Neonatal Fc Receptor Regulation of Lung Immunoglobulin and CD103⁺ Dendritic Cells Confers Transient Susceptibility to Tuberculosis

Alexis Vogelzang, Laura Lozza, Stephen T. Reece, Carolina Perdomo, Ulrike Zedler, Karin Hahnke, Dagmar Oberbeck-Mueller, Anca Dorhoi, Stefan H. E. Kaufmann
Max Planck Institute for Infection Biology, Berlin, Germany

The neonatal Fc receptor (FcRn) extends the systemic half-life of IgG antibodies by chaperoning bound Fc away from lysosomal degradation inside stromal and hematopoietic cells. FcRn also transports IgG across mucosal barriers into the lumen, and yet little is known about how FcRn modulates immunity in the lung during homeostasis or infection. We infected wild-type (WT) and FcRn-deficient (fcgrt⁻/⁻) mice with *Pseudomonas aeruginosa* or *Mycobacterium tuberculosis* to investigate whether recycling and transport of IgG via FcRn influences innate and adaptive immunity in the lung in response to bacterial infection. We found that FcRn expression maintains homeostatic IgG levels in lung and leads to preferential secretion of low-affinity IgG ligands into the lumen. Fcgrt⁻/⁻ animals exhibited no evidence of developmental impairment of innate immunity in the lung and were able to efficiently recruit neutrophils in a model of acute bacterial pneumonia. Although local humoral immunity in lung increased independently of the presence of FcRn during tuberculosis, there was nonetheless a strong impact of FcRn deficiency on local adaptive immunity. We show that the quantity and quality of IgG in airways, as well as the abundance of dendritic cells in the lung, are maintained by FcRn. FcRn ablation transiently enhanced local T cell immunity and neutrophil recruitment during tuberculosis, leading to a lower bacterial burden in lung. This novel understanding of tissue-specific modulation of mucosal IgG isotypes in the lung by FcRn sheds light on the role of mucosal IgG in immune responses in the lung during homeostasis and bacterial disease.

Secretary immunoglobulin A (IgA) and IgM are exported by a dedicated receptor to the mucosa, where they coat pathogens and nurture beneficial microbiota (1). Yet the idea of the immune predominance of IgA in mucosal secretions has been challenged by the identification of the neonatal Fc receptor (FcRn) at mucosal sites in adult mammals (2). FcRn binds to conserved elements of the Fc portion of IgG at low pH, transports IgG across polarized epithelial barriers, and chaperones IgG between endocytic and degradative compartments (3). Selective expression of FcRn in epithelium was also found to enable antigens bound to IgG to be retrieved from the lumen to initiate systemic immunity to *Citrobacter rodentium* (4). FcRn guides monomeric IgG to the endocytic excretion pathway, extending its serum half-life by preventing lysosomal degradation (5). Finally, intracellular trafficking of antibody complexes within antigen-presenting cells via FcRn can influence lymphocyte activation with respect to opsonized particles (6, 7). Despite FcRn being the primary means by which IgG is transported across epithelial barriers, conflicting reports in a variety of infection models have questioned whether FcRn is required for the protective action of mucosal IgG, implying immune roles at the mucosal interface beyond neutralization of pathogenic determinants (8–10).

Inhaled air passes constantly through the lungs, continually introducing particulate and microbial matter into the lower airways, where it interacts with secreted antibodies. Many of the studies cited above have shed light on the role of FcRn-mediated IgG regulation and transport specifically in the gastrointestinal tract. Knowledge of the role of FcRn in the lung remains limited, although expression has been identified in parenchyma and recruited immune cells (11, 12). Indeed, FcRn has been harnessed to provide inhaled Fc-fusion vaccines access to systemic immunity (13, 14). We are interested in how IgG interactions with FcRn may be involved in homeostatic immune development in the lung and also in defense against respiratory infections by two bacterial species that require orchestrated multicomponent immune responses. The first, *Pseudomonas aeruginosa*, is an opportunistic pathogen that rapidly recruits innate immune cells to the lung parenchyma and is sensitive to complement deposition (15). The second, *Mycobacterium tuberculosis*, is a slow-growing pulmonary pathogen that disseminates to peripheral organs. Exponential growth in the lung is arrested by the arrival of specific Th1 T cells in the lung approximately 2 to 3 weeks after infection which promote formation of granulomas composed of diverse myeloid and lymphoid cells (16). Antibodies are thought to play a limited protec- tive role in natural *M. tuberculosis* infection despite strong local recruitment of B cells to granulomas (17, 18). By studying fcgrt⁻/⁻ mice during homeostasis and bacterial infection, we aimed to...
characterize specific immune roles for IgG at the lung interface with the external environment.

MATERIALS AND METHODS

Bacteria. Mycobacterium bovis BCG SSI 1331 (American Type Culture Collection [ATCC]; no. 35733), VPM1002 (recombinant BCG [rBCG]), and M. tuberculosis H37Rv (ATCC; no. 27294) stock was prepared as described previously (19). To generate green fluorescent protein (GFP)-expressing M. tuberculosis, competent H37Rv was transformed with plasmid pGFP-11 by electroporation and selected with kanamycin (20). P. aeruginosa strain PAO1 (PAO1-DSM; German Collection for Microorganisms and Cell Cultures) was grown in Luria broth (LB) at 37°C to the late exponential/early stationary phase and then subcultured for 2 h at 37°C to the mid-exponential phase before use. For CFU enumeration of the two pathogens, serial dilutions were performed in phosphate-buffered saline—0.05% Tween 80 (PBST) and plated onto Middlebrook 7H11 agar or LB agar and incubated at 37°C for 3 to 4 weeks or 16 h, respectively.

Experimental animals and manipulations. FcγR−/− and P25 transgenic mice were purchased from Jackson Laboratories (USA) and bred in-house alongside C57BL/6 controls. Female mice were 8 to 16 weeks of age at the beginning of the experiments and were matched for age and sex and distributed randomly between treatment groups. Mice were kept under specific-pathogen-free (SPF) conditions at the Max Planck Institute for Infection Biology in Berlin, Germany, and experiments were carried out according to institutional guidelines approved by the local ethics committee of the German authorities (Landesamtes für Gesundheit und Soziale). For vaccination studies, P. aeruginosa strain PAO1 was killed by incubation for 2 h in 1% paraformaldehyde at 37°C and then washed 3 times in phosphate-buffered saline (PBS). Killing was confirmed by plating this preparation, and then the solution was adjusted to deliver 1 × 10² bacteria per 6 µl dose into both nostrils at days 0, 8, and 14. Mice were infected intratracheally (i.t.) 40 days after vaccination or sham PBS treatment with 4 × 10³ P. aeruginosa strain PAO1 cells under mild anesthesia. A Glasc-Col inhalation exposure system was used for aerosol infection of mice with low-dose (100 CFU) M. tuberculosis, confirmed by plating lung homogenates 1 day after infection. For generation of bone marrow chimeras, mice were irradiated with 9 Gy and immune reconstituted with 1 × 10⁶ bone marrow cells the following day. Chimeric mice were offered drinking water containing 125 mg/liter enrofloxacin (Bayer) for 4 weeks and allowed to recover for 12 weeks.

Sample preparation. Bronchoalveolar lavage was performed with 500 µl PBS supplemented with protein inhibitors (Roche) for antibody analysis. For antibody and CFU analysis, lungs were homogenized in PBST and protein inhibitors using GentleMACS dissociator M tubes (Miltenyi Biotec). For flow cytometry, lungs were harvested from perfused animals and digested with collagenase (0.7 mg/ml collagenase IV [Sigma-Aldrich]—0.3 mg/ml collagenase D [Roche])—RPML 1640 medium at 37°C in 5% CO₂ for 30 min. Single-cell suspensions were prepared by mechanical dissociation through a 70-µm-pore-size nylon mesh using RPMI 1640 medium with 10% fetal calf serum (Gibco).

Flow cytometry. Single-cell suspensions were stained on ice with the following eBioscience antibodies: B220 (RA3-6B2), Siglec-F lyzed with FlowJo (TreeStar). or FACSCanto II (Becton Dickinson) flow cytometers. Data were analyzed, as previously described (3), FcγR−/− animals displayed reduced serum IgG levels, reflecting the well-described role of the FcRn extending the half-life of systemic IgG. This reduction in IgG isotypes was likewise observed in FcγR−/− lung parenchyma, implying that systemic IgG supplies tissues via diffusion from the circulation. Our analysis of antibody in BALF of WT and FcγR−/− mice revealed that, unlike IgM and IgA, luminal IgG strongly depended on FcRn expression, as FcγR−/− mice had 10- to 100-fold lower concentrations of locally dominant IgG isotypes. Intriguingly, the ratio of isotypes found in the airways of WT mice diverged from that observed in serum, such that IgG2b and IgG3 were the dominant antibody isotypes secreted into the mucosa. This discordant Ig isotype ratio depended on FcRn expression, as the levels of IgG isotypes in FcγR−/− lung more closely mirrored serum and tissue levels (Fig. 1A). Thus, FcRn in the lung qualitatively and quantitatively influences IgG in mucosal secretions.

FcRn modifies homeostatic myeloid immune subsets in lung. We performed immune phenotyping of lung-resident immune cells in WT and FcγR−/− animals to determine whether this receptor influences local leukocyte homeostasis. While most immune cell populations were comparable between strains, in the absence of FcRn we observed increased numbers of lung-resident XC chemokine receptor 1-positive (XCR1⁺) CD103⁺ dendritic cells (DC) compared to the results seen with WT animals (Fig. 1B and C). This correlated with increased homeostatic mRNA transcription of irga (CD103) in lung, the DC growth factor flt3 and CD103⁺ DC-specific transcription factors batf3 and irf8 (see Fig. S1 in the supplemental material). However, expression of CD11b⁺ DC-associated transcription factor irf4 was not influen...
FCRn is expressed by myeloid cells and modulates lung DC populations, and thus may influence local immunity to infection. In order to assess the role of luminal and intracellular IgG transport during acute bacterial infection requiring efficient recruitment of innate leukocytes, we first vaccinated WT and FCRn-deficient (fgrt-/−) mice intranasally with formalin-killed P. aeruginosa to generate a pool of specific IgG in the lung as previously described (21). Vaccinated and sham-treated animals were challenged via intratracheal administration with virulent P. aeruginosa to determine whether local immunity was compromised in the absence of mucosal IgG and in the presence of lower levels of homeostatic IgG circulation in fgrt-/− animals. WT and fgrt-/− mutants were equally susceptible to acute bacterial pneumonia, suggesting that local IgG binding to FCRn is dispensable for the recruitment and antimicrobial action of neutrophils in a model where acute vascular leakage drives a rapid inflammatory response that is independent of adaptive immunity (Fig. 2) (22). Intriguingly, vaccinated fgrt-/− mice lacking mucosal IgG transport were protected from bacterial growth to the same extent as WT, suggesting that protective IgG acts within tissues or reaches the lumen due to paracellular antibody diffusion in P. aeruginosa infection (Fig. 2B).

**FIG 2** FCRn is not required for recruitment of neutrophils or bactericidal activity in P. aeruginosa infection. WT and fgrt-/− mice were vaccinated intranasally 3 times with paraformaldehyde-fixed P. aeruginosa. At 40 days post-vaccination, vaccinated mice (i.n vac) and sham controls (PBS) were intratracheally infected with 4 × 10^6 P. aeruginosa cells. (A) At 24 h postinfection, the percentage of lung-infiltrating neutrophils of live cells was measured by flow cytometry. (B) CFU of P. aeruginosa recovered from lungs 24 h postinfection. Data show medians and interquartile ranges of representative data from 2 experiments (n = 5 to 10 per group in each experiment). Asterisks indicate P values (*, P < 0.05; **, P < 0.01).

FCRN maintains intracellular stores of IgG in myeloid cells. We determined the role of FCRn in a model of chronic bacterial infection where specific IgG levels and adaptive immunity develop gradually over the course of infection. WT and fgrt-/− mice were vaccinated intranasally 3 times with paraformaldehyde-fixed P. aeruginosa. At 40 days post-vaccination, vaccinated mice (i.n vac) and sham controls (PBS) were intratracheally infected with 4 × 10^6 P. aeruginosa cells. At 24 h postinfection, the percentage of lung-infiltrating neutrophils of live cells was measured by flow cytometry. (B) CFU of P. aeruginosa recovered from lungs 24 h postinfection. Data show medians and interquartile ranges of representative data from 2 experiments (n = 5 to 10 per group in each experiment). Asterisks indicate P values (*, P < 0.05; **, P < 0.01).
subjected to aerosol infection with *M. tuberculosis*, and the local and systemic antibody response was measured in serum, homogenized lung tissue, and BALF 28 days after infection. Although *fcgrt*−/− mice exhibited reduced systemic and lung parenchymal IgG levels compared to WT mice, we observed similar 10- to 100-fold increases in the levels of IgG isotypes compared to their respective baseline levels, suggesting that a B cell response of equivalent magnitude had developed during tuberculosis (TB) (Fig. 3A). Indeed, local numbers of B cells in lung tissue increased equally in the two strains over the course of infection (Fig. 3B). IgG1 and IgG2b levels in BALF from infected *fcgrt*−/− mice approached WT levels, indicating that the local antimycobacterial humoral immune response compensated for the reduced half-life of IgG in the absence of FcRn-mediated rescue from lysosomal degradation. However, levels of IgG2c and IgG3 in BALF from WT mice increased 10-fold over the baseline levels in an FcRn-dependent manner, suggesting that these isotypes require active export to the lung lumen across a functional epithelial barrier during TB (Fig. 3A). Immunofluorescence staining of infected WT lung tissue revealed intracellular IgG2c (green) in addition to nuclear DAPI staining (blue). Scale bars, 100 μm (left and middle panels) and 30 μm (far right panel). (D) Median fluorescence intensity (MFI) of intracellular IgG2c measured by flow cytometry in various lung-resident and infiltrating immune cells on day 28 postinfection. Data are representative of the results from 2 experiments (n = 4 to 6 animals per group in each experiment).

**FcRn modifies cell recruitment to granulomas during TB.** In order to investigate whether global FcRn expression affects the kinetics of leukocytes infiltrating the lung during TB, we performed flow cytometric analysis of *M. tuberculosis*-infected WT and *fcgrt*−/− lung tissue. At the early day 14 time point, migratory innate immune cells such as neutrophils, monocytes, and macrophages as well as resident alveolar macrophages were numerically similar in the presence or absence of FcRn. In contrast, discrepancies in the numbers and proportions of neutrophils, inflammatory monocytes, and macrophages developed between *fcgrt*−/− and control animals 1 month postinfection, implying that the influx or survival of inflammatory myeloid cells declined over the course of infection (Fig. 4A). Professional antigen-presenting cells, particularly the enlarged CD103+ DC population, were also more frequent in *fcgrt*−/− lung during infection (Fig. 4A). Infection with *M. tuberculosis* GFP revealed higher proportions and numbers of infected CD103+ DC, suggesting that opsonized *M. tuberculosis* interacts with FcRn to inhibit phagocytic or bactericidal capacity in this population (Fig. 4B; see also Fig. S3 in the supplemental material).

We measured upregulation of CD69, an adhesion molecule that is upregulated following T cell receptor stimulation particularly in the enlarged CD103+ DC population, were also more frequent in *fcgrt*−/− lung during infection (Fig. 4A). Infection with *M. tuberculosis* GFP revealed higher proportions and numbers of infected CD103+ DC, suggesting that opsonized *M. tuberculosis* interacts with FcRn to inhibit phagocytic or bactericidal capacity in this population (Fig. 4B; see also Fig. S3 in the supplemental material). We measured upregulation of CD69, an adhesion molecule that is upregulated following T cell receptor stimulation particularly in the enlarged CD103+ DC population, were also more frequent in *fcgrt*−/− lung during infection (Fig. 4A). Infection with *M. tuberculosis* GFP revealed higher proportions and numbers of infected CD103+ DC, suggesting that opsonized *M. tuberculosis* interacts with FcRn to inhibit phagocytic or bactericidal capacity in this population (Fig. 4B; see also Fig. S3 in the supplemental material).
mice at day 14 postinfection (Fig. 4C). To verify the increased T cell immunogenicity of fcgrt−/− lungs, purified transgenic T cell receptor CD4+ T cells specific for the mycobacterial antigen 85B (Ag85B) were transferred into WT and fcgrt−/− recipients 12 days following infection with M. tuberculosis. At 72 h after the transfer, the fcgrt−/− lung environment drove increased proliferation of transferred cells as measured by carboxyfluorescein succinimidyl ester (CFSE) dilution (Fig. 4D).

Gene expression analysis of selected gene transcripts over the course of infection showed that ongoing inflammation gradually inhibited fcgrt expression in WT lung tissues. Numerically increased CD103+ DC and the corresponding influx of activated CD4+ T cells in fcgrt−/− mice correlated with increased mRNA levels of lymphocyte chemoattractant transcripts such as itgae, encoding the protein CD103, and the lymphocyte- and DC-recruiting chemokines ccl19 and ccl21 (see Fig. S1 in the supplemental material).

FcRn deficiency leads to a transient decrease in M. tuberculosis load and pathology in lung. The idea of the impact of FcRn on pulmonary immunity was supported by analysis of M. tuberculosis growth over the course of disease, which revealed that fcgrt−/− animals were significantly protected from bacterial burden at 2 weeks and to a greater extent at 1 month after infection (Fig. 5A). Perhaps due to the ongoing suppression of fcgrt transcription during inflammation (see Fig. S1 in the supplemental material), this transient effect was lost 2 months postinfection (Fig. 5A). Peripheral organs of fcgrt−/− animals were not similarly protected, suggesting that FcRn does not play a role in dissemination of M. tuberculosis (Fig. 5B). FcRn was also redundant for M. tuberculosis uptake by alveolar macrophages in the bronchoalveolar space following infection, as similar numbers of live M. tuberculosis could be retrieved from lavaged lung parenchyma from WT and fcgrt−/− animals 1 day after aerosol administration (Fig. 5C).

Histological analysis of lung tissue showed equivalent levels of dispersed cellular infiltration in fcgrt−/− mice, despite the reduced M. tuberculosis burden in fcgrt−/− animals (Fig. 5D). In addition to the reduced numbers of infiltrating neutrophils observed in fcgrt−/− animals, the immunostaining pattern of myeloperoxidase was diminished within the lesions of fcgrt−/− lungs, suggesting that early control of mycobacterial growth was associated with limited neutrophilic recruitment to granulomas (Fig. 5E). Thus, FcRn shapes the efficacy of the unfolding adaptive immune response to M. tuberculosis infection in the pulmonary mucosa.

FcRn expressed by hematopoietic and stromal cells inhibits immunity in TB. Last, we generated bone marrow chimeras to investigate whether IgG transport via FcRn to and from airways, or intracellular IgG trafficking via the FcRn within immune cells, correlated with susceptibility to tuberculosis. Both immune and nonimmune cells contributed to maintenance of inflammatory IgG levels in mixed bone marrow chimeras (Fig. 6A). However, the protective phenotype of fewer neutrophils and decreased M. tuberculosis burden of fcgrt−/− mice at 1 month postinfection was recapitulated only when both hematopoietic and nonhematopoietic cells lacked FcRn (Fig. 6B and C). FcRn modulated immunity in both tissues and within mucosal secretions over the course of M. tuberculosis infection. Our results suggest that both functions of the FcRn, ongoing IgG transport across tissue barriers and within intracellular compartments, enhance pathology and inhibit bacterial killing during tuberculosis.

FIG 4 FcRn regulation of IgG during tuberculosis favors infiltration of myeloid cells and restricts T cell priming. Leukocyte infiltration in WT and fcgrt−/− lung was analyzed by flow cytometry after aerosol infection with M. tuberculosis. (A) Quantification of the indicated myeloid cell subsets post-infection. (B) Quantification of the number of innate immune cells in the lung. (C) Quantification of the indicated myeloid cell subsets post-infection. (D) A total of 1 × 10^6 transgenic TCR P25 CD4 T cells stained with carboxyfluorescein succinimidyl ester (CFSE) were intravenously transferred into M. tuberculosis-infected WT and fcgrt−/− mice on day 12 following infection. On day 15 postinfection, the proportion of transferred T cells that underwent cell division was measured as shown on the representative flow cytometry histograms. All graphs show the median and interquartile range and cumulative individual data from 2 pooled experiments where n = 5 animals per experiment, except panel D, which shows results from one experiment of two performed. Asterisks indicate the P value (*, P < 0.05).
This report reveals the nuanced balance between IgG recycling and transport by FcRn under homeostatic and inflammatory conditions in the lung. Analysis of the relative levels of FcRn-dependent transport of Ig subclasses in murine gut led to the proposal of the affinity hierarchy IgG2a/IgG2c > IgG1 > IgG2b > IgG3, and the relative concentrations of IgG isotypes matched this descending order of FcRn binding (23). However, we observed increased levels of IgG2b and IgG3 compared to IgG2c in the lung, though these isotypes were present at similar concentrations in the serum and lung parenchyma. Thus, the IgG isotypes with the lowest affinity for FcRn are found at the highest concentration in lung secretions, leading counterintuitively to a divergence from serum isotype ratios in an inverse relationship to affinity for the receptor.

A similar pattern was described in mammary secretions and healthy human lungs (24–26), and it has been suggested that high-affinity IgG1 and IgG2c antibodies locally saturate FcRn, resulting in preferential release of low-affinity IgG3 and IgG2b into the lumen. The prevalent IgG2b and IgG3 isotypes have relatively increased bactericidal and opsonizing properties (27), and perhaps their dominance represents an adaptation maintained by FcRn in order to maximize the sterilizing efficacy of mucosal antibody in the lung. However, the same processes could damage sensitive tissues during the persistent inflammation that characterizes tuberculosis. Parallels may be drawn between the detrimental actions of FcRn in this model and previous findings showing that it can promote IgG complex-mediated tissue pathology in autoimmune inflammatory disorders (28–30). Indeed, specific antagonists of FcRn are being pursued as novel therapeutic measures for this reason (31).

Our data from murine P. aeruginosa infection support the notion that FcRn-mediated IgG transport into the lumen becomes redundant under inflammatory conditions because paracellular transport and vascular leakage bypass the receptor (8, 10).
compromised immunity to P. aeruginosa infection in fcgrt−/− lung and equivalent uptake of M. tuberculosis GFP by neutrophils suggest a limited cell-intrinsic role for FcRn in neutrophil function in vivo during acute infection. This contrasts with in vitro studies showing that FcRn is required for optimal granulocyte phagocytosis of IgG-coated, heat-killed Streptococcus pneumoniae (32), but perhaps other phagocytic receptors or pathogen-specific determinants can compensate for FcRn-Ig interactions in vivo. The slight difference in M. tuberculosis burdens in lung at day 14, before the arrival of CD4+ T cells and production of specific IgG, is therefore surprising and leaves room for the possibility that FcRn can influence intracellular bacterial replication or local production of bactericidal peptides.

Numerous studies have shown that the early recruitment of CD4+ T cells to the lung is the dominant factor determining bacterial control and lung pathology during tuberculosis (16, 33). Our studies join a growing body of literature showing that failed T cell responses result in excessive inflammation during tuberculosis in humans, primates, and mice, in association with increased bacterial load (34, 35). CD103+ DC have been shown to promote increased Th1 and Th17 CD4+ T cell responses in vivo compared to monocyte-derived CD11b+ DC, T helper subsets that have been shown to be particularly beneficial during TB (36, 37).

It is tempting to speculate that the numerical increase of both infected and bystander CD103+ DC in fcgrt−/− mice maintains improved T cell priming in tuberculosis. Perplexingly, FcRn has previously been reported to aid uptake and presentation of IgG-complexed antigen for CD4+ and CD8+ T cells by CD11b+ DC in several models (38, 39). Yet, during tuberculosis, we observed the inverse phenomenon of increased infection rates and T cell activation in the absence of the receptor at the early day 14 time point after infection, and, even when bacterial growth rates diverged between WT and fcgrt−/− mice by day 28, similar levels of CD4+ T cell activation could be observed despite the lower antigenic burden in protected fcgrt−/− animals. Perhaps DC in various tissue compartments have different requirements for the receptor, or perhaps the solubility of Ig antigen complexes and their interactions with FcRn differ between model systems. The striking increase in resident lung DC observed in fcgrt−/− mice during homeostasis emphasizes that the role of FcRn during myeloid cell development in tissues should be targeted in future studies.

We observed that increased flt3 transcript expression correlated with an expanded CD103+ DC population in fcgrt−/− animals, supporting previous publications showing the systemic importance of this growth factor for DC development (40, 41). The explanation of how local IgG or commensal microbes may inhibit lung DC development or recruitment via FcRn remains elusive. Analysis of contrived commensal colonization of germfree fcgrt−/− animals or acquisition of lung immunity in neonates will have to address this issue in the future. Interestingly, an elegant study by Koch et al. recently showed a strikingly similar phenomenon in neonatal gut, where maternal IgG2b and IgG3 in the mucosa suppressed downstream T cell priming to mucosal antigens (42). Our results confirm that local dendritic cells are influenced by the prevailing signals derived from mucosal IgG during development. Mucosal IgG may limit access of antigens to lung or draining lymph nodes, as was previously shown to be the case in the context of barrier integrity of the gut epithelium in neonates (43).

Our results clearly demonstrate an indirect role for IgG in modulating cell recruitment to granulomas. We have shown here that the quantity and quality of IgG in airways are maintained by FcRn, which negatively impacts homeostatic populations of DC in the lung and subsequent T cell immunity and neutrophil recruitment during tuberculosis. This novel understanding of tissue-specific modulation of mucosal IgG isotypes in the lung by FcRn sheds light on the role of mucosal IgG for immune responses in the lung during homeostasis and bacterial disease.

ACKNOWLEDGMENTS
A.V. and S.H.E.K. conceived the study, and A.V., A.D., and L.L. designed the experiments. S.T.R. created M. tuberculosis GFP. A.V. wrote the manuscript, and A.D., L.L., and S.H.E.K. contributed editorial comments and interpretation of results. A.V., C.P., K.H., and U.Z. performed the experiments.

This work was supported by the Max Planck Society. We declare that no conflict of interest exists.

FUNDING INFORMATION
This work, including the efforts of Stefan H. E. Kaufmann, was funded by Max Planck Society.

REFERENCES
1. Sutherland DB, Suzuki K, Fagarasan S. 2016. Fostering of advanced mutualism with gut microbiota by immunoglobulin A. Immunol Rev 270:20–31. http://dx.doi.org/10.1111/imr.12384.
2. Blumberg RS, Koss T, Story CM, Barisani D, Polischuk J, Lipin A, Pablo L, Green R, Simister NE. 1995. A major histocompatibility complex class I-related Fc receptor for IgG on rat hepatocytes. J Clin Invest 95:2397–2402. http://dx.doi.org/10.1172/JCI117934.
3. Wellen AW, Baier N, Tang QJ, Van den Hof M, Blumberg RS, Lencer WI, Massol RH. 2013. Multivalent immune complexes divert FcRn to lysosomes by exclusion from recycling sorting tubules. Mol Biol Cell 24:2398–2405. http://dx.doi.org/10.1091/mbc.E13-04-0174.
4. Yoshida M, Kobayashi K, Kuo TT, Bry L, Glickman JN, Claypool SM, Kaser A, Nagashi T, Higgins DE, Mizoguchi E, Wakatsuki Y, Roope-nian DC, Mizoguchi A, Lencer WI, Blumberg RS. 2006. Neonatal Fc receptor for IgG regulates mucosal immune responses to luminal bacteria. J Clin Invest 116:2142–2151. http://dx.doi.org/10.1172/JCI27821.
5. Stapleton NM, Einarsson PT, Hendersen AF, Morson B, Orkin S, Adar M, Girard S, Hemeryck L, Minuth J, Smith PM, Tsai RP, Ribeiro R, Orkin S, Morison J. 2008. Severe congenital hyperbilirubinemia and neonatal hepatitis. J Pediatr 153:86–91. http://dx.doi.org/10.1016/j.jpeds.2008.04.046.
6. Funke M, O’Leary D, Woods A, Mellinghoff KI, Blumberg RS. 2009. The “asymmetric” dimethylarginine in 2019. Immunol Rev 236:44–60. http://dx.doi.org/10.1111/j.1600-065X.2009.00661.x.
7. Liu X, Li L, Yang Z, Farkas CM, Millar C, Chmiel J, Hsu H, Fredman P, Zhou P, Glimcher LH. 2010. FcRn regulates murine IgG homeostasis from birth and is essential for survival. J Exp Med 207:859–874. http://dx.doi.org/10.1084/jem.20091473.
8. Zhang Z, Chen X, Hernandez LD, Lipari P, Flattery A, Chen SC, Kramer S, Bolshov JD, Racine F, Cape H, Kelly CP, Thien Hien AG. 2015. Toxicin-mediated extracellular transport of anthrax antibodies facilitates protection against Clostridium difficile infection. Infect Immun 83:416–424. http://dx.doi.org/10.1128/IAI.02550-14.
9. Armitage CW, O’Meara CP, Harvie MC, Timms P, Blumberg RS, Beagley KW. 2014. Divergent outcomes following transcytosis of IgG targeting intracellular and extracellular chlamydial antigens. Immunol Cell Biol 92:417–426. http://dx.doi.org/10.1038/icb.2013.110.
10. Morel A, Passot C, Arnould C, Dumans A, Beaumont E, Goulieux-Gruart V, Roinegard P, Blanchard E. 2015. The neonatal Fc receptor does not modulate hepatitis C virus neutralization. J Gen Virol 96:1062–1066. http://dx.doi.org/10.1099/vir.0.000060.
11. Sakagami M, Omidi Y, Campbell L, Kandalaft LE, Morris CJ, Barar J, Gumbleton M. 2006. Expression and transport functionality of FcRn within rat alveolar epithelium: a study in primary cell culture and in the

isolated perfused lung, Pharm Res 23:270–279. http://dx.doi.org/10.1007/s11152-005-9226-0.
12. Spiekermann GM, Finn PW, Ward ES, Dumont J, Dickinson BL, Blumberg RS, Lencer WJ. 2002. Receptor-mediated immunoglobulin G transport across mucosal barriers in adult life: functional expression of FcRn in the mammalian lung. J Exp Med 196:303–310. http://dx.doi.org/10.1084/jem.20020400.
13. Lu L, Palaniyandi S, Zeng R, Bai Y, Liu X, Wang Y, Pauza CD, Roopenian DC, Zhu X. 2011. A neonatal Fc receptor-targeted mucosal vaccine strategy effectively induces HIV-1 antigen-specific immunity to genital infection. J Virol 85:10542–10553. http://dx.doi.org/10.1128/JVI.00541-11.
14. Ye L, Zeng R, Bai Y, Roopenian DC, Zhu X. 2011. Efficient mucosal vaccination mediated by the neonatal Fc receptor. Nat Biotechnol 29:158–163. http://dx.doi.org/10.1038/nbt.1742.
15. Saeland S, Aase A, Fae K, Houthuys E, Dorhoi A, Kreuchwig A, Furkert J, Cripps AW, Dunkley ML, Clancy RL. 2009. Mucosal and systemic immunization with killed Pseudomonas aeruginosa protect against acute respiratory infection in rats. Infect Immun 77:2652–2659. http://dx.doi.org/10.1128/IAI.00545-11.
16. Cooper AM. 2009. T cells in mucosal infection and disease. Curr Opin Immunol 21:378–384. http://dx.doi.org/10.1016/j.coi.2009.06.004.
17. Kozakiewicz L, Chen Y, Xu J, Wang Y, Dunussi-Joannopoulos K, Ou Q, Flynn JL, Porcelli SA, Jacobs WR, Jr, Chan J. 2013. B cells regulate neutrophilia during Mycobacterium tuberculosis infection and BCG vaccination by modulating the interleukin-17 response. PLoS Pathog 9:e1003472. http://dx.doi.org/10.1371/journal.ppat.1003472.
18. Kozakiewicz L, Phuh J, Flynn J, Chan J. 2013. The role of B cells and humoral immunity in Mycobacterium tuberculosis infection. Adv Exp Med Biol 783:225–250. http://dx.doi.org/10.1007/978-1-4614-6111-1_12.
19. Grode L, Seiler P, Baumann S, Hess J, Brinkmann V, Nasser Eddine A, Mann P, Goossen C, Bandermann S, Smith D, Bancroft GJ, Reyrat JM, van Soolingen D, Raupach A, Kaufmann SH. 2005. Increased vaccine efficacy against tuberculosis of recombinant Mycobacterium bovis bacillus Calmette-Guérin mutants that secrete lipoarabinomannan. J Clin Invest 115:2472–2479. http://dx.doi.org/10.1172/JCI24167.
20. Kremer L, Baulard A, Estajquier P, Poulain-Godefroy O, Locht S. 2005. Green fluorescent protein as a new expression marker in mycobacteria. Mol Microbiol 58:506–516. http://dx.doi.org/10.1111/j.1365-2958.2005.04936.x.
21. Cripps AW, Dunkley ML, Clancy RL. 1994. Mucosal and systemic immunizations with killed Pseudomonas aeruginosa protect against acute respiratory infection in rats. Infect Immun 62:1427–1436.
22. Moura-Alves P, Fae K, Houthuys E, Sela A, Fae K, Houthuys E, Stemerding AM, Stapleton NM, Janssen H, Verheul ML. 2014. Neutrophil Fc receptor-mediated immune complex uptake by human CD11b+ dendritic cells promotes immune complex-mediated glomerular disease. J Am Soc Nephrol 25:918–925. http://dx.doi.org/10.1681/ASN.2013050498.
23. Seher TD, Ludington WB, Barton GM. 2011. Competition for FcRn-mediated immunoglobulin G transport gives rise to short half-life of human IgG3 and offers therapeutic potential. J Immunol 187:5991–5998. http://dx.doi.org/10.4049/jimmunol.1100171.
24. Ginnoux F, Liu K, Helft J, Bogunovic M, Greter M, Hashimoto D, Price Y, Nussenzweig M, Merad M. 2009. The origin and development of nonlymphoid tissue CD103+ cells. J Exp Med 206:1159–1173. http://dx.doi.org/10.1084/jem.20082716.
25. Merrill WW, Naegel GP, Olszewski JS, Reynolds HY. 1985. Immunoglobulin G subclass proteins in serum and lavage fluid of normal subjects. Quantitation and comparison with immunoglobulins A and E. Am Rev Respir Dis 131:387–392. http://dx.doi.org/10.1164/ajrccm.131.3.387.
26. Guyer RL, Koshland ME, Knopf PM. 1976. Immunoglobulin binding by mouse intestinal epithelial cell receptors. J Immunol 119:577–583.
27. Stapleton NM, Andersson JT, Steerming AM, Bjarnarson SP, Verheul ML, Aase A, Fae K, Houthuys E, Stemerding AM, Stapleton NM, Janssen H, Verheul ML. 2014. Neutrophil Fc receptor-mediated immune complex uptake by human CD11b+ dendritic cells promotes immune complex-mediated glomerular disease. J Am Soc Nephrol 25:918–925. http://dx.doi.org/10.1681/ASN.2013050498.
28. Seher TD, Ludington WB, Barton GM. 2011. Competition for FcRn-mediated immunoglobulin G transport gives rise to short half-life of human IgG3 and offers therapeutic potential. J Immunol 187:5991–5998. http://dx.doi.org/10.4049/jimmunol.1100171.
29. Ginnoux F, Liu K, Helft J, Bogunovic M, Greter M, Hashimoto D, Price Y, Nussenzweig M, Merad M. 2009. The origin and development of nonlymphoid tissue CD103+ DCs. J Exp Med 206:1159–1173. http://dx.doi.org/10.1084/jem.20091756.
30. Waskow C, Liu K, Dassari-Jee G, Guernoprez P, Ginnoux F, Merad M, Shengelia T, Yao K, Nussenzweig M. 2008. The receptor tyrosine kinase Flt3 is required for dendritic cell development in peripheral lymphoid tissues. Nat Immunol 9:658–666. http://dx.doi.org/10.1038/ni.1615.
31. Koch MA, Reiner GL, Lugo KA, Kreuk LS, Stanbery AG, Ansaldo E, Seher TD, Ludington WB, Barton GM. 2016. Maternal IgG and IgA antibodies dampen mucosal T helper cell responses in early life. Cell 165:827–841. http://dx.doi.org/10.1016/j.cell.2016.04.055.
32. Gomez de Aguero M, Galán-Varonburg SC, Fuhrer T, Rupp S, Uchimura Y, Li H, Steinert A, Heikenwalder M, Helft J, Bogunovic M, Greter M, Hashimoto D, Price Y, Nussenzweig M, Merad M. 2009. The origin and development of nonlymphoid tissue CD103+ DCs. J Exp Med 206:1159–1173. http://dx.doi.org/10.1084/jem.20091756.