in the BAL or lungs (Figure 2, E and F), regardless of their TB outcome. This result was confirmed by multilabel confocal immunofluorescence microscopy (Figure 2, G–K). Thus, our results show that coinfection of macaques with LTBI with a nonpathogenic virus that fails to deplete CD4⁺ T cells (SIVAGY) and depletion of CD4⁺ T cells using antibodies does not result in similar rates of TB reactivation observed upon coinfection with pathogenic SIVmac239.

As early as week 2 after SIV challenge or CD4R1 administration, NHPs displayed significant changes in the distribution of their residual CD4⁺ T cells. Mtb infection alone can impact the distribution of effector and memory cells, and memory CD4⁺ T cells are associated with protection (11–13). We compared 4 experimental groups of NHPs: our SIVmac239-coinfection reactivators and nonreactivators, SIVAGY-coinfected, and CD4R1-administered NHPs. Although CD4⁺ T cells were massively depleted in both experimental groups of NHPs, it was vital
Figure 3. Peripheral and BAL CD4+ T cells reflect a proportional shift toward effector profile during CD4R1 administration. The proportions of residual effector (CD95+CD28-, circles), memory (CD95+CD28-, squares), and naive (CD95+CD28-, triangles) (A) CD4+ and (B) CD8+ T cells in the peripheral blood and (C) CD4+ and (D) CD8+ T cells in the BAL of SIVmac239-coinfected (n = 15, black) and CD4R1-administered NHPs (n = 8, red) compared with LTBI NHPs (n = 10, blue) at week 11 after TB. *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001, 2-way ANOVA with Tukey’s multiple testing correction. BAL (E) effector and (F) memory CD4+ T cells quantification every 4 weeks starting at week 7 after TB. Coinfection and antibody administration shown by the dotted line. (G) qRT-PCR analysis of IL-2, IL-12B, and IL-17A collected from BAL cells at week 11 and week 15 after TB from SIV reactivators (n = 3, yellow) and CD4R1-administered NHPs (n = 3, red). *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001, multiple t tests with Holm-Sidak method for multiple comparison correction. Data represent mean ± SEM.

to understand the contribution of the remaining CD4+ T cells to the host’s immune defenses. Coinfection with SIVmac239 resulted in a significant reduction of memory (CD95+CD28-) CD4+ T cells, which were replaced by naive (CD95+CD28-), not effector (CD95+CD28-), CD4+ T cells (Figure 3A). Most remaining CD4+ T cells in the blood of SIV-coinfected animals were naive, consistent with findings in human studies that Mtb/ HIV-coinfected patients harbor less differentiated CD4+ memory T cells compared with TB-only patients (14). HIV infection can lead to an activated and exhausted memory CD4+ T cell phenotype (15), which likely contributes to the more significant loss of memory CD4+ T cells. Compared with animals with LTBI, CD4R1-administered NHPs exhibited significantly higher frequencies of effector and reduced naive and memory CD4+ T cells in the peripheral blood (Figure 3A). In the peripheral blood, CD8+ T cells did not differ between groups (Figure 3B). In the BAL, changes were observed only in the CD4+ (Figure 3C) and not the CD8+ T cell compartment (Figure 3D). Memory CD4+ T
cells were replaced by naive cells upon SIV coinfection in the BAL (Figure 3C). Interestingly, the distribution of BAL CD4+ T cells following CD4+ T cell depletion by SIVmac239 coinfection did not significantly differ from the distributions observed during LTBI, indicating that the virus affected all compartments equally within the BAL. BAL effector CD4+ T cell populations were significantly increased in CD4R1-administered NHPs compared with both LTBI and SIVmac239-coinfected NHPs, whereas memory CD4+ T cell populations were significantly reduced (Figure 3C). Throughout CD4R1 administration, CD4+ T cells in the BAL reflected a bias toward effector (Figure 3E) rather than memory (Figure 3F) populations. Greater proportions of effector T cells in the lungs of CD4R1-administered, versus SIV-coinfected, NHPs strongly suggest that early
pathogenic events associated with SIV coinfection suppress the natural homeostatically driven CD4+ T cell proliferation and downstream effector T cell functions. As these differences were unique to the BAL and not the lung tissue (Supplemental Figure 3, A and B), they suggest that T cells in the lung, and not circulating BAL cells, are capable of retaining their typical functions.

Gene-expression analyses in the BAL from SIV-coinfected versus CD4R1-administered NHPs confirmed our assertion that CD4+ T cell maturation states differed in these 2 groups. Real-time RT-PCR analysis revealed that at week 11 and 15 after Mb, NHPs administered CD4R1 had significantly higher levels of transcripts involved in T cell activation, differentiation, and function (e.g., IL-2, key signal for Th1 differentiation; IL-12B, important for sustaining memory effector Th1 responses against Mb; and IL-17A, expressed on Th17+ cells) compared with SIV reactivators (Figure 3G). These cytokines indicate that although CD4R1 depleted the CD4+ T cell compartment, it did not inhibit T cell maturation and signaling. On the contrary, SIV coinfection inhibited T cell maturation and signaling.

Gene-enrichment analyses on lung transcriptome confirmed that SIV coinfect reactivators experienced chronic immune activation and a cytokine storm. CD4+ T cell homeostasis was enhanced in CD4R1-administered NHPs and unperturbed during SIV coinfection. The gene terms “immunoglobulin/major histocompatibility complex (MHC),” “MHC class I/class II antigen recognition protein,” etc., were significantly enriched only in CD4R1-administered NHPs (Figure 4A and Supplemental Table 2, InterPro). Several categories encoding proteins interacting with RAS (Figure 4A), involved in cell signaling pathways that control cell growth and cell death, were only significantly enriched in CD4R1-administered NHPs. This underscores that normal immune homeostasis is observed in CD4R1-administered NHPs, despite severe lymphopenia. Categories such as “inflammatory response,” “innate immunity,” and “endoplasmic reticulum” were only enriched in SIV reactivators, suggesting chronic immune activation and an ensuing cytokine storm in that group (Figure 4B). Unsurprisingly, SIV reactivators were characterized by the enrichment of gene categories corresponding to chronic immune activation and inflammation, including “TNF-signalizing” and “Fc-gamma receptor mediated phagocytosis” (Figure 4C). Reactivator lungs substantially expressed high levels of genes involved in proinflammatory cytokine/chemokine signaling (CXCL12, IL6), myeloid cell/phagocyte/complement/innate immune signaling (CD68, CFH, C1S, TXNDC11), matrix degradation critical for granuloma formation (IGFBP7), and antigen presentation (HLA-B) (Supplemental Table 2). Genes belonging to the pathway “regulation of lymphocyte differentiation,” including NDFIP1, FAS, and RNF41, were induced in lungs of reactivators. The genes only expressed in nonreactivators’ lungs included those involved in the restoration of normal lung and cytoskeletal function (WWOX, TRIOBP, PEAK3), B cell function (SPANXAI) (16), repair of acute tissue injury (MYLK2) (17), inhibition of cytokine storm (TIP39, SLC8A2) (18), viral restriction (APOBEC3) (19), induction of apoptosis (IFT57), and induction of robust cellular immune responses (CLECG5/KLRG2, BTNL2A3, TSPEAR, MRAI).

Real-time RT-PCR and multiplex cytokine analysis strongly confirmed genome-wide transcriptome results. Cytokines associated with inflammation, including TNF-α, IL1-α, IL1-β, and IL-6, were highly induced in the lungs of SIV-coinfected reactivators compared with CD4R1-administered NHPs (Figure 4D). Multiplex analysis supported this trend, showing that in the BAL supernatant at necropsy, TNF-α, IL1-α, IL1-β, and IL-6 proteins were more abundant in SIV-coinfected reactivators (Figure 4E). Significantly increased IFN-γ was detected in the plasma (Supplemental Figure 4A) and BAL (Supplemental Figure 4B) of reactivators compared with other groups. Other markers of increased cell signaling and migration associated with inflammatory responses were significantly increased in the plasma (CXCL9; Supplemental Figure 4C) or BAL (CXCL13 [Supplemental Figure 4D], eotaxin [Supplemental Figure 4E], I-TAC [Supplemental Figure 4F], and IL-6 [Supplemental Figure 4G]) of reactivators. SIVA6Y-coinfected NHPs had significantly increased IL-8 and MPI1-β in the plasma (Supplemental Figure 4, H and I) compared with all other groups, and LTBI NHPs had significantly increased eotaxin in the plasma (Supplemental Figure 4J) and SDF-1α in BAL (Supplemental Figure 4K).

Chronic immune activation in SIV-coinfected animals was supported by immunophenotyping. T cells in the lungs of reactivators had a significantly more activated phenotype (HLA-DR+) compared with those from NHPs that received CD4R1 (Figure 4F). HLA-DR+ (Figure 4G) and HLA-DR+ (Figure 4H) T cells in reactivators exhibited significantly higher CCR5 (recruitment) and CD69 (early activation) markers compared with CD4R1 administration. This suggests higher recruitment and entrapment of activated lymphocytes at the primary site of infection in reactivators. CCR5-associated recruitment to the immune synapse induces stronger T cell-APC attraction, which results in reduced T cell responsiveness to chemokines (20). This response is typically followed by increased T cell proliferation and upregulated effector responses (20, 21), which was observed through increased CD69 expression by SIV reactivators. Furthermore, CCR7+ T cell populations were significantly reduced in reactivators compared with CD4R1-administered NHPs (Figure 4, G and H), suggesting that migration to secondary lymph nodes was impaired in T cells from the lungs of reactivators (22). Significantly reduced PD-1 expression in reactivators compared with CD4R1-administered NHPs indicated that HLA-DR+ (Figure 4G), but not HLA-DR+ (Figure 4H), T cells in reactivators were subject to regulation, strongly supporting a pattern of chronic immune activation in reactivators. As this pattern was also observed in the CD8+ T cell compartment (Supplemental Figure 5, A–C), SIV-mediated CD4+ T cell depletion is likely not the sole cause of immune imbalance leading to chronic activation, as T cell recruitment, activation, and homeostatic regulation was maintained in NHPs with antibody-mediated CD4+ T cell depletion. As can be seen in the reactivators with reduced PD-1 expression, some of these inhibitory responses are not being maintained, thus allowing increased immune activation and the cytokine storm seen in our transcriptomics data to occur.

Although it is known that the functions of mycobacterial-specific CD4+ T cells are impaired during HIV infection, the immunological mechanisms underlying how CD4+ T cells contribute to maintaining immune control during LTBI and how HIV perturbs
this control are not understood (23). Not all NHPs with LTBI/SIV coinfection exhibit reactivation despite comparable CD4+ T cell depletion (5), indicating that CD4+ T cell-independent mechanisms exist for the maintenance of LTBI. Coinfection of animals with SIVAGY showed that CD4+ T cells are important for preventing reactivation during Mtb/SIV. However, the mere depletion of CD4+ T cells is insufficient for reactivation of LTBI, since only 1 of 8 CD4RI-administered NHPs developed overt TB disease. The key difference in the lungs of primates where lung CD4+ T cells were depleted by antibody or by pathogenic SIV was that the signatures of chronic immune activation and blockade of normal lymphocytic homeostasis were detected in the lungs of only reactivators.

Our results show that lung-specific chronic immune activation may be important for SIV-mediated (and perhaps HIV-mediated) reactivation of LTBI. Since CD4RI administration resulted in greater depletion of memory CD4+ T cells and preservation of effector CD4+ T cells, it is possible that this homeostasis drives the lack of reactivation in these NHPs. It is further acknowledged that the lack of chronic immune activation may be a byproduct of the corresponding lack of reactivation. Importantly, it appears that SIV pathogenicity drives mechanisms such as chronic immune activation that disturb the normal homeostasis of the effector CD4+ population. Understanding how SIV and chronic immune activation impact immunity to TB could lead to the development of new treatment paradigms aimed at the cure of Mtb/HIV coinfection.

Methods

Study approval. All procedures were approved by the Tulane National Primate Research Center Institutional Animal Care and Use Committee and Tulane Institutional Biosafety Committee.

Author contributions

ANB, SAK, and DK designed the study. ANB, AC, TWF, XA, BT, MKG, and NG performed experiments. ANB, AC, TWF, XA, DKS, THL, SAK, and DK analyzed results. JAH, JR, SM, and MA interpreted results and provided vital reagents. ANB, JR, SM, MA, SAK, and DK prepared the manuscript.

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