MR1-restricted mucosal-associated invariant T (MAIT) cells respond to mycobacterial vaccination and infection in nonhuman primates

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Studies on mucosal-associated invariant T cells (MAITs) in nonhuman primates (NHP), a physiologically relevant model of human immunity, are handicapped due to a lack of macaque MAIT-specific reagents. Here we show that while MR1 ligand-contact residues are conserved between human and multiple NHP species, three T-cell receptor contact-residue mutations in NHP MR1 diminish binding of human MR1 tetramers to macaque MAITs. Construction of naturally loaded macaque MR1 tetramers facilitated identification and characterization of macaque MR1-binding ligands and MAITs, both of which mirrored their human counterparts. Using the macaque MR1 tetramer we show that NHP MAITs activated in vivo in response to both Bacillus Calmette-Guerin vaccination and Mycobacterium tuberculosis infection. These results demonstrate that NHP and human MR1 and MAITs function analogously, and establish a preclinical animal model to test MAIT-targeted vaccines and therapeutics for human infectious and autoimmune disease.

INTRODUCTION

Mucosal-associated invariant T cells (MAITs) are non-classical T cells that recognize microbial-derived vitamin B metabolites in the context of major histocompatibility complex related protein 1 (MR1).1,2 MAITs comprise up to 10% of CD8+ T cells in the peripheral blood of humans and accumulate in tissues like the liver, lung, and gut mucosa where these cells rapidly respond to microbial threats.3–5 MAIT cells utilize a semi-invariant T-cell receptor (TCR) to probe ligands bound by the highly conserved MR1 molecule. Like classical major histocompatibility complex class I molecules, MR1 is expressed by all nucleated cells, but its surface expression is often limited in the absence of appropriate ligand.6 MAIT TCRs generally utilize a TRAV1-2–TRAJ33 alpha chain combined with a more diverse array of beta chains, although novel MAIT cell populations utilizing other TCRs than TRAV1-2 have recently been identified.7,8 Upon recognition of their cognate antigen, MAITs secrete inflammatory cytokines like interferon-γ (IFNγ) and tumor necrosis factor-α (TNFα). Moreover, MAITs are capable of this effector function immediately upon egress from the thymus.9 Taken together, these features indicate that MAIT cells are an innate-like T-cell subset primed to respond rapidly to bacterial pathogens.

Indeed, these innate like T cells play a role in the initial host response to Mycobacterium tuberculosis (Mtb) and other microbial pathogens including fungi and yeast.10–12 MAITs
RESULTS
Nonhuman primate MR1 ligand identification
Mouse and human MR1 tetramers have facilitated the characterization of MAITs in each species. Given that many human-specific reagents cross-react with the corresponding target on macaque cells, we were surprised to find that human MR1 tetramers folded with multiple ligands, as described below, exhibited a limited ability to stain rhesus macaque MAITs (see discussion of Figure 1e). To assess whether differences in the MR1 coding sequence might explain these results, we sequenced the functionally critical MR1 \( \alpha_1 \) and \( \alpha_2 \) domains from human, chimpanzee, rhesus macaque, cynomolgus macaque, Japanese macaque, pigtailed macaque, sooty mangabey, and baboon. Sequences were highly conserved among the different species, with greater than 95% nucleotide identity to human MR1 for all species examined (see Supplementary Figure S1 online). Despite this high level of sequence conservation, all nonhuman primate species, except chimpanzees, possessed three amino acid substitutions in previously described MAIT TCR contact residues (Figures 1a,b). In contrast, sites known to form hydrogen bonds with the well-characterized MR1 ligand rRL-6-CH_2OH were conserved in every species. These results suggested that the human MR1 tetramer might not effectively bind macaque MAITs due to variation at residues critical for TCR docking, but MR1 molecules from distinct primate species should bind and present identical ligands.

To test these hypotheses, we synthesized human and macaque MR1 tetramers using insect cells and the baculovirus expression system (see Supplementary Figure S2). The MR1 molecule was naturally loaded with ligand by infecting MR1-expressing Hi5 cells with *Escherichia coli* (*E. coli*), and the secreted MR1 molecule subsequently purified via affinity chromatography. Because there is an array of potential ligands using this system, we hypothesized that we could use this monomer to identify the nearly full complement of putative MR1 ligands. Indeed, using liquid chromatography-mass spectrometry to identify macaque MR1 ligands as previously described, we identified approximately 40 putative MR1 ligands (data not shown).

Although we are currently unable to assign structures to the majority of these ligands without confirmed spectra of the synthetic molecule, we identified the known activating ligand rRL-6-CH_2OH and a second ligand, 7-hydroxy-6-methyl-8-ribityl lumazine (Figures 1c,d). rRL-6-CH2OH and 5-OP-RU, both MAIT cell-activating MR1 ligands, have the same chemical formula \( \text{C}_{12}\text{H}_{19}\text{N}_{4}\text{O}_{7} \) and are thus isoelemental. Additionally, these compounds have indistinguishable MS2 fragmentation spectra (PMID 23051753, 24695216, and 24108697). Thus, it is possible that the compound identified here as rRL-6-CH2OH is 5-OP-RU or a mixture of both.

Nonhuman primate MR1 tetramers exhibit crossreactive staining
We then used this newly synthesized human tetramer to stain peripheral blood mononuclear cell (PBMC) samples from the same primate species used in our MR1 sequencing analysis above and found that the human tetramer was not universally crossreactive and strictly stained human and chimpanzee MAITs (Figure 1e). This staining pattern is consistent with the results from our MR1 sequencing in which human and chimpanzee MR1 sequences are identical at the previously noted TCR contact residues (Figures 1a,b). In contrast to the results from the human MR1 tetramer, the macaque tetramer effectively stained MAITs from all eight species tested. The ability of macaque MR1 to stain human and chimpanzee MAITs is likely due to the amino acid substitutions at positions 72 and 151, residues that are known to mediate MR1 xenoreactivity. Although the anti-human \( V_{\alpha}7.2 \) antibody exhibited variable crossreactivity against nonhuman primate species, both macaque and human MR1 tetramer cells were \( V_{\alpha}7.2 + \) in humans and chimpanzees confirming the tetramer specifically identifies MAITs (Figure 1e; see Supplementary Figure S3a). Indeed, the human and macaque MR1 tetramer cells were also CD161 + in humans, although the anti-human CD161 antibody did not sufficiently crossreact with nonhuman primate CD161 to function as a marker in these species (see Supplementary Figure S3b). The unloaded macaque MR1 tetramer control did not stain cells from any species, confirming that an appropriate ligand is required for the macaque MR1–TCR interaction (Figure 1e). The MR1 tetramer staining in human samples mirrors previous studies in which MAITs are present in both the CD8 + and the CD8 – CD4 – populations; however, this double-negative MAIT population was absent in every nonhuman primate species tested (see Supplementary Figure S3). These results provide the first demonstration of staining by a macaque MR1 tetramer and explain the limited crossreactivity observed with the human tetramer. Moreover, they demonstrate that the MR1 gene is highly conserved across several species but diverges at critical residues that affect TCR–MR1 docking. Finally, because this tetramer is loaded with a mixture of ligands, it is possible that it will identify additional MAIT populations that would be missed by a single ligand-loaded tetramer.

Functional and phenotypic characterization of nonhuman primate MAITs
Next, we sought to phenotypically and functionally characterize MR1 tetramer + cells from rhesus macaques. In order to adapt the single-cell, unbiased TCR\( \alpha_{2} \beta \) analysis technique to macaques, we first extracted and characterized the TRA
gene sequences from the rhesus macaque genome based on homology to the human TRA genes (see Supplementary Figures S4 and S5). Using these data and the TRB gene sequences we previously extracted from the rhesus macaque genome, we then performed paired sequencing of CDR3α and CDR3β from MR1 tetramer + cells sorted from the liver and blood of a healthy macaque. Consistent with results from humans, we found that the majority of MAITs utilized TRAV1-2 in conjunction with TRAJ33 (Figure 2a). Additionally, most TCRα chains contained the canonical ligand-binding
tyrosine (Figure 2b and see Supplementary Figure S6).27 Previous studies of TCRz sequences across multiple species have demonstrated that these clonotypes can be efficiently produced by the gene recombination process due to “overlap” between TRAV and TRAJ genes.28 This means that the germline-encoded nucleotide sequences for these clonotypes can be produced by a variety of mechanisms. TRAV19 TCRz chains were also enriched in our sequence analysis (see Supplementary Figure S6). These TRAV19 sequences contained a common “NE” amino acid motif that may indicate shared recognition of the same MR1–ligand complex. This identification of TRAV1-2 negative MAITs is consistent with recent studies in humans.7,8 The TCRz chains were relatively promiscuous in pairing with a variety of TCRβ chains (see Supplementary Figure S6a). Tetramer + cells from the liver exhibited reduced diversity of TRAV and TRAJ usage compared with the blood MAITs (Figure 2a). Additionally, there was liver-specific usage of TCRβ V regions suggesting tissue-specific differences in macaque MAITs parallel to previous results examining human MAIT TCRs26 (data not shown). These differences may be due, in part, to local clonal expansion or differential establishment of tissue residency, but will require additional in-depth study.

Previous studies examining human and mouse MAITs demonstrated that these cells express PLZF, a regulator of innate like T-cell development, and RORγt, a master regulator of TH17 differentiation.7,15 A proportion of MAITs have also been described as T-bet +, a marker associated with IFNγ production. We found that the majority of MR1 tetramer+ cells express PLZF and RORγt (Figures 2d,e). MAITs expressed variable levels of T-bet that were indistinguishable from MR1 tetramer− CD8+ T cells, consistent with previous studies characterizing MAITs in mice and humans.19 These data suggest that macaque MAITs might also produce higher levels of IFNγ compared with mice, and further confirm that the macaque MR1 tetramer is identifying macaque MAITs. Finally, we assessed MR1 tetramer + cell function in an E. coli-based stimulation assay.3 MAIT activation is MR1 dependent, and should not be blocked by the pan-major histocompatibility complex-I blocking antibody W6/32.29 MR1 tetramer + cells produced significant amounts of IFNγ and TNFα, and upregulated CD69 in response to E. coli (Figures 2f,g). This activation was reduced by the MR1 blockade, but not by the isotype control or pan-major histocompatibility complex-I blocking antibody (Figure 2f). In contrast, MR1 tetramer − CD8+ cells produced very little IFNγ and/or TNFα in response to E. coli stimulation and this was not significantly reduced by anti-MR1 antibody (Figure 2g). Interestingly, MAITs did not produce significant amounts of IL-17 in these assays (data not shown). These results demonstrate that macaque MR1 tetrarsers specifically stain macaque MAITs and macaque MAITs are activated in response to bacteria in an MR1-dependent fashion.

Next, we assessed the distribution and phenotype of MAITs across 23 different tissues in 11 Indian rhesus macaques. We found that MAITs were most abundant in the epithelium of the lung (measured by bronchoalveolar lavage, BAL) and in liver (Figures 3a,b). We did not detect high frequencies of MAITs in lymph nodes or other primary and secondary lymphoid organs. We also examined MAIT activation by Ki-67 and CD69 staining, and observed nearly equivalent frequencies of Ki-67 + MAITs and non-MAIT CD8+ T cells in most tissues (Figure 3c). We focused our statistical analysis on BAL, liver, and blood, and found small differences in Ki-67 expression in BAL and liver compared with non-MAIT CD8+ T cells. In contrast, MAITs expressed higher levels of CD69 than non-MAITs in most primary lymphoid tissues, secondary lymphoid tissues and select extralymphoid tissues (Figure 3d). We observed significantly more CD69 + MAITs than non-MAIT CD8+ T cells in the liver while these frequencies were essentially equivalent in the blood and BAL. These results suggest that MAITs may actively encounter and respond to bacterial antigens in the liver, particularly as the liver filters blood from the intestinal tract.60 We also examined the memory phenotype of MAITs in these tissues and found that the majority of MAITs were CD28+CD95+ memory T cells (see Supplementary Figure S7a). Further analysis revealed that MAITs in the primary and secondary lymphoid tissues, particularly the lymph nodes, exhibited a mostly central memory (CCR7+, CD28+) phenotype while in extralymphoid tissues they exhibited a transitional effector memory (CCR7−, CD28+) or effector memory (CCR7−, CD28−) phenotype (see Supplementary Figure S7b). Human blood MAITs are traditionally considered effector memory T cells as defined by CD45RA and CD62L expression.31 While

Figure 1 MR1 sequence polymorphisms explain species-specific tetramer reactivity. (a) The amino acid alignment of the MR1 x1 and x2 domains is displayed. Identity to the human reference sequence is indicated by a “.". Differences are indicated by the specific amino acid. Lowercase indicates heterozygosity. (b) A ribbon diagram of the x1 and x2 domains of the MR1 molecule (from above). TCR contact residues are highlighted in blue. (c) Left panel—negative ion MS1 extracted ion chromatogram of rRL-6-CH2OH [M–H]− = 329.1100) from macaque MR1 co-cultured with E. coli (black) or the macaque MR1 mock control (blue). Middle panel—composite MS1 spectra from 11.25–12.00 min of ion 329.1100. The black trace is macaque MR1 co-cultured with Escherichia coli and the macaque MR1 mock control is blue. Right panel—fragment spectrum of ion 329.1105 at 11.51 min from the E. coli sample. The precursor and product ions are consistent with the published fragment spectra for synthetic rRL-6-CH2OH. Possible fragment ion structures are shown with their corresponding chemical formula, calculated mass, and observed ion. (d) Left panel—negative ion MS1 extracted ion chromatogram of HMRL [M–H]− = 327.0946) from macaque MR1 co-cultured with E. coli (black) or the macaque MR1 mock control (blue). Second from left panel—Composite negative ion MS1 spectra from 10.00–10.80 min of ion 327.0948. The black trace is macaque MR1 co-cultured with E. coli and the macaque MR1 mock control is blue. Second from right panel—Fragment spectrum of ion 327.0948 at 10.38 min from the E. coli sample. The precursor and product ions are consistent with the fragment spectra of synthetic HMRL. Possible fragment ion structures are shown with their corresponding chemical formula, calculated mass, and observed ion. Right panel—Fragment spectrum of synthetic HMRL. (e) Tetramer staining of PBMC from each of the indicated species is plotted against staining by the anti-Vx7.2 antibody. Plots progressively gated on lymphocytes, singlets, live, CD3+, CD8+ cells. HMRL, 7-hydroxy-6-methyl-8-ribityl lumazine; MR1, major histocompatibility complex related protein 1.
Figure 2 Macaque MR1 tetramer + cells are phenotypically and functionally similar to human and mouse MAITs. (a) TCRβ V (top) and J (bottom) region usage from MR1 tetramer + cells from both the blood (n = 61 cells) and liver (n = 44 cells) of a healthy macaque. (b) Select CDR3α sequences of blood MAITs. The canonical ligand-binding tyrosine is noted in bold/underlined. (c) TCRβ V (top) and J (bottom) region usage from MR1 tetramer + cells from blood (n = 73 cells) and liver (n = 64 cells). TRBV4-1, 4-2, and 4-3 were combined together since they are indistinguishable based on the sequence length. (d) Representative nuclear staining for PLZF, RORγt, and T-bet. Plots were progressively gated on lymphocytes, singlets, live, CD3+, and CD8+ cells. (e) PBMC from eight animals were stained for the nuclear proteins PLZF, RORγt, and T-bet. Mean, s.e.m., and P-values (Wilcoxon rank-sum test) are shown. (f) Representative staining of MAITs (top) and non-MAIT CD8+ T cells (bottom) in a functional assay. (g) Functional assay results examining activation of MAITs from the spleen of eight animals to PFA fixed E. coli. Cells were stained with IFNγ and TNFα (left) and CD69 (right). Results from MR1 tetramer-positive cells are shown in blue and MR1 tetramer negative cells in red. All blocking antibodies were added at a concentration of 100 μg·ml⁻¹. Conditions were plated in duplicate and averaged. Mean, s.e.m. and P-values (Kruskal Wallis and Dunn’s multiple comparison post-test) are shown. IFNγ, interferon-γ; MAITs, mucosal-associated invariant T cells; MR1, major histocompatibility complex related protein 1; TNFα, tumor necrosis factor-α.

T-cell memory markers differ between the two species, we find that macaque MAITs exhibit a spectrum of memory phenotypes largely dependent on their anatomical location. It is currently unclear why a higher frequency of macaque MAITs exhibit a central memory phenotype compared with human MAITs. These results highlight the differences in MAITs.

ARTICLES

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among distinct tissues, possibly reflecting differences in antigen exposure at these sites.

**MAITs respond to mycobacterial infection in vivo**

We next utilized the macaque model to examine the functional response of MAITs to bacterial challenge in vivo. Following intradermal vaccination with Bacillus Calmette-Guerin (BCG) to the left and right chest of each animal, we failed to observe a significant and consistent change in MAIT frequency in peripheral blood (**Figure 4a**). In contrast, MAITs became specifically activated in response to BCG vaccination in the blood at days 14 and 21 post-vaccination compared with...
Nine weeks after the initial BCG vaccination, we re-vaccinated the macaques, and 2 weeks later collected skin from the sites of vaccination on the chest (pectoral) and skin from the inner thighs (inguinal) as a non-vaccination control site, as well as spleen, blood, BAL, axillary lymph node, and inguinal lymph node. While not significant, we observed a trend toward higher frequencies of MAITs in skin at the site of vaccination compared with skin from a distal site (Figures 4c,d). Further, we measured significantly higher frequencies of Ki-67 + MAITs in the vaccine site compared with the distal inguinal skin (Figure 4e). Moreover, MAITs from the vaccine site were also significantly more activated than non-MAIT CD8 + T cells from the same site (Figure 4g). In contrast, non-MAIT CD8 + T cells tended to be more activated in the distal inguinal skin (Figure 4g). Measurements of activated MAITs in the draining axillary lymph node compared with the distal inguinal lymph nodes mirrored those in the vaccination site skin and distal skin (Figures 4e,g). While contemporaneous uninfected controls were not available, the Ki-67 expression in MAITs from axillary lymph nodes of BCG infected macaques was nearly 10-fold higher than MAITs from axillary lymph nodes of healthy, non-vaccinated animals (Figures 3c and 4e). Finally, we also found a higher frequency of granzyme B-loaded MAITs in skin at the vaccination site and the draining axillary lymph node compared with the distal skin and lymph nodes.

Figure 4  MAITs respond to bacterial pathogens in vivo. Four macaques were challenged with BCG intradermally and monitored for 6 weeks. (a) Frequency of MAITs in the blood of BCG vaccinated macaques. Mean and s.e.m. are shown. Kruskal Wallis test revealed no significant differences in MAIT frequencies. (b) Frequency of Ki-67 + MAITs (blue) and non-MAIT CD8 + T cells (red) in the blood. Mean, s.e.m, and P-values (Wilcoxon rank-sum test) are shown for (d,g). (c) Representative MR1-tetramer, Ki-67, and GzmB staining from the pectoral (top, vaccine site) and inguinal (bottom, non-vaccine site) skin sections. (d) Frequency of MAITs of CD8 + T cells in the chest and thigh skin sections (left) and in the axillary and inguinal lymph nodes (right). (e) Percentage of Ki-67 + MAITs in the chest and thigh skin (left) and axillary and inguinal lymph nodes (right). (f) Frequency of GzmB + MAITs in the skin (left) and lymph nodes (right). (g) Comparison of Ki-67 + frequencies among MAITs and non-MAIT CD8 + T cells in the chest (left), thigh (middle), and axillary lymph node (right). Mean, s.e.m., and P-values (Friedman test with Dunn’s multiple comparison post-test) are shown for (d,g). (h) Ki-67 + frequency of MAITs (blue) and not MAITs (red) after re-vaccination. (i) MAIT frequencies of CD8 + T cells in blood after TB infection. (j) MAIT and non-MAIT CD8 + T cell Ki-67 frequencies after Mtbi infection. Mean, s.e.m., and P-values (Friedman test with Dunn’s multiple comparison post-test) are shown for (d-g).
(Figure 4f), demonstrating that BCG vaccination licenses MAITs via granzyme B.\(^3\) Longitudinal analysis after the revaccination showed that, while not significant, blood MAITs were activated again at day 14 post-vaccination (Figure 4h). These results demonstrate that MAITs are specifically activated in response to bacterial threats in the nonhuman primate. Additionally, these data demonstrate the need to examine MAIT cells in the relevant tissues because measurements from blood alone may miss MAIT cell activation, and further reveal the unique advantages of the nonhuman primate model, where longitudinal tissue biopsies may be taken.

Finally, we examined the MAIT response to Mt infection. We monitored nine intrabronchially Mt-infected macaques for 8 weeks post challenge for changes in MAIT frequency and activation in blood. Similar to BCG vaccination, we did not observe a consistent change in MAIT frequencies in peripheral blood (Figure 4i). It is possible that MAITs expanded locally in the lung in response to infection and that measurements in the blood missed this local expansion. We again observed an increase in Ki-67 + MAITs in the blood over the course of infection; however, we observed a similar increase in activated non-MAIT CD8 + T cells as well (Figure 4j). The activation of non-MAIT CD8 + T cells is consistent with the higher pathogenicity of Mt infection compared with BCG vaccination.\(^{32}\) These results further illustrate that macaque MAITs are activated and respond rapidly to bacterial infection. Future studies examining lung MAITs in Mt infection may reveal more rapid activation compared with blood MAIT responses.

**DISCUSSION**

Here, we thoroughly characterized nonhuman primate MAITs and demonstrated their capacity to respond to bacterial threats. We utilized a novel strategy to synthesize macaque and human MR1 molecules that bind a wide breadth of different ligands. Elution studies from these naturally loaded MR1 molecules revealed a surprising spectrum of previously unidentified ligands, which will serve to further our understanding of MR1 function. Importantly, however, we found that MR1 molecules from different primate species bind identical ligands. This conservation indicates an important role for MAIT cells in the primate immune response, and suggests that studies of MR1 in macaques are directly applicable to human MR1. When the naturally loaded MR1 molecule is used as a tetramer reagent, it likely stains a greater diversity of MAIT cells as indicated by the variety of TCRs we identified. While the TRAV1-2 and TRAJ33 were common among the MAIT cells we sequenced, we also identified additional, non-canonical TCRs (TRAV19) that a single-ligand tetramer might not bind.

In addition to establishing MR1 and MAIT function in macaques, we have annotated the rhesus macaque TCRx genes in order to perform single-cell paired TCRxβ analysis. This technique will now facilitate a more detailed investigation of the TCR repertoire in macaque studies of aging, host defense, and autoimmunity, thereby increasing the utility of this animal model. Indeed, paired with the macaque MR1 tetramer, this technique can now elucidate whether the MAIT TCR repertoire changes in response to vaccination and infection, and whether tissue- or pathogen-specific MAIT TCR signatures exist.

MAITs are an important component of host immunity to Mt infection, and represent a potential new vaccine target. Using the MR1 tetramer made here, future macaque studies can directly examine if MAITs play a role in controlling Mt or other bacterial infections, and test whether their function, phenotype, and anatomical distribution can be modulated through the rational design of novel vaccine immunogens. While we did not observe MAIT cell expansion in the blood of Mt-infected macaques, we hypothesize that expansion occurred at the site of infection in the lung. Unfortunately, we were unable to specifically measure lung immunity in the current study. However, given the ability to perform repeated longitudinal lung biopsies or BAL in macaques, future Mt studies in macaques can definitively investigate pulmonary MAIT immunity. Ultimately, this study demonstrates the power of the nonhuman primate model to interrogate MAIT biology and provides a foundation for studying the role of MAITs in both infectious disease and autoimmune disorders.

**METHODS**

**Live subjects and bacteria.** Seventeen nonhuman primates were followed longitudinally in this study. All animals were cared for at the Oregon National Primate Research Center (ONPRC) with the approval of the ONPRC Animal Care and Use Committee using the standards of the NIH Guide for the Care and Use of Laboratory Animals. Four Indian rhesus macaques were challenged intradermally with \(2 \times 10^5\) live units of BCG vaccine SSI. Half of the dose was delivered to the left and right chest of each animal. Nine rhesus macaques were infected intrabronchially with 10 CFU of Mt Erdmann strain. Details on the clinical progression of the animals can be found in Hansen et al. (unpublished data). Cyropreserved or fresh samples from an additional 43 nonhuman primates were used in this study, which were assigned to other, unrelated studies. Cyropreserved samples were used in the cross species stain, TCR sequencing experiments, and functional assay. We observed no loss of function by cyropreserved splenocytes compared with fresh PBMCs (data not shown).

**Expression of soluble macaque MR1 protein, ligand loading, and biotinylation for tetramer preparation.** *Macaca mulatta* MR1 and \(\beta_m\) genes were expressed as a single-chain construct in insect cells using the baculovirus expression system (see Supplementary Figure S2a,b). The \(\beta_m\) gene was fused with the ectodomain of the MR1 gene with a flexible glycine serine linker. The BirA biotinylation sequence (avitag), 3C protease digestion sequence and 6X histidine tag were added at the C-terminal end of MR1 and cloned into the pACgp67A vector (BD Biosciences, San Jose, CA) downstream of the gp67 secretion signal sequence. A second construct was made only with a 6X histidine tag at the C-terminal end (see Supplementary Figure S2a).

Baculovirus containing MR1 constructs were generated following the transfection and amplification in Sf9 cells. MR1 was loaded with *E. coli* ligands following a co-infection method during MR1 expression in Hi5 cells. Specifically, Hi5 cells were grown in antibiotic-free media and infected with MR1 containing baculovirus. After 18–20 h of incubation, Hi5 cells were co-infected with *E. coli* strain BL21 (40 cells per ml) from an overnight culture. Hi5 cells were harvested after 65–68 h and supernatant containing the secreted MR1 was collected by centrifugation. The pH of the supernatant was adjusted to 7.2 with HEPES or Tris (Fisher Scientific, Waltham, MA) buffer and buffer...
exchanged with HEPES buffered saline (HBS: 10 mM HEPES, pH 7.2, 150 mM NaCl, and 0.02% azide). MR1 proteins were purified from supernatant using Ni-NTA affinity chromatography (Qiagen, Valencia, CA). Ni-NTA agarose and imidazole (final concentration of 20 mM) were added to the supernatant and incubated overnight on a rotator. The next day, protein was eluted with HBS and 200 mM imidazole, and fractions were checked on sodium dodecyl sulfate polyacrylamide gel electrophoresis. MR1 containing fractions were pooled, buffer exchanged to HBS with 20 mM imidazole and treated with 3C protease overnight at 4°C to remove the histidine tag. Protein was again passed through the Ni-NTA column to remove contaminants and flow through was collected. MR1 was buffer exchanged three times with 20 mM Tris pH 8.0, 50 mM NaCl and concentrated.

MR1 was biotinylated using the BirA enzyme in the presence of buffer-A (50 mM bicine buffer, pH 8.3) and buffer-B (10 mM ATP, 10 mM MgOAc, and 50 μM d-biotin) (Avidity, LLC, Aurora, CO). After 5 h of incubation at room temperature MR1 was purified using S200 size exclusion chromatography (GE Healthcare, Fairfield, CT) and checked for biotinylation in non-reducing sodium dodecyl sulfate polyacrylamide gel electrophoresis with streptavidin (see Supplementary Figure S2c).

For ligand identification experiments, MR1 histidine-tagged protein was loaded similarly with E. coli ligands, purified with Ni-NTA and S200 size exclusion chromatography.

Sequencing and sequence analysis. DNA and RNA were isolated from approximately 3 × 10^6 PBMCs from all primate species using the Qiagen DNA/RNA Allprep Kit. cDNA was then synthesized using the SuperScript III Reverse Transcriptase (ThermoFisher Scientific, Waltham, NJ). Amplimers of MR1 sequences were generated via amplification of cDNA by PCR using high-fidelity Phusion polymerase (New England Biolabs, Ipswich, MA) and the primers below for amplification of cDNA by PCR using high-fidelity Phusion polymerase (ThermoFisher Scientific, Waltham, MA). cDNA was then synthesized using the SuperScript III Reverse Transcriptase (ThermoFisher Scientific, Waltham, MA). cDNA was then synthesized using the Transcriptor High-Fidelity cDNA Synthesis Kit (Roche, Basel, Switzerland). After 1 h, colon pieces were pressed over a 70-μm cell strainer and rinsed thoroughly with HBSS. Colon pieces were then shone for 1 h at 225 r.p.m. in R3 containing 0.5 μM EDTA. Colon pieces were then poured over a tea strainer and rinsed thoroughly with HBSS. Colon pieces were then shone for 1 h at 225 r.p.m. in R3 containing 0.5 μM EDTA. Colon pieces were then poured over a tea strainer and rinsed thoroughly with HBSS. Colon pieces were then shone for 1 h at 225 r.p.m. in R3 containing 0.5 μM EDTA. Colon pieces were then poured over a tea strainer and rinsed thoroughly with HBSS. About 500 μl of 0.5 μM EDTA was added to stop collagenase activity. Colon cells were pelleted and then resuspended in approximately 40 ml of isotonic percoll (GE Biosciences) diluted to 70% with R3. This cell suspension was then overlaid below 7 ml of isotonic percoll diluted to 57% with 1 × PBS in eight tubes and spun at 550 g for 20 min with no brake and acceleration minimized. After spinning, cells at the interface were removed using a transfer pipette and placed in R10. Cells were centrifuged at 830 g for 4 min and then resuspended in an appropriate volume for counting. Liver was treated in EDTA, digested, and underlaid in percoll as described above. Lung and skin were diced, digested, and centrifuged in percoll as described above.

Antibodies and staining. Approximately 1–2 × 10^6 cells were placed in 50–100 μl of R10 media for tetramer staining purposes, with or without 50 nM dasatinib. Tetramer was added at a final concentration of 100 nM and cells were incubated in the dark at room temperature for 1 h. Surface staining antibodies were then added and cells were incubated for an additional 30 min in the dark at room temperature. Cells were then washed once with 1 × PBS and fixed with 2% PFA for surface stains or the fix/perm reagent from the eBioscience Foxp3/Transcription Factor Staining Kit. Antibodies used in this study included: anti-CD3 (clone: SP34-2, Pacific Blue; BD Biosciences), anti-CD28 (clone: CD3-1, unconjugated, Mabtech, Nacka Strand, Sweden), anti-CD8 (clone: SK1, TrueRed; BD Biosciences), anti-CD4 (clone: L200, PE-Cy7; BD Biosciences), anti-CD28 (clone: CD28.2, PE; BD Biosciences), anti-CD95 (clone: DX2, FITC; BD Biosciences), anti-CD69 (clone: FN50, ECD, Biolegend, San Diego, CA), anti-CCR7 (clone: 150503, Pacific Blue, R&D Systems), anti-IFNγ (clone: B27, FITC; BD Biosciences), anti-TNF (clone: Mab11, Alexa 700; BD Biosciences), anti-IL-17A (clone: eBio64DEC17; eBioscience, San Diego, CA), anti-PLZF (clone: Mag2.1F7, PE; eBioscience), anti-RORγt (clone: AKFJS-9, PE; eBioscience), anti-T-bet (clone: 4B10, PE-Cy7; eBioscience) V97.2 (clone: 3C10, APC-Cy7; Biolegend), CD161 (clone: 3G10, FITC; Biolegend). LIVE/DEAD Fixable Yellow Dead Cell Stain (Life Technologies, Carlsbad, CA) was used to assess cell viability. Intracellular stains used the Foxp3/Transcription Factor Staining Kit. Stains were run on the LSR II (BD Biosciences) and then analyzed using FlowJo, Version 10.0.8 or 9.9 (TreeStar, Ashland, OR).

Design of TCR segment-specific primers. To design a nested multiplex PCR approach to analyze the paired TCRββ repertoire at the single-cell level in macaque, we adapted a similar strategy to that described for mouse and human single-cell PCR for TCR repertoire analysis. Unlike mouse and human, an annotated TCRα/d locus was not available for macaque in any database. It has been reported that the humans and chimpanzees share 98.6% identity of their genomes and a previous comparative analysis of the TCRβ chain of rhesus macaques and humans reported that their TCRβ chains are highly similar (92.9%) at the nucleotide level. With this knowledge we first assessed the cross-priming of published individual human TRAV and TRBV forward primers with their corresponding reverse primers (TRAC and TRBC reverse primers) using the total RNA harvested from macaque CD8 + T cells. As shown in Supplementary Figure S8, the majority (96.55%) of the macaque TRBV families were amplified by the human external and internal set of forward primers. However, only 35% of the macaque TRAV families showed amplification by the published human TRAV external and internal primers (data not shown). Therefore, we downloaded the macaque ARTICLES
The PCR conditions were 95°C for 5 min, followed by 34 cycles of 95°C for 20 s, 52°C for 20 s, and 72°C for 45 s, followed by a final extension of 72°C for 7 min. At the end of the PCR cycle, the PCR products were visualized on 2% agarose gel (see Supplementary Figure S10). The PCR products were purified by Exonuclease I/Shrimp Alkaline Phosphatase treatment and sequenced as described with the relevant TRAC or TRBC internal primer using a ABI Big Dye sequencer (Applied Biosystem) at the Hartwell Center, St Jude Children’s Research Hospital. The sequence data were analyzed using a custom-built macro-enabled Microsoft Excel sheet and IMGT to derive CDR3 nucleotide and amino acid sequences. To determine the corresponding TRAV-TRAJ or TRBV/TRBJ usage, the sequence data derived from single cells were compared with the extracted macaque-specific TRAV/TRAJ and the downloaded TRBV/TRBJ genes (from IMGT) by using the BLAST algorithm locally (CLCbio; Qiagen).

**Single-cell sorting.** PBMCs and the lymphocytes from the liver biopsy were resuspended at 2 x 10⁷ per ml of PBS. 3 x 10⁵ cells were stained with Fixable Live/Dead Aqua (Molecular Probes, Eugene, OR) in 200 μl volume for 20 min at room temperature in the dark. The cells were centrifuged and the cell pellet was resuspended in 150 μl of sort buffer (PBS containing 0.1% BSA, Fraction V (Gibco, Hampton, NH)) containing 100 ng MR1-APC tetramer. The cells were incubated with the tetrramer for 1Gibco hr at room temperature in the dark. Following incubation, an antibody mixture was mixed consisting of CD3-PacBlue and CD8-PerPCy5.5 in sort buffer was added to the tubes and pulsed by vortexing 2-3 times. The cells were further incubated on ice for 30 min in dark. After the incubation, the cells were washed twice with sort buffer by centrifuging at 500 g for 5 min. The cell pellet was resuspended in sort buffer (containing 200 units of RNasin per ml (Promega)) and filtered through 40 μm cell strainer prior to sorting. The MR1-tetramer + CD8 + CD3 + T cells were single cell sorted into the wells of a 96-well PCR plate (Eppendorf) using a 1μl Synergy cell sorter (Sony, Tokyo, Japan) with following settings: Multi-drop sort OFF, Multi-drop exclude OFF, Division 10, Center sort%: 90. The last two columns of the plate (11 and 12) were left unsorted to use as negative controls for the PCR. After sorting, the plates were sealed using plate sealer film (MicroAmp; Applied Biosystems, Foster City, CA) and centrifuged at 500 g for 3 min and stored at −80°C until reverse transcription and PCR.

**TCR RT-PCR and sequencing.** The cDNA synthesis was performed directly from single cells without any RNA extraction step using the SuperScript VILO cDNA synthesis kit (Thermo Scientific) with minor modification of the manufacturer’s instructions. The cDNA synthesis was carried out in 2.5 μl of reaction mix comprising 0.5 μl of 5 X VILO reaction mix, 0.25 μl of 10 X SuperScript Enzyme mix, 0.25 μl of 1% Triton X-100 (Sigma). The reaction was performed by using the following program in a thermocycler: 25°C for 10 min, 42°C for 60 min, and 85°C for 5 min. Following reverse transcription, a multiplex-nested PCR was carried out using a Taq polymerase-based PCR kit (Qiagen). The PCR reaction recipe for first and second round PCR were performed as previously described except for the primer components.24-26,35,36 For the first-round PCR, a cocktail of TRAV and TRBV specific forward primers (each primer at 0.1 μM final concentration) and both TRAC and TRBC specific anti-sense primers (0.2 μM final concentration) was used (see Supplementary Figure S9). For the nested PCR, the reaction for the TCRβ chains and TCRδ chains was setup separately using 2.5 μl of the first-round PCR products to serve as templates. The PCR mix for TCRα and TCRβ consisted of the same recipe as described with corresponding internal sense TCRα primers and internal antisense TCR β primer or internal sense TRBV primers and internal antisense TRBC primer, respectively (see Supplementary Figure S8).24,35 The PCR conditions were 95°C for 5 min, followed by 34 cycles of 95°C for 20 s, 52°C for 20 s, and 72°C for 45 s, followed by a final extension of 72°C for 7 min. At the end of the PCR cycle, the PCR products were visualized on 2% agarose gel (see Supplementary Figure S10). The PCR products were purified by Exonuclease I/Shrimp Alkaline Phosphatase treatment and sequenced as described with the relevant TRAC or TRBC internal primer using a ABI Big Dye sequencer (Applied Biosystem) at the Hartwell Center, St Jude Children’s Research Hospital.24 The sequence data were analyzed using a custom-built macro-enabled Microsoft Excel sheet and IMGT to derive CDR3 nucleotide and amino acid sequences. To determine the corresponding TRAV-TRAJ or TRBV/TRBJ usage, the sequence data derived from single cells were compared with the extracted macaque-specific TRAV/TRAJ and the downloaded TRBV/TRBJ genes (from IMGT) by using the BLAST algorithm locally (CLCbio; Qiagen).

**Extraction of the TRAV, TRDV, and TRJ genes from the rhesus macaque genome.** The published rhesus macaque (Macaca mulatta) genome is available from the National Center for Biotechnology Information (NCBI) Rhesus Macaque Genome Resources website (http://www.ncbi.nlm.nih.gov/projects/assembly/rgp/assembly/macaca_mulatta/).36 The TRA and TRD gene locus are located on chromosome 7 (Accession number: NC_007864.1). The human TR reference genes (i.e., ‘01 alleles) were obtained from the IMGT/GENEDB database (downloaded on 9 December 2010).36 To extract the TRA and TRD genes from the rhesus macaque genome we used a method similar to the one we previously used to extract the TRB genes from the rhesus macaque genome.25 The rhesus macaque chromosome 7 sequence was queried against all human TRA and TRD reference gene sequences using BLAST (Basic Local Alignment Search Tool, version 2.2.20) to identify the rhesus sequence that resembled the human TR genes.37 Results were filtered to those with e-value ≤ 0.001, a match to ≥ 35% of the query human reference gene, and total percent identity ≥ 75% with the human reference gene. Overlapping regions from the BLAST searches against all TR genes were then merged and the resultant regions were extended by 300 base pairs in both the 5’ and 3’ directions to account for regions that may have been missed due to the BLAST local alignment. These regions of the rhesus macaque genome were then compared against each of the human TR reference genes using EMBOS Needle 6.0.0 (gap open: 10, gap extend: 4), which uses the Needleman-Wunsch algorithm.38 These alignments were used to identify the best human match to each region of the rhesus macaque genome, to determine the terminal ends of the rhesus macaque gene sequences, and to determine the various gene regions (i.e. intron, exon, and recombination signal sequences). TRA and TRD sequences available from the Dryad Digital Repository: http://dx.doi.org/10.5061/dryad.vm36c.

**MAIT ICS.** A small seed culture of DH5α E. coli was inoculated and incubated on a shaker at 37°C overnight. The following day, a 1:50-1:100 dilution was prepared and shaken until it reached approximately 5 x 10⁸ cells per ml as estimated by OD600 on a spectrophotometer. The E. coli were then fixed using 2% PFA for 20 min. The E. coli were then washed and resuspended in complete media for addition to the assay at a 15:1 bacteria to cell ratio. PBMC or splenocytes were resuspended at 2 x 10⁷ cells per ml and 50 μl (1 x 10⁶) cells were added to each well of a 96-well round bottom plate in R10. Blocking antibodies were added 1 h prior to the addition of E. coli at a concentration of 100μg ml⁻¹ and cells were incubated at 37°C and 5% CO₂ for 1 h. After the addition of E. coli, cells were incubated for another hour before the addition of Brefeldin A (Sigma Aldrich) and co-stimulatory molecules CD28 and CD49d (BD Biosciences). Cells were incubated for 8 h at 37°C and 5% CO₂, and then refrigerated until the following day. Cells were washed five times to remove blocking antibodies and stained with tetramer and surface staining antibodies. Then intracellular stains were performed using the eBioscience Fix/Perm/Transcription Factor Staining Buffer Set.

**Statistical analysis.** Owing to small sample sizes, we could not assess the normality of our data and utilized nonparametric tests throughout our analysis. For tissue MAIT frequency and functional assay analysis, we used the Kruskal Wallis test to evaluate differences between each column, and pre-planned contrasts test using Dunn’s test was performed to correct the overall type I error rate. We used the Wilcoxon
rank-sum test to compare Ki-67 and CD69 frequencies in select tissues. We used the Friedman test in the longitudinal BCG and TB study to detect differences between time points. When significant, we used Dunn’s multiple comparison post-test to compare each time point to day 0.

SUPPLEMENTARY MATERIAL is linked to the online version of the paper at http://www.nature.com/mi

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AUTHOR CONTRIBUTIONS
J.M.G. contributed to writing of the manuscript, performing experiments, analysis and figures. P.D. and C.M. contributed to performing experiments, analysis, writing, and figures. S.R. contributed to reagent development, writing, and figures. W.A., B.F.R., D.F.M., and M.C.G. contributed to performing experiments and analysis. J.S.R., S.A., H.L.W., B.J.B., J.C.F., and G.X. contributed to performing experiments. K.B.H., contributed to performing experiments, analysis, and figures. H.C., A.W.L., and M.K.A. contributed to coordinating experiments and animal care. T.H.D., H.G.G., M.K., M.J.H., M.C.G., and D.A.L. contributed to analysis and conception of experiments. B.S.P. contributed to statistical analysis of experiments. S.H.G. contributed to coordinating and performing experiments and their analysis. J.J.S. contributed to veterinary care and experimental advice. L.J.P. contributed to conception of the experiments. V.V. contributed to analysis and writing. W.H. contributed to performing experiments and their analysis. P.G.T. contributed to conception of experiments and their analysis. D.M.L. contributed to conception of the experiments and their analysis. E.J.A. contributed to conception of the experiments, reagent development, analysis, and figure production. J.B.S. contributed to conception of the experiments, performing the experiments, analysis, writing, and figure production.

DISCLOSURE
The authors declared no conflict of interest.

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