Regionally compartmentalized resident memory T cells mediate naturally acquired protection against pneumococcal pneumonia

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As children age, they become less susceptible to the diverse microbes causing pneumonia. These microbes are pathobionts that infect the respiratory tract multiple times during childhood, generating immunological memory. To elucidate mechanisms of such naturally acquired immune protection against pneumonia, we modeled a relevant immunological history in mice by infecting their airways with mismatched serotypes of *Streptococcus pneumoniae* (pneumococcus). Previous pneumococcal infections provided protection against a heterotypic, highly virulent pneumococcus, as evidenced by reduced bacterial burdens and long-term sterilizing immunity. This protection was diminished by depletion of CD4+ cells prior to the final infection. The resolution of previous pneumococcal infections seeded the lungs with CD4+ resident memory T (TRM) cells, which responded to heterotypic pneumococcus stimulation by producing multiple effector cytokines, particularly interleukin (IL)-17A. Following lobar pneumonias, IL-17-producing CD4+ TRM cells were confined to the previously infected lobe, rather than dispersed throughout the lower respiratory tract. Importantly, pneumonia protection also was confined to that immunologically experienced lobe. Thus regionally localized memory cells provide superior local tissue protection to that mediated by systemic or central memory immune defenses. We conclude that respiratory bacterial infections elicit CD4+ TRM cells that fill a local niche to optimize heterotypic protection of the affected tissue, preventing pneumonia.

INTRODUCTION

Pneumonia remains a serious public health burden both in the United States and globally. More than 1 million children under the age of 5 years die worldwide from pneumonia and associated complications each year.1 In the United States, pneumonia is the most common reason for the hospitalization of children2 and accounts for nearly half of the infectious disease-related hospitalizations and deaths of older adults.3 Pneumonia disproportionally affects the youngest and older members of the population, with differential underlying immunological explanations.4 The most common cause of community-acquired bacterial pneumonia at both ends of the age spectrum is *Streptococcus pneumoniae* (pneumococcus). Colonization of the upper airways by pneumococcus is prevalent and recurrent for children and a precursor for pneumococcal disease, which in addition to pneumonia can also include meningitis, sepsis, and otitis media.1,5 Widespread vaccination programs with the pneumococcal conjugate vaccine have significantly reduced the incidence of pneumococcal disease; however, this vaccine is by design only capable of protecting against a small subset of pneumococci (so-called “vaccine type”) and some studies report an increase in disease caused by non-vaccine serotypes.5 Challenges with current vaccines highlight the need for a better understanding of...
Protective immune mechanisms in order to develop new vaccines that provide broader protection.

Pneumococcal carriage decreases during the first 2 years of life due in part to the development of naturally acquired adaptive immune memory. To provide protection against respiratory pathogens that exhibit substantial diversity within species, such as the seasonal variation in influenza viruses or the >90 different serotypes of pneumococcus currently circulating, naturally acquired adaptive immune protection must involve heterotypic responses to epitopes widely conserved within a species. Humans have heterotypic memory T cells and serum antibodies that recognize diverse strains of influenza virus as well as multiple serotypes of pneumococcus. Both epidemiological and experimental evidence in mice and in humans demonstrate that this naturally acquired heterotypic immunological memory provides substantial protection against respiratory infection with newly encountered influenza viruses. Very recently, naturally acquired heterotypic immunity against pneumococcus has been modeled in mice, revealing that CD4 T helper type 17 (Th17) cells can help protect the lung against pneumococcal infection. It remains unclear which types of memory T cells may provide such heterotypic immunity against pneumococcus in the lung and how they enhance lung defense.

In addition to systemic immune responses, the mucosal surfaces also contain resident memory T (T RM) cells that can be elicited by viral and chronic infections. The first evidence for T RM cells in the lung came from mouse studies which demonstrated that influenza infections result in lung-localized, non-circulating, influenza-specific memory CD4 T cells that provide superior host defense against subsequent infections compared with the circulating influenza-specific central memory CD4 T cells. Adult human lungs contain large numbers of CD4 TRM cells based on surface staining with CD69, and at least some of these cells respond to influenza, which suggests that they resulted from prior respiratory infection. Whether and how the bacterial causes of pneumonia elicit or are influenced by lung CD4 TRM cells is, to our knowledge, largely unexplored. The types of pathogens recognized by lung CD4 TRM cells, the responses of lung CD4 TRM cells to relevant activation stimuli, and the functional capabilities of lung CD4 TRM cells require further study, with knowledge gaps especially significant for bacterial pneumonia.

RESULTS
Repetitive respiratory infections establish heterotypic protection against pneumococcal pneumonia

In order to advance understanding of immune mechanisms protecting normal healthy adults from pneumococcal pneumonia, we endeavored to model naturally acquired heterotypic lung immunity in mice. We caused mild and self-limiting respiratory infections with pneumococcus, allowed 4–8 weeks for any inflammation to subside, and then infected the lungs of these mice with Sp3, a serotype to which they had not previously been exposed. In naive mice, this Sp3 challenge causes a severe pneumonia that includes inexorable growth of the bacteria in the lungs and disseminated extrapulmonary infection. The initial infections were with live pneumococcus via an intranasal (i.n.) instillation to mimic natural infection, using a volume and delivery designed to distribute throughout the upper and lower airways of the mice. When mice were infected i.n. with one dose of Sp19F 4 weeks prior to Sp3 pneumonia challenge, the bacterial burden in the lungs of the mice did not differ significantly from the bacterial burden in the lungs of the control mice treated i.n. with sterile saline (Figure 1a). However, when mice were infected i.n. with two doses of Sp19F 1 week apart prior to the intratracheal (i.t.) Sp3 challenge, there was a multi-log reduction in lung bacterial burden compared with the saline-treated control group (Figure 1a). During early childhood, multiple different pneumococcal serotypes infect the airways and potentially generate immune responses. We tested whether diversified pneumococcal serotypes were similarly capable of providing heterotypic immune protection against pneumonia. Serial i.n. injections with Sp19F followed by Sp35B or with Sp19F followed by Sp35B followed by Sp23A were each capable of improving pulmonary defense against a heterotypic pneumococcus compared with control mice that received sterile saline previously instead of pneumococcus (Figure 1a). Bacteremia was also assessed, and mice that received serial pneumococcal exposures had significantly less Sp3 in their blood than control mice previously receiving saline (Figure 1b). These data indicate that prior pneumococcal exposures elicit immune memory that can control and contain a highly virulent pneumococcus after respiratory infection.

In order to determine whether this heterotypic protection was durable and life-saving, we followed cohorts of mice for a longer period of 7 days after Sp3 infection. The saline-treated mice began to decline around day 2 of pneumonia, with no mice living by day 5 (Figure 1c). In contrast, 94% of the Het Imm (i.n.) mice survived to the end of the experiment on day 7 (Figure 1c), at which point they were killed so that lung and blood bacterial burdens could be determined. No Sp3 colonies were recovered from the lungs or blood of nearly all (16/17) mice that had been previously exposed to heterotypic pneumococci. These data demonstrate that heterotypic lung protection generated by mixed exposures to pneumococci can result in long-term sterilizing immunity leading to a full recovery from an otherwise lethal infection.

Although our goal was to generate a mouse model of naturally acquired heterotypic anti-pneumococcal pulmonary immunity, there was the potential that respiratory exposures to pneumococcus could bolster non-specific innate immune responses in the lung. In particular, although more transient than the protection observed here (lasting days instead of weeks or months), it has been previously demonstrated that intrapulmonary delivery of non-viable extracts from Haemophilus influenzae bacteria can remodel lung innate immune responses to provide improved host defense that is effective...
against a wide range of microbes, including Gram-positive bacteria, Gram-negative bacteria, viruses, and fungi. Therefore, we tested whether our pneumococcus-elicited defense against lung infection with heterotypic pneumococci could be generated by dead pneumococci and whether the protective immunity elicited by respiratory pneumococcal infections extended to bacterial species other than pneumococcus. We did not see the same heterotypic protection against Sp3 pneumonia when mice were treated with equivalent numbers of Sp19F that were heat-killed instead of living (Figure 1d). These data suggest that elements of active infection are essential to the generation of heterotypic immunity in the respiratory tract. Furthermore, prior infections with live pneumococci did not provide lung protection against the unrelated Klebsiella pneumoniae in the lungs (Figure 1e). These data suggest that pneumococcal infections generate protection against pneumonia that is restricted in microbial specificity. Altogether, these results argue against the respiratory
exposures to pneumococcus remodeling pulmonary innate immunity alone to protect against a wide range of pathogens but suggest instead that repeated respiratory infections with pneumococci generate new mechanisms of lung defense that are heterotypic but pneumococcus specific.

**Heterotypic anti-pneumococcal immunity accelerates lung neutrophil recruitment**

To gain a first glimpse into the mechanisms behind heterotypic lung protection during pneumococcal pneumonia, we began by examining histological views of susceptible and resistant lungs, both with and without Sp3 pneumonia. One mechanism by which viral infections have been demonstrated to remodel adaptive immunity and protect against viral pneumonia is via the generation of inducible bronchus-associated lymphoid tissues, characterized by organized lymphoid aggregates in the subepithelium and airway interstitium. However, hematoxylin and eosin staining did not reveal inducible bronchus-associated lymphoid tissues or other types of tertiary lymphoid organs, or any consistent histological differences between non-pneumonic lungs of mice with heterotypic immunity compared with those of saline control mice (Figure 2a). After 24 h of infection with Sp3, lungs from both groups showed histological evidence of acute pneumonia, including leukocyte recruitment to the air spaces and interstitia (Figure 2a). To quantify the airspace recruitment of leukocytes, bronchoalveolar lavage was performed 7 and 24 h after infection. After 7 h, there were significantly more neutrophils in the airspaces of the protected lungs compared with susceptible controls (Figure 2b). At 24 h, a time at which there were multi-log decreases in bacteria in the lungs of mice with heterotypic protection, there were no longer differences in the numbers of airspace neutrophils (Figure 2b). We observed no differences in airspace macrophage numbers between control and protected lungs at either time point. Thus prior respiratory infections with pneumococcus remodel the lung immune response to accelerate neutrophil recruitment.

**Increased lung-to-neutrophil signaling due to heterotypic immunity**

Because neutrophil recruitment was accelerated, we examined the production of neutrophil-directed cytokines in the lungs. During pneumococcal pneumonia, various neutrophil-directed cytokines have been shown to come from distinct cellular sources. In the airspaces of mice previously exposed to pneumococcus, neutrophil chemokines (C-X-C motif chemokine ligand 1 (CXCL1), CXCL2, and CXCL5) and colony stimulating factors (granulocyte colony-stimulating factor (G-CSF) and granulocyte-macrophage colony-stimulating factor (G-MCSF)), and their receptors, were analyzed in BALF. Neutrophil-directed cytokines were significantly increased in the lungs of mice with heterotypic immunity compared with saline controls (Figure 2b). These findings suggest that heterotypic anti-pneumococcal immunity accelerates lung neutrophil recruitment through increased lung-to-neutrophil signaling.

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**Figure 2** Heterotypic anti-pneumococcal immunity accelerates lung neutrophil recruitment. (a) Representative hematoxylin and eosin images of Het Imm and saline control lungs that were collected, fixed with 4% paraformaldehyde, and paraffin embedded 0 and 24 h after an infection with $1 \times 10^6$ Sp3 colony-forming units (CFU). Images represent $\times 10$ magnification. (b) Bronchoalveolar lavage fluid (BALF) was collected from Het Imm (i.n.) and saline lungs 7 and 24 h following infection with $1 \times 10^6$ Sp3 CFU. Differential cell counts were obtained and data are expressed as the total number of neutrophils and macrophages. Bars represent means with s.e.m. displayed. Data were log transformed ($Y = \log(Y)$) and significance was determined by two-way analysis of variance followed by Sidak’s post hoc test ($n = 8–9$ for each group). $^*P<0.05$ for saline vs. Het Imm comparisons. Two independent experiments were performed in every case.
factor (GM-CSF)) were significantly increased in the lungs of mice with heterotypic immunity within 7 h of the pneumonia challenge (Figure 3). Prior studies indicate the sources of these to be predominantly epithelial cells (CXCL5 and GM-CSF), myeloid cells (CXCL2), or other lung structural cells (CXCL1 and G-CSF). These data suggest that during pneumococcal pneumonia in lungs with heterotypic immunity multiple cell types are stimulated to produce higher levels of neutrophil-targeting cytokines.

**CD4+ Th17 cells in lungs with heterotypic immunity during pneumococcal pneumonia**

The proinflammatory cytokine interleukin (IL)-17, which can be produced by CD4+ T cells (Th17), is a pivotal factor in host defense at mucosal surfaces. A major effector function of IL-17 is the stimulation of epithelial and stromal cells to produce cytokines that induce the emigration of neutrophils into mucosal sites. Owing to the increase in various neutrophil chemotactic factors in the airspaces of mice with heterotypic protection, we examined the Th17 signal in the lungs of pneumonic-naïve and Het Imm mice. We were not able to detect any appreciable IL-17A or IL-23p19, a stabilization factor for Th17 responses, in the bronchoalveolar lavage fluid of either treatment group (data not shown). CD4+ T cells are typically found in the lung parenchymal tissue more than in the airspaces. Therefore, IL-17A protein was measured in whole-lung homogenates. Here IL-17A was detected, and pneumonic lungs from mice with heterotypic immunity contained significantly more IL-17A protein than those from pneumonic-naïve mice (Figure 4a). Interferon

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**Figure 3** Increased lung-to-neutrophil signaling due to heterotypic immunity. (a–f) Het Imm (i.n.) and saline control mice were infected with 1 × 10^6 Sp3 colony-forming units and bronchoalveolar lavage fluid (BALF) was collected 7 h later. Cytokine concentrations were measured in the BALF using a multiplex Luminex assay. Data are expressed as pg ml⁻¹ and bars represent means with s.e.m. Dashed lines indicate the limit of detection for the particular analyte. Significance was determined by Student’s t-test (n = 14 per group), *P < 0.05. Three independent experiments were performed. CXCL1, C-X-C motif chemokine ligand 1; G-CSF, granulocyte colony-stimulating factor; GM-CSF, granulocyte macrophages colony-stimulating factor; TNF, tumor necrosis factor.
IFN-γ can also enhance neutrophil-mediated defenses against pneumococcus in the lungs and, similar to IL-17A, can come from memory T cells. There was no significant difference in IFN-γ content in the pneumonic lungs due to heterotypic immunity (Figure 4a). IL-17A can be generated by multiple cellular types, so we tested whether heterotypic immunity caused pneumonic lungs to contain more Th17 cells specifically. Single-cell suspensions of pneumonic lungs were stimulated with phorbol myristate acetate/Ionomycin plus Golgi blockade to allow for intracellular cytokine staining (ICS). Although the percentage of Th1 (defined as CD4⁺ IFN-γ⁺) cells did not change in pneumonic lungs due to previous pneumococcal exposures, the percentage of Th17 (CD4⁺ IL-17A⁺) cells was substantially greater in pneumonic lungs with heterotypic immunity (Figure 4b). This impacted the overall abundance of these cytokine-secreting T cells, with significant increases in Th17 cells due to prior serial exposures to pneumococci but no change in the total number of Th1 cells during active heterotypic protection (Figure 4c). Thus, during pneumonia, lungs with heterotypic immunity contain more Th17 cells and IL-17A than do lungs of mice that had never previously experienced pneumococcal respiratory infections.

CD4⁺ Th17 cells can provide defense against pneumococcus or other bacteria such as Klebsiella, making them strong candidates for mediating the protection against pneumonia in the present model of naturally acquired immunity.
heterotypic anti-pneumococcal immunity. To test whether CD4⁺ cells made essential contributions to the lung protection afforded by heterotypic anti-pneumomococcal immunity, we depleted CD4⁺ cells after full immunity had been established but immediately prior to the final Sp3 challenge. At 72 and 24 h before the Sp3 pneumonia challenge, mice with heterotypic immunity received administrations of either GK1.5 or immunoglobulin G (IgG) i.n. and intraperitoneally (i.p.) in order to ensure complete depletion of CD4⁺ cells. Mice that received GK1.5 to deplete CD4⁺ cells had significantly more Sp3 colony-forming units (CFU) in their lungs compared with the mice treated with non-targeting IgG (Figure 5a). In addition, the depletion of CD4⁺ cells resulted in significantly less IL-17A and IFN-γ protein in the lungs of the GK1.5-treated mice compared with IgG controls (Figure 5b,c). These data indicate that CD4⁺ cells are critical sources of the increased IL-17A (and also IFN-γ) during pneumonia, and they make essential contributions to the improved lung defense of mice with naturally acquired heterotypic anti-pneumococcal immunity.

To test whether this CD4⁺ T-cell-mediated lung defense required lymph node egress characteristic of recirculating memory T cells, we treated mice with the S1P receptor inhibitor FTY720, beginning 4–6 weeks after the initial infections and 2 weeks prior to the final Sp3 challenge (Figure 5d,e). Mice that received FTY720 to block lymph node egress had significantly more Sp3 CFU in their lungs compared with control vehicle-treated mice (Figure 5d). In addition, the depletion of CD4⁺ cells resulted in significantly less IL-17A and IFN-γ protein in the lungs of the FTY720-treated mice compared with control vehicle-treated mice (Figure 5e). These data indicate that CD4⁺ cells are critical sources of the increased IL-17A (and also IFN-γ) during pneumonia, and they make essential contributions to the improved lung defense of mice with naturally acquired heterotypic anti-pneumococcal immunity.

**Figure 5** CD4⁺ cells are required for optimal heterotypic protection against pneumococcal pneumonia. (a) Het Imm mice (intratracheal (i.t.)) were administered either a CD4-depleting GK1.5 antibody 72 and 24 h before a pneumonia challenge both intranasally (100 μg) and intraperitoneally (500 μg) or a control immunoglobulin G (IgG) antibody at the same concentrations. Mice were infected with 1 × 10⁶ Sp3 colony-forming units (CFU) and whole lungs were collected after 24 h of pneumonia to determine lung bacterial burden. (b, c) Interleukin (IL)-17A and interferon (IFN)-γ protein levels were measured in the whole-lung homogenates from a. Data are expressed as pg per lung. Significance was determined by Student’s t-test. (d) FTY720 or vehicle control was administered to Het Imm (i.t.) mice at a dose of 1 mg kg⁻¹ 2 days prior and the day of a pneumonia challenge with 1 × 10⁶ Sp3 CFU. Bacterial burden was assessed in whole lungs after 24 h. Significance was determined using Mann–Whitney test. (e) IL-17A protein levels were measured in the whole-lung homogenates from panel (d). Significance was determined by Student’s t-test. Individual dots represent a single mouse and horizontal lines represent medians. Dashed line indicates Sp3 CFU infection input. Bars represent means with s.e.m. Two independent experiments were performed. *P < 0.05.
days prior to and the day of the final pneumonia challenge. FTY720 had no effect on lung bacterial burdens or on lung IL-17A protein levels (Figure 5d,e). Therefore, we conclude that lymph node egress and CD4+ T-cell recirculation are not essential for the immune protection in the lungs that is elicited by prior lower respiratory infections.

**Recovery from respiratory infection seeds the lungs with pneumococcus-specific Th17 and Th1 cells**

The nature and activities of pneumococcus-specific memory CD4+ T cells elicited by respiratory infections are unknown. Because healthy normal human lungs contain T_{RM} cells and mice who recover from influenza infection have lungs containing influenza-specific CD4+ T_{RM} cells, we hypothesized that recovery from respiratory infections with pneumococcus may leave behind a population of pneumococcus-specific CD4+ T cells in the lungs, poised to respond to subsequent infections with the production of protective cytokines.\(^{20,22,23,41}\) Therefore, we investigated pneumococcus-specific CD4+ T cells from healthy lungs, in which no pneumonia or other infectious process was underway.

We collected CD4+ T cells from the healthy and histologically normal (Figure 2a) lungs of mice with or without heterotypic immunity and stimulated these cells with pneumococcus-pulsed antigen-presenting cells (APCs) to assess reactivity. To monitor cell-specific expression of cytokines that define the lungs, we used ICS to measure IL-17A and IFN-γ in CD4+ T cells. Non-specific stimulation of T cells using anti-CD3/anti-CD28 beads demonstrated that both types of lungs contained CD4+ T cells that were capable of elaborating these cytokines (Figure 6a, left), consistent with prior analyses of mouse and human lungs.\(^{19,23}\) Neither cytokine was produced by CD4+ T cells from either type of lung when they were cultured with vehicle-pulsed APCs (Figure 6a, middle). Importantly, both IL-17A and IFN-γ were expressed by CD4+ T cells from the lungs of previously infected mice that were cultured with pneumococcus-pulsed APCs (Figure 6a, right). These data demonstrate for the first time that pneumococcus-specific CD4+ T cells are in the lungs of mice with heterotypic immunity, even without an ongoing infection. Of interest, these cells are not polyfunctional with regard to the two cytokines analyzed but instead appear to separate into distinct sets with Th-1 or Th-17-like properties.

We next sought to determine whether other cytokines that are hallmarks of T-cell activation may also be produced by pneumococcus-specific CD4+ T cells, in addition to IL-17A and IFN-γ. To answer this question, cultures of lung CD4+ T cells plus APCs that were pulsed with vehicle or pneumococcus were incubated for 72h, after which supernatants were analyzed for concentrations of Th cell effector cytokines. Blocking antibody targeting major histocompatibility complex (MHC)-II was used to test whether antigen presentation to CD4+ T cells was specifically essential to the cytokine responses. Figure 6b,c show the results from CD4+ T cells cultured with vehicle-pulsed APCs (unstim), pneumococcus-pulsed APCs (killed pneumococcus), pneumococcus-pulsed APCs plus MHC-II blockade, or anti-CD3/anti-CD28 as a positive control. CD4+ cells from mice without heterotypic immunity were functionally capable of making cytokines as evidenced by anti-CD3/anti-CD28 stimulation, but they produced none of these cytokines in response to pneumococcus-pulsed APCs (Figure 6b). In sharp contrast, CD4+ cells from the lungs of mice with heterotypic immunity produced multiple cytokines in response to pneumococcus-pulsed APCs, including IL-17A, IFN-γ, IL-22, IL-2, and tumor necrosis factor (TNF)-α (Figure 6c). The induction of these cytokines was completely ablated by the MHC-II blockade (Figure 6c), confirming that this cytokine expression resulted from antigen-specific MHC-II-dependent activation of CD4+ T cells. However, MHC-II-dependent activation of pneumococcus-specific CD4+ cells from lungs with heterotypic immunity did not result in expression of either IL-4 or IL-10 (Figure 6c), demonstrating that T cells residing in the lungs after respiratory infections with pneumococcus have distinct Th cell phenotypes. Overall, these results demonstrate that respiratory infections seed the lungs with pneumococcus-specific memory CD4+ T cells that include Th17 and Th1 phenotypes.

**Lung CD4+ T_{RM} cells and heterotypic pulmonary protection imprint locally**

In addition to being non-circulating CD4+ T cells that reside in tissues, resident memory CD4+ T_{RM} cells are defined by high cell surface expression of CD11a and CD69 in the unactivated state.\(^{19,20}\) In mice, CD4+ T_{RM} cells also have high expression of the memory marker CD44 and low expression of the lymphoid homing receptor CD62L.\(^{20}\) Although T_{RM} cells disperse throughout the skin following a localized viral infection,\(^{42}\) nothing is known about regional localization of lung T_{RM} cells after the resolution of a lung infection. This is of particular interest for pneumococcal respiratory infections because they typically result in lobar pneumonia. We modeled this in mice by generating pneumococcal infections in a single lung lobe and then analyzing CD4+ T_{RM} cells and integrated antibacterial defense in the same (ipsilateral) lung lobe compared with the previously uninfected contralateral lobes from the other side of the thoracic cavity. The ipsilateral lobe showed clear evidence of lung CD4+ T_{RM} cells with a distinct population of CD3+ CD4+ CD11ahigh CD69+ cells appearing in mice with heterotypic immunity but not in those without (Figure 7a). There were significantly more lung CD4+ T_{RM} cells in the previously infected mice compared with previously uninfected mice (Figure 7b), demonstrating that pulmonary infections with pneumococcus generate lung CD4+ T_{RM} cells. These CD3+ CD4+ CD11ahigh CD69+ cells from the lungs were also CD44high and CD62Llow, whereas CD3+ CD4+ CD11alow CD69- cells displayed lower CD44 expression and higher CD62L expression (Figure 7c), further confirming the lung CD4+ T_{RM} cell phenotype. In striking contrast to the ipsilateral lobe, the contralateral lung lobes did not show accumulation of CD3+ CD4+ CD11ahigh CD69+ cells (Figure 7a,b). This leads to the unexpected conclusion...
Figure 6  Recovery from respiratory infection seeds the lungs with pneumococcus-specific T helper type 17 (Th17) and Th1 cells. The left lobes of Het Imm (intratracheal) and saline control mice were digested with collagenase and CD4\(^+\) cells were stimulated with killed Sp3-pulsed splenocytes. (a) After 12 h of stimulation in the presence of a protein transport inhibitor, cells were harvested and CD3\(^+\) cells were analyzed for the production of interleukin (IL)-17A and interferon (IFN)-\(\gamma\) by intracellular cytokine staining. Flow cytometric plots show the percentage of IL-17A and IFN-\(\gamma\) production from CD3\(^+\) cells in three groups: pan T-cell activator \(\alpha\)-CD3/CD28 (left), vehicle-pulsed antigen-presenting cells (APCs, middle), and killed pneumococcus-pulsed APCs (APCs-pneumococcus, right). Two independent experiments were performed. (b and c) CD4\(^+\) cells from Het Imm and saline lungs were stimulated with killed Sp3-pulsed splenocytes for 72 h. Supernatants from saline (b) and Het Imm (c) stimulations were collected and cytokine concentrations were measured using a multi-plex luminex assay. In addition to killed pneumococcus, three controls are displayed: vehicle-pulsed splenocytes (unstim), pan T-cell activator \(\alpha\)-CD3/CD28, and \(\alpha\)-major histocompatibility complex (MHC)-II-blocking antibody added to killed pneumococcus-stimulated samples. IL-22 protein was measured independently using an enzyme-linked immunosorbent assay. Cytokine concentrations were measured in three separate stimulation experiments and are displayed as pg ml\(^{-1}\). Bars represent means with s.e.m. The limit of detection for each assay is indicated. Data were log transformed (\(Y = \text{Log}(Y)\)) and significance was assessed by one-way analysis of variance followed by Sidak’s post hoc test, *\(P<0.05\) vs. APCs-vehicle (unstim) for each cytokine. Three independent experiments were performed. TNF, tumor necrosis factor.
that lung CD4⁺ T<sub>RM</sub> cells localize to and remain in the previously infected tissue site rather than distributing widely throughout the respiratory mucosa.

To determine whether IL-17A-producing CD4⁺ T cells were also differentially localized to the site of previous pneumococcal infections, cells from the left and right lung lobes of mice with or without heterotypic anti-pneumococcal immunity were analyzed by ICS. A distinct population of IL-17A-producing cells was present only in the ipsilateral left lobes from mice with heterotypic immunity but not in the same lobe from control mice nor in the contralateral lobes from those mice with heterotypic immunity (Figure 7d,e). Cells that expressed IFN-γ were not compartmentalized the same way (Figure 7d,f).

Although previous literature has demonstrated a distribution of memory cells throughout a surface such as the skin following transfer immune activities. Extravascular CD4⁺ T cells from the lungs had distinct abilities to transfer immune activities. Extravascular CD4⁺ T cells were sorted from the lungs and spleens of mice with heterotypic immunity and were injected intravenously (i.v.) into naive recipient mice. A heterotypic strain of pneumococcus was subsequently delivered to the lungs of these naive recipients, and after 24 h, IL-17 mRNA was measured in the infected lung. The transfer of spleen CD4⁺ T cells did not change lung IL-17 expression compared with negative vehicle control, but the transfer of lung CD4⁺ T cells was sufficient to significantly increase IL-17A expression in the pneumonic lung (Figure 7h). These results provide evidence that lung-resident CD4⁺ T cells are superior to spleen-derived CD4⁺ T cells in providing heterotypic pulmonary immunity during pneumonia.

Given our findings that lobar pneumonia led to a lobe-specific accumulation of lung CD4⁺ T<sub>RM</sub> cells, we hypothesized that heterotypic protection against pneumococcus might also be restricted within the respiratory tract. In order to test whether the compartmentalization of T<sub>RM</sub> cells within the lung could impact heterotypic lung protection, we designed a unique animal model in which the initial and final infections could be matched to the same lobe or delivered instead contralaterally (Figure 7i). To determine whether anatomical or other differences between the left and right lobes of the mouse lung might impact lung defense in the absence of prior infections, naive mice were examined. Anatomical differences between the left and right lung lobes did not affect bacterial growth as Sp3 lung burdens were equivalently high in naive mice whether the infection was in the left or right lobes (Figure 7i). Groups of mice were infected twice with Sp19F i.t. in their left lobe selectively, separated by a week, and then after 4–8 weeks, one group was challenged with Sp3 i.t. in the ipsilateral (left) and the other group in the contralateral (right) lobes (Figure 7i). After 24 h of lobar pneumonia, lungs were harvested and bacterial burdens were measured. In stark contrast to the naive mice (Figure 7j), in mice which had recovered from prior heterotypic infections of the left lobe, infections of the left (ipsilateral) lobe led to significant decreases in bacterial burden compared with infections of the right (contralateral) lobes (Figure 7k). The previously infected lobe but not the contralateral lobes were capable of initiating a rapid and robust heterotypic recall response that protected against further infection. We conclude that a localized pneumococcal infection in the lung results in the regional deposition of CD4⁺ T<sub>RM</sub> cells that protect the immediate but not distant respiratory mucosa against a subsequent heterotypic pneumococcal pneumonia.

**DISCUSSION**

Repeated respiratory infections with pneumococci establish heterotypic immune protection against pneumococcal pneumonia, mediated by regionally compartmentalized lung CD4⁺ T<sub>RM</sub> cells. We propose that this mimics the naturally acquired immune protection that is afforded by most healthy young adult humans but which wanes over time due to aging and comorbidities, such as smoking, alcohol abuse, and chronic disease. Incorporating the model of naturally acquired heterotypic immune protection against pneumonia with other mouse models of aging and comorbidity will facilitate the investigation of mechanisms underlying pneumonia susceptibility in older adults, which is a major contributor to unhealthy aging. The present studies suggest that the transition from a susceptible young child to a more protected adolescent or young adult involves the regional establishment of lung CD4⁺ T<sub>RM</sub> cells recognizing respiratory pathogens.

Prior to pneumococcal infection, adult laboratory mice do not have CD4⁺ T<sub>RM</sub> cells in their lungs, as seen both here and in previous studies. This contrasts markedly from adult human lungs, in which CD4⁺ T<sub>RM</sub> cells are consistently abundant. Repeated pneumococcal infections of the respiratory tract seed the lungs with antibacterial T<sub>RM</sub> cells, and similar consequences result from respiratory virus infections. It appears that lower respiratory tract infection, rather than the microbiome (which is present in laboratory mice), is essential to generating lung CD4⁺ T<sub>RM</sub> cells.

The cleanliness of the environments in which laboratory mice are raised and studied is an important limitation to mouse
models. Immune responses in standard laboratory mice may most closely model the immune responses of infants and young children, which are important to understand as early childhood is an especially vulnerable time. However, to use mice for modeling and studying immune responses to pathogens in adult humans, it may be necessary to first create a relevant infection history. In this communication, we utilize models that are relevant to studying the naturally acquired heterotypic immune mechanisms that protect most young healthy adults against pneumococcal infection. Similar adaptations for other common respiratory pathogens should also be established, so that the research community will be better empowered to study the immunoprotective mechanisms most important to the majority of the population.

Consistent with our own work, a recent study demonstrates that recovery from prior pneumococcal infections generates memory CD4+ cells that are essential for cross-serotype host protection. Here we demonstrate for the first time that CD4+ T RM cells have a predominant role in mediating host defense during pneumococcal pneumonia. Studies of both human and mouse lungs have observed influenza-specific T RM cells, but the data presented here are to our knowledge the first evidence of lung T RM cells specific to any of the bacterial causes of pneumonia. Similar to the pneumococcus, most of the microbes that most commonly cause pneumonia are ubiquitous pathogens. We postulate that lung T RM cells may be critical to preventing pneumonias from diverse etiologies.

The localization of T RM cells at the anatomical site of possible reinfection is a fundamental aspect of tissue-resident memory and rapid recall responses, but the distribution of these cells within a complex organ such as the lung is largely unknown at present. One study of localized vaccinia virus skin infection in mice found that a population of CD8+ non-circulating T RM cells resided not only at the initial site of infection but also distributed throughout the entire skin surface. Another study found that memory CD8+ cells preferentially persisted in the epidermis at the site of prior infection in a mouse model of cutaneous herpes simplex virus-1 infection. The differences in T RM distribution between these two studies could be due to the different infection courses by two different viruses. To our knowledge, we are the first to show the regional localization of lung T RM cells, and we have leveraged this to demonstrate a profound regional impact on memory recall responses and the outcome of reinfection. Het Imm mice that were challenged contralateral to the initial infections were unable to control the infection and displayed very little heterotypic protection, while mice challenged in the ipsilateral lobe possessed full heterotypic protection. Systemic mechanisms of protection, such as central memory CD4+ T cells and circulating anti-pneumococcal antibodies, should be equally protective for ipsilateral and contralateral lung lobes. Although the actions of anti-pneumococcal T RM cells likely synergize with systemic arms of immunity, the co-localization of heterotypic lung protection with lung T RM suggests that tissue-resident immune memory is an essential mechanism of protection that results from the resolution of respiratory infection.

The effector functions of T RM cells generated following resolution of pneumococcal infections appear complex and to involve multiple parallel protective pathways. When CD4+ T RM cells from the lungs of mice previously exposed to

Figure 7 Lung CD4+ resident memory T (T RM) cells and heterotypic pulmonary protection imprint locally. The left (L) and right (R) lobes of uninfected Het Imm (intratracheal (i.t.)) and saline control mice were digested separately with collagenase. (a) Representative flow cytometric plots show lung T RM cells that were identified as CD4+CD8−/C211+CD69+CD62L+CD44+. (b) T RM cells were quantified in the left (ipsilateral) and right (contralateral) lobes of Het Imm and saline mice. Data are displayed as % CD11a+CD69+ of CD4+ cells. Bars represent means with s.e.m. Significance was determined by two-way analysis of variance (ANOVA) followed by Tukey’s post hoc test (n=4–5 per group, experiment repeated three times). (c) Representative flow cytometric plot of CD44 and CD62L expression on CD11a+CD69− lung T RM cells (red) and lung CD4+ CD11a+CD69− cells (blue). (d) Single cells from the left (ipsilateral) and right (contralateral) lobes of Het Imm (i,t.) and saline mice were stimulated with phorbol myristate acetate/ ionomycin for 6 h in the presence of a protein transport inhibitor. Intracellular cytokine staining and flow cytometry was performed to monitor cytokine production. Representative flow plots show interleukin (IL)-17A- and interferon (IFN)-γ-producing CD4+ cells from Het Imm and saline lungs. (a and f) IL-17A- and IFN-γ-producing CD4+ cells were quantified in the left (ipsilateral) and right (contralateral) lobes of Het Imm and saline mice. Bars represent means with s.e.m. Significance was determined by two-way ANOVA followed by Tukey’s post hoc test (n=5–6 per group). Three independent experiments were performed. (g) Mice on a mixed background were infected twice with saline, intranasal Sp19F (Het Imm i.n.), or i.t. Sp19F (Het Imm i.t.). After 4 weeks, mice from each group were challenged with 1 × 106 Sp3 colony-forming units (CFU). Lungs were harvested after 24 h and bacterial burden was determined. Individual dots represent a single mouse and horizontal lines represent medians. Dashed line indicates Sp3 CFU infection input. Significance was determined using Mann–Whitney test. Two independent experiments were performed. *P<0.05.
pneumococcus were stimulated in an antigen-specific manner, they produced not only IL-17A but also IFN-\(\gamma\), IL-2, TNF-\(\alpha\), and IL-22. IL-17A has been shown in many circumstances to be protective during both colonization and infection with pneumococcus, through the induction of neutrophil recruitment.\(^{14,38,39}\) In addition to Th cells, there are also innate sources
of IL-17 like gamma-delta ($\gamma\delta$) T cells and group 3 innate lymphoid cells (ILC3). In a Klebsiella vaccination study, $\gamma\delta$ T cells were the main source of lung IL-17 during a primary infection, but respiratory tract vaccination led to CD4$^+$ T cells emerging as the dominant source during Klebsiella pneumonia.\textsuperscript{53} In the context of pneumococcal pneumonia, lung group 3 innate lymphoid cells have been shown to produce IL-17 during a primary infection.\textsuperscript{51} Although we cannot at this time rule out the possibility that there are additional cellular sources of IL-17, our CD4$^+$ cell depletion data suggest that CD4$^+$ T cells are the major source of IL-17 in the lungs of mice with heterotypic immunity.

Many of the other cytokines we identify here as products of pneumococcus-specific lung CD4$^+$ T$_{\text{RM}}$ cells may further contribute to protection. IFN-$\gamma$ stimulation of macrophages induces antimicrobial programs and enhances antigen processing and presentation.\textsuperscript{57} In addition, IFN-$\gamma$ facilitates bacterial clearance during pneumococcal pneumonia through the augmentation of neutrophil extracellular trap formation.\textsuperscript{36} IL-22 stimulates the production of antimicrobial peptides by the airway epithelium during Gram-negative bacterial pneumonia.\textsuperscript{52} TNF-$\alpha$ signaling during pneumococcal pneumonia amplifies nuclear factor (NF)-$\kappa$B activation throughout the infected lung, improving neutrophil recruitment and bacterial clearance.\textsuperscript{54} IL-2 is an important cytokine for T-cell proliferation, homeostasis, and differentiation,\textsuperscript{53} and it may be critical in the setting for helping establish and maintain lung-resident memory cells, as has been observed in the setting of allergic airway disease.\textsuperscript{54} A shared downstream mechanism for many from this group of inflammatory modulators involves the acceleration of neutrophil recruitment and amplification of phagocyte function. This functional redundancy during heterotypic lung protection may prevent bacteria from easily developing resistance against these immune mechanisms.

The goal of the work presented here was to increase our understanding of the naturally acquired immune mechanisms that mediate rapid recall responses in the lung. This led to improved mouse models for studying immunity in the lungs and the demonstration that the regional accumulation of lung CD4$^+$ T$_{\text{RM}}$ cells has a predominant role in preventing the most common cause of bacterial pneumonia. Many exciting and important questions remain. The biology of lung CD4$^+$ T$_{\text{RM}}$ cells is only beginning to be elucidated, and the signals that recruit and retain these cells in the tissue remain to be determined. Tracking the number and function of lung T$_{\text{RM}}$ cells in settings of susceptibility such as aging or comorbidities will provide valuable mechanistic insights into pneumonia risk. Future vaccine strategies that generate lung T$_{\text{RM}}$ cells against potential respiratory pathogens such as pneumococcus may be novel approaches for protecting individuals developing susceptibility due to advancing age or comorbidities. Activating these lung T$_{\text{RM}}$ cells pharmacologically to produce downstream immune effectors that provide antimicrobial defense could be an additional innovative approach to pneumonia therapy. Taking informed steps in these exciting directions will require improved knowledge of antimicrobial lung CD4$^+$ T$_{\text{RM}}$ cells.

**METHODS**

**Mice.** All animal protocols were approved by the Boston University Institutional Animal Care and Use Committee. Experiments were initiated when mice were 6–15 weeks of age. Female C57BL/6j mice were used unless otherwise specified and were purchased from Jackson Laboratories (Bar Harbor, ME). Mice were maintained in specific pathogen-free conditions at the Boston University Medical Center.

**Heterotypic immunity.** Bacteria were grown on blood agar plates (Trypticase Soy Agar (TSA) II) with 5% Sheep Blood, BD Biosciences, San Jose, CA) for 14h at 37 °C with 5% CO$_2$ and suspended in sterile saline prior to infecting mice. For heterotypic immunity, mice were serially infected with 1–3 (as specified) doses of 1–3 $\times$ 10$^6$ CFU Streptococcus pneumoniae serotype 19F (EF3030, Sp19F), serotype 35B (Sp35B), and/or serotype 23A (Sp23A), with 1 week intervals between such serial infections. Such infections were self-limiting. Sp23A and Sp35B were clinical isolates provided by Dr Stephen Pelton (Boston University). Injections were performed by anesthetizing mice with a ketamine (50 mg kg$^{-1}$) and xylazine (5 mg kg$^{-1}$) mixture via i.p. injection and applying a 10 µl bacteria suspension to each naris until the inoculation was inhaled completely. These mice are referred to as HetImm (i.n.); control mice received i.n. instillations of sterile saline. In some experiments, as specified, mice were infected i.t. as previously described\textsuperscript{25} in the left lobes with two doses of Sp19F 1 week apart; control mice received i.t. sterile saline instillations. Infections were with 1–3 $\times$ 10$^6$ CFU per mouse for all serotypes and routes of delivery. For heat-killed Sp19F, bacteria were suspended at a dose of 1–3 $\times$ 10$^6$ CFU per mouse and incubated in a 55 °C water bath for 45 min prior to i.n. instillation; plating confirmed no living Sp19F. Mice were rested 4–8 weeks after the final infection or saline instillation before being analyzed further or enrolled in pneumonia experiments.

**Pneumonia.** Streptococcus pneumoniae serotype 3 (Sp3, ATCC 6303, Manassas, VA) was instilled i.t. at a dose of 1 $\times$ 10$^6$ CFU unless otherwise stated for pneumonia challenges. K. pneumoniae (ATCC 43816) was instilled i.t. at a dose of 5 $\times$ 10$^6$ CFU per mouse. To assess bacterial burden, mice were killed at specified time points by isoflurane overdose, and CFU were enumerated in the lungs that were homogenized (Bullet blender, Next Advance, Averill Park, NY) in water containing a protease inhibitor cocktail (Roche, Schisch-Rotkreuz, Switzerland) as well as in blood samples collected from the inferior vena cava. When indicated, mice received FTY720 i.p. at a dose of 1 mg kg$^{-1}$, 2 days prior to and on the day of final pneumonia challenge.\textsuperscript{56} The efficacy of the FTY720 regimen was confirmed by observation of decreased peripheral blood CD4$^+$ T cells.

**Lung histology.** After heart ligation, the lungs were fixed by inflation with 4% paraformaldehyde at 23 cm H$_2$O pressure. Left lobes were paraffin embedded, cut into 5-µm sections, and stained with hematoxylin/eosin.\textsuperscript{57}

**Bronchoalveolar lavage.** Bronchoalveolar lavage fluid was collected as previously described.\textsuperscript{55} Briefly, lungs were removed and secured to a blunted 20-gauge needle via the trachea and inflated with 1 ml ice cold phosphate-buffered saline, after which the liquid was withdrawn and collected. The process was repeated until the lungs had been lavaged with a total of 10 ml phosphate-buffered saline. The first milliliter was collected in a separate syringe and saved for protein analysis. Cells were counted using a hemocytometer (Hauser Scientific, Horsham, PA) and cell differentials were obtained using a cytocentrifuge (Thermo Scientific, Waltham, MA) followed by staining with Diff-Quik solutions (Dade-Behring, Deerfield, IL).

**Cytokine concentrations.** Cytokine concentrations in bronchoalveolar lavage fluid were assessed using a Bio-Plex magnetic bead assay (Luminex, R&D Systems, Minneapolis, MN) and analyzed using a Bio-Plex 200 workstation (Bio-Rad, Hercules, CA). The cytokine panel included GM-CSF, G-CSF, CXCL1, CXCL2, CXCL5, CCL20,
IL-23p19, IL-17A, and TNF-α. Whole-lung IL-17A and IFN-γ protein concentrations were measured by enzyme-linked immunosorbent assay (R&D Systems) following protein extraction from lung homogenates. Cytokine concentrations in ex vivo CD4⁺ T-cell stimulation supernatants were measured by Luminex (for IL-17A, IFN-γ, IL-2, TNF-α, IL-4, and IL-10; R&D Systems) or enzyme-linked immunosorbent assay (for IL-22; R&D Systems).

Lung cell suspensions. To generate single-cell suspensions, mouse lungs were digested in type 2 collagenase (Worthington Biochemicals, Lakewood, NJ). Lung lobes were isolated and stored in RPMI-1640 media (Life Technologies, Carlsbad, CA) on ice until minced in digestion solution (type 2 collagenase 1 mg ml⁻¹, DNase 1 150 μg ml⁻¹, CaCl₂ 2.5 mM in phosphate-buffered saline), incubated at 37 °C while shaking for 1 h, and passed through a 70-μm cell strainer (Fisher, Grand Island, NY). Erythrocytes were removed using red blood cell lysis buffer (Sigma, St Louis, MO) and single-cell suspensions were counted using a hemocytometer.

CD4⁺ depletion. For CD4⁺ depletion studies Het Imm (i.t.) mice were administered an anti-CD4⁺ monoclonal antibody (clone GK1.5) or a rat IgG2b isotype control antibody (BioXcell, West Lebanon, NH). Antibodies were diluted in sterile saline and delivered at doses of 500 μg i.p. and 100 μg i.n. (under anesthesia) both 72 and 24 h prior to the i.t. Sp3 pneumonia challenge.

Ex vivo phorbol myristate acetate/ionomycin stimulation of lung cells. Cells from the lungs digested using the collagenase single-cell suspension protocol were resuspended in RPMI-1640 media containing 10% fetal bovine serum, penicillin (100 U ml⁻¹)/streptomycin (100 μg ml⁻¹), 1 mM sodium pyruvate, 2 mM L-glutamine, 0.2 mM 2-mercaptoethanol, 2 μg MβC while shaking for 1 h, and passed through a 70-μm strainer cell (Fisher, Grand Island, NY). Erythrocytes were removed using red blood cell lysis buffer (Sigma, St Louis, MO) and single-cell suspensions were counted using a hemocytometer.

CD4⁺ depletion. For CD4⁺ depletion studies Het Imm (i.t.) mice were administered an anti-CD4⁺ monoclonal antibody (clone GK1.5) or a rat IgG2b isotype control antibody (BioXcell, West Lebanon, NH). Antibodies were diluted in sterile saline and delivered at doses of 500 μg i.p. and 100 μg i.n. (under anesthesia) both 72 and 24 h prior to the i.t. Sp3 pneumonia challenge.

Adoptive transfer of Het Imm CD4⁺ lymphocytes. Single-cell suspensions of the lungs from uninfected Het Imm mice (CD45.2) were prepared as described above with the following modification. Prior to killing, 2.5 μg of anti-CD45-APC antibody (clone 30-F11) was administered i.v. to label intravascular cells, as previously described. Mice were killed 3 min after i.v. antibody administration, and the lungs were cleared of blood via the right ventricle using a 24-gauge angiocatheter and perfusing with 10 ml of Hank's Balanced Salt Solution. Splenocytes from uninfected Het Imm mice were also harvested and single-cell suspensions were generated by passage through a 70-μm filter. All cell suspensions were labeled with CD45-PE/Cy7 (clone 30-F11), CD4-PE (clone GK1.5), and a viability dye. A FACSAriaII was used to sort cells that were live CD45⁺ but not labeled by the intravascular marker and CD4⁺. Naïve mice received saline, lung-derived CD4⁺ cells, or spleen-derived CD4⁺ cells from Het Imm mice, transferred via i.v. tail vein injection. After 3 days, recipient mice were challenged i.t. with Sp3 as described above. Twenty-four hours later, left lungs tissues were harvested and RNA was extracted as previously described. Quantitative reverse transcriptase–PCR for IL-17A was performed using a Taqman probe set (Mm00439618_m1) (Life Technologies).

Pneumococcal-specific T-cell responses. CD4⁺ cells were positively selected using the EasySep Positive Selection Kit (StemCell Technologies, Vancouver, BC, Canada) from left lobes of saline and Het Imm (i.t.) mice that were digested according to the collagenase single-cell suspension protocol; cells from three saline mice or Het Imm (i.t.) mice were pooled for each experiment. APCs were collected from the spleens of naïve C57BL/6 mice that were euthanized through a 70-μm cell strainer, washed, and rid of red blood cells. APCs were pulsed with killed-Sp3 (kSp3) by incubating at a ratio of 3:1 (bacteria:cells) at 37 °C with 5% CO₂, for 1 h with gentle agitation every 20 min. The kSp3 preparation was generated using beta-propiolactone as a bactericidal agent, as previously described. Some splenocytes were vehicle-pulsed (no killed Sp3) as unstimulated APC controls. Mitomycin C (Fisher Scientific, Waltham, MA) was added to prevent splenocyte proliferation. After washing three times, pulsed splenocytes were mixed with CD4⁺ cells at a 10:1 ratio of splenocytes:CD4⁺ cells. For positive controls, CD4⁺ cells were stimulated with mouse T-activator CD3/CD28 Dynabeads (Life Technologies) at a 3:1 bead:cell ratio. Purified anti-mouse MHC-II clone M5/114.15.2 and isotype control clone RTK4530 (Biolegend) were added to indicated wells at a concentration of 10 μg ml⁻¹. For cytokine concentrations, cells were incubated for 72 h, after which supernatants were collected and stored at −80 °C until multi-plex protein analyses. For ICS assays, cells were cultured 12 h in the presence of GolgiStop, and splenocytes had a mismatched CD45 allele (CD45.1) that allowed separation from the CD45.2⁺ Het Imm CD4⁺ cells during flow cytometric analyses. After the 12-h incubation, cells were harvested, stained for cell surface markers, and permeabilized for ICS using fluorescent antibodies to stain for CD45⁺ (clone BM8), IFN-γ, IL-17A, IL-23p19, IL-10, TNF-α, and IL-6 (Biolegend). All antibodies were from eBioscience except for the CD4-APC antibody, which was from Biolegend. Cells were initially gated for live (using eFluor506; eBioscience) CD45⁺ CD3e⁺ cells.

Lung-resident memory cells. Left (ipsilateral) and right (contralateral) lobes of uninfected Het Imm (i.t.) and saline mice were digested separately using collagenase as described above. Single-cell suspensions were prepared for flow cytometry and labeled with the following antibodies: CD45-BV510 (clone 30-F11), CD3e-APC/Cy7 (clone 145-2C11), CD3e-APC/Cy7 (clone GK1.5), CD11a-APC (clone M17/4), CD69-PE (clone H1.2F3), CD62L-FTTC (clone MEL-14), and CD44-PEFluor610 (clone IM7) (Biolegend). Lung-resident memory CD4⁺ T cells were defined as living cells (L/D⁻) that were CD45⁺ /CD3e⁺ /CD4⁺ /CD11a⁺ /CD69⁺ /CD62L⁻/CD44⁻. Data were expressed as % CD11a⁺CD69⁺ cells of CD4⁺ T cells.

Statistics. All statistical analyses were performed using GraphPad Prism 6.0 (GraphPad, La Jolla, CA). CFU data are expressed as individual values with medians, while all other data are shown as means ± s.e.m. Two groups were compared using either Student’s t-test or Mann–Whitney U-test for non-parametric data. Multiple groups were compared using a one- or two-way analysis of variance followed by post hoc test for multiple comparisons (specified in figure legends). Multiple non-parametric data sets were compared using Kruskal–Wallis test. Data sets included results pooled from multiple independent experiments. Differences were considered significant if P < 0.05.

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AUTHOR CONTRIBUTIONS
All authors participated in the interpretation of experiments and review of
the manuscript. N.M.S.S. led the performance of all experiments, collected
and analyzed data, performed statistical analyses, and drafted the
manuscript. G.A.W. assisted with experimental design, experimental
execution, and data analysis. F.T.C. contributed to the technical design
and execution of the in vivo animal model. K.L.H. assisted with the design of
the flow cytometry experiments. K.Y. contributed to the experimental
design and interpretation of the results. E.L. provided technical assistance
with all aspects of the project. R.M. aided in the formation of the manuscript
and contributed to data analysis and interpretation. H.D. assisted with
experimental design of stimulation assays and contributed to data analysis
and interpretation. M.R.J and L.J.O. contributed to the experimental
design, data analysis, and interpretation. J.P.M. conceptualized the line
of investigation, supervised experiments, assisted in experimental design
and analysis, and contributed to the drafting of the manuscript.

DISCLOSURE
The authors declared no conflict of interest.

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