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## **Graphical abstract**



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# Allergen presensitization drives an eosinophildependent arrest in lung-specific helminth development

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This study investigates the relationship between helminth infection and allergic sensitization by assessing the influence of preexisting allergy on the outcome of helminth infections, rather than the more traditional approach in which the helminth infection precedes the onset of allergy. Here we used a murine model of house dust mite-induced (HDM-induced) allergic inflammation followed by *Ascaris* infection to demonstrate that allergic sensitization drives an eosinophil-rich pulmonary type 2 immune response (Th2 cells, M2 macrophages, type 2 innate lymphoid cells, IL-33, IL-4, IL-13, and mucus) that directly hinders larval development and reduces markedly the parasite burden in the lungs. This effect is dependent on the presence of eosinophils, as eosinophil-deficient mice were unable to limit parasite development or numbers. In vivo administration of neutralizing antibodies against CD4 prior to HDM sensitization significantly reduced eosinophils in the lungs, resulting in the reversal of the HDM-induced *Ascaris* larval killing. Our data suggest that HDM allergic sensitization drives a response that mimics a primary *Ascaris* infection, such that CD4<sup>+</sup> Th2-mediated eosinophil-dependent helminth larval killing in the lung tissue occurs. This study provides insight into the mechanisms underlying tissue-specific responses that drive a protective response against the early stages of the helminths prior to their establishing long-lasting infections in the host.

#### Introduction

Helminth parasites are well known for their capacity to modulate the parasite-directed host immune response (1, 2); with long-standing infection these parasites modulate the host response to bystander antigens/pathogens (3–5), to allergic diseases (6, 7), and to both orally and parenterally administered vaccines (8, 9). More controversially and largely on the basis of murine models, helminths have been implicated in ameliorating the severity of inflammatory bowel disease (10, 11), diabetes (12, 13), and arthritis (14), findings that have driven the concept of using these parasites or their excreted/secreted products, including their extracellular vesicles (15), for the treatment of inflammatory diseases (16, 17). However, the mechanisms by which helminths mediate these protective functions remained poorly defined.

Other studies, in contrast, have indicated that infection by helminths, more specifically by *Ascaris lumbricoides*, is associated with an increased incidence of allergic reactions (18–20) and that the presensitization to *Ascaris* antigens accelerates mite-specific IgE response upon mite antigen inhalation in murine models (21). More recently, it has been suggested that *Ascaris* larval migration in the lungs induces severe allergic airway disease in mice (22).

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Studies to date have attempted to elucidate the mechanisms underlying the regulation of allergic diseases by helminth infection (23). Helminth infections induce IL-10 responses in the host, which have been hypothesized to be a central mediator in regulating allergic diseases (24). The helminth-induced IL-10 directly modulates allergen-specific CD4+ effector responses (25, 26) or indirectly alters the IgG4/IgE balance in favor of IgG4 (27, 28). Both allergic disorders and helminth infections share an association with type 2 immune responses, characterized by the production of IL-4, IL-5, IL-9, IL-10, and IL-13, induction of IgG4 and IgE antibodies, and tissue and peripheral blood eosinophilia (29, 30). This type 2-associated immune response that drives an eosinophildominated environment has been implicated in protection against helminth infection but has also been shown to play a key role in the pathogenesis of allergic diseases (26, 31). The ability of eosinophils to control parasite levels has been extensively discussed in the literature over the last few decades (32), mainly in the context of primary, concomitant, or secondary infections (reinfection) using irradiated Onchocerca volvulus larvae (33), Trichinella spiralis larvae (34), Nippostrongylus brasiliensis and Heligmosomoides polygyrus adult worms (35), and Brugia malayi microfilariae (36) as models. In most cases of eosinophil-mediated killing of helminth larvae, the mechanisms underlying the helminth killing appear to involve either antibody-induced or complementinduced release of toxic eosinophil granule proteins and/or reactive oxygen intermediates by activated eosinophils (37) or in conjunction with other innate cells such as neutrophils (38).



**Figure 1. HDM sensitization followed by** *Ascaris* **infection limits the parasite burden by impairing larval development in the lungs.** (**A**) Experimental design scheme for HDM allergic sensitization followed by *Ascaris* infection. (**B**–**G**) Parasite burden in the liver (**B**) and lungs at different time points of evaluation (**C**–**F**), as well as lung-stage larval development by morphometric analysis of larvae recovered from HDM-*Ascaris*<sup>+</sup> and HDM+*Ascaris*<sup>+</sup> BALB/c mice (**G**). (**H**) Representative bright confocal image of the larvae recovered in the BAL at day 8 of infection (scale bars: 200 μm). Each symbol represents a single mouse, and the horizontal bars are the geometric means (GMs). *P* values are indicated on each graph. Three independent experiments were performed. Differences between HDM-*Ascaris*<sup>+</sup> (*n* = 17 mice total) and HDM+*Ascaris*<sup>+</sup> (*n* = 17 mice total) groups were considered statistically significant at *P* < 0.05 by unpaired Mann-Whitney test. Kruskal-Wallis test followed by Dunn's multiple-comparisons test was used for **G**.

Human population shifts have led to a reexamination of how helminth infections and allergy interface given that for many people allergic sensitization has preceded the acquisition of the helminth infection. Indeed, we have previously demonstrated that allergic sensitization to house dust mites (HDM), an environmental allergen, prior to the acquisition of a human filarial infection induces a markedly exaggerated parasite antigen–specific Th2-associated immune response (39). We hypothesized that this exaggerated type 2–dominated immune response could play a vital role in limiting parasite burden at the expense of parasite-associated morbidity.

Using the HDM model of murine allergic asthma inflammation and *Ascaris* spp. infection, we sought to identify the underlying mechanisms of how preexisting allergic sensitization influences the outcome of *Ascaris* infection. We found that sensitization with HDM impairs *Ascaris* migration and development through the underlying type 2-dominated, eosinophil-rich inflammation. Here, we dissect the mechanism by which allergic sensitization protects the host from helminth infection, highlighting how tissue-specific responses may control the early stages of helminth parasites before they establish long-lasting infections in the host.

#### Results

HDM sensitization followed by Ascaris infection inhibits larval development and reduces numbers of parasites in the lungs. It has been shown quite convincingly that the early Ascaris larval migration through the organs (termed larval ascariasis) in mice mimics with a certain degree of rigor what happens in acute infection of pigs and humans (definitive hosts), in terms of both the pattern of larval migration (40–42) and the highly polarized Th2 immune response in the lung (43, 44).

To determine whether pre-established pulmonary allergic inflammation driven by HDM alters the nature of a lung-migrating helminth infection, we examined larval migration to the liver (day 5) and to the lungs 5, 8, and 18 days postinfection (dpi) (Figure 1A). Our data show that at day 5 after *Ascaris* infection, HDM sensitization prior to infection (HDM<sup>+</sup>*Ascaris*<sup>+</sup>) did not affect the numbers of migrating larvae to the liver (Figure 1B) or to the lungs (Figure 1C). By 8 dpi, in contrast, there was a 70% reduction in the total worm burden in the lungs/bronchoalveolar lavage (BAL) of HDM<sup>+</sup>*Ascaris*<sup>+</sup> mice compared with HDM-unsensitized *Ascaris* infected mice (HDM<sup>-</sup>*Ascaris*<sup>+</sup>) (Figure 1, D–F). Moreover, larvae

isolated from the lungs and BAL fluid of the HDM<sup>+</sup>Ascaris<sup>+</sup> mice displayed significant reduction in size compared with larvae from HDM<sup>-</sup>Ascaris<sup>+</sup> (Figure 1, G and H), suggesting that HDM sensitization followed by Ascaris infection inhibits larval development and reduces numbers of parasites in the lungs. Notably, at 18 dpi there were no more larvae in the lungs, as all animals typically clear the infection by day 14 dpi.

Increased leukocyte infiltration, mucus production, and type 2 immune profile in the lungs of HDM+Ascaris+ mice. We hypothesized that the HDM-driven lung inflammation was limiting the parasite burden and larval development in HDM<sup>+</sup>Ascaris<sup>+</sup> mice. To test this hypothesis, we first evaluated the changes in the lung tissue by examining the infiltration of inflammatory cells into the lungs, the levels of chemokines that might underlie some of the cell migration into the lung tissue, the mucus production by epithelial cells in the airways, and the cytokine signature in the lung tissue. Sensitization with HDM followed by Ascaris infection (HDM+Ascaris+) caused a marked influx of inflammatory cells (leukocytes) to the lung tissue, in comparison with HDM-Ascaris<sup>+</sup> mice at 5 and 8 dpi. Interestingly, the HDM-induced cellularity in the lungs returned to baseline 18 days after the last sensitization. The number of leucocytes in the lungs (Figure 2A) as well as the histological analysis of the lung tissue, at 8 dpi, revealed an intense perivascular and peribronchial infiltrate of inflammatory cells (Figure 2B). This robust influx of leukocytes into the lungs of HDM+Ascaris+ compared with HDM-Ascaris+ mice was associated with a significant increase of CCL-11 (eotaxin) (Supplemental Figure 1A; supplemental material available online with this article; https://doi. org/10.1172/JCI127963DS1) and CCL-5 (RANTES) (Supplemental Figure 1B) in the lungs at 5 dpi, which continued to be elevated at the peak of larval migration 8 dpi. These levels returned to basal levels by 18 dpi. In contrast, CCL-2 (MCP-1) was driven in the Ascaris-infected mice at all 3 time points (Supplemental Figure 1C). Finally, there was no difference in CXCL-5 (LIX) among the groups and time points (Supplemental Figure 1E).

Lung tissue from HDM<sup>+</sup>Ascaris<sup>+</sup> mice had increased intensity of Alcian blue (AB)/periodic acid–Schiff (PAS) staining at 5 and 8 dpi in comparison with HDM<sup>+</sup>Ascaris<sup>-</sup> mice (Figure 2C), indicating a marked increased of mucus production by the goblet cells in the lung epithelium. In agreement with our AB/PAS staining intensity, HDM<sup>+</sup>Ascaris<sup>+</sup> mice had elevated gene expression of *Muc5a* (Figure 2D).

The levels of the major type 2-associated cytokines were measured in lung homogenates, including IL-4 (Figure 2E), IL-13 (Figure 2F), IL-33 (Figure 2G), and IL-5 (Figure 2H), as well as those other cytokines belonging to regulatory and type 1 inflammatory phenotypes (Supplemental Figure 2). Our data demonstrate that allergic sensitization with HDM prior to *Ascaris* infection induced a markedly significant increase in IL-4, IL-13, and IL-33 levels at day 5 and day 8 in HDM<sup>+</sup>*Ascaris*<sup>+</sup> mice compared with HDM<sup>-</sup>*Ascaris*<sup>+</sup> mice. At 8 dpi, HDM<sup>+</sup>*Ascaris*<sup>+</sup> mice also had a significant increase in the levels of the regulatory cytokine IL-10 (Supplemental Figure 2B) and the regulatory soluble decoy receptor IL-13Ra2 (Supplemental Figure 2C) when compared with HDM<sup>-</sup>*Ascaris*<sup>+</sup> mice. Finally, HDM allergic sensitization followed by *Ascaris* infection also induced an increase of TNF- $\alpha$  levels at day 5 and day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 2E), as well as IFN- $\gamma$  levels at day 8 (Supplemental Figure 3E). tal Figure 2F), in comparison with HDM-*Ascaris*<sup>+</sup> mice. Notably, only *Ascaris* infection induced a significant increase of IL-4, IL-13, IL-13Ra2, IL-1 $\beta$ , and TNF- $\alpha$  levels in comparison with naive mice. Interestingly, the IL-5 and IL-6 levels showed a quite different pattern from the IL-4, IL-13, IL-33 in that a marked downregulation of that cytokine occurred at day 8 in HDM<sup>+</sup>*Ascaris*<sup>+</sup> compared with only HDM<sup>-</sup>*Ascaris*<sup>+</sup> mice.

Allergic sensitization coincident with Ascaris infection drives strong innate and adaptive type 2 immune responses in the lungs associated with an eosinophil- and M2 macrophage-rich environment. The robust inflammatory response induced by the HDM allergic sensitization followed by Ascaris infection was characterized both phenotypically and functionally. Initially, a CD4<sup>+</sup> T cell differentiation-flow cytometry panel was used to assess the nature and frequency of Th1, Th2, and Th17 cells among the lung-purified leukocytes in each group (Figure 3A). Our data demonstrated that at 5 dpi, HDM allergic sensitization drove a significant increase in the frequency of IL-5- or IL-13producing CD4+ T cells (Th2) in HDM+Ascaris- mice, as well as in HDM<sup>+</sup>Ascaris<sup>+</sup> mice, compared with HDM<sup>-</sup>Ascaris<sup>+</sup> mice and also with the naive HDM-Ascaris- mice. The magnitude of Th2 cell differentiation driven by allergic sensitization continued to increase at 8 dpi in HDM+Ascaris- mice and in HDM+Ascaris+ mice compared with HDM-Ascaris+ and naive mice. At 8 dpi, the migrating larvae induced an increase of Th2 cells in the lungs of HDM-Ascaris+ mice compared with naive mice. Finally, at 18 dpi, the frequency of Th2 cells driven by HDM allergic sensitization in the HDM+Ascaris- mice progressively decreased, with frequencies near the baseline levels seen in naive mice. At 18 dpi, migrating Ascaris larvae alone drove a marked increase in the frequency of Th2 cells in HDM-Ascaris+ mice compared with naive animals. However, this robust upregulation of Th2 cells mediated by Ascaris was not as dramatic in HDM<sup>+</sup>Ascaris<sup>+</sup> mice, suggesting a tolerance to the parasite-induced Th2 response in HDM-sensitized animals or that the lower parasite burden in the lungs contributed to the attenuated Ascaris-induced CD4+ T cell response compared with HDM-Ascaris+ (Figure 3B). Interestingly, the HDM allergic sensitization was associated with a modest increase in IFN-γ-producing (Th1) and IL-17A-producing (Th17) CD4<sup>+</sup> T cells in the HDM<sup>+</sup>Ascaris<sup>-</sup> and HDM<sup>+</sup>Ascaris<sup>+</sup> mice compared with HDM-Ascaris+ and naive mice at 5 dpi and 8 dpi. At 18 dpi, the frequency of Th1 and Th17 response driven by the allergic inflammation had also returned to baseline. When the pulmonary Th1 and Th17 responses in the HDM-Ascaris+ mice were evaluated, there were no significant increases in the frequency of IFN-y- or IL-17-producing CD4+ T cells in comparison with naive mice, suggesting that Ascaris infection itself induces a polarized CD4<sup>+</sup> Th2 response in the lungs at day 18 (Figure 3, C and D).

To assess the drivers of the augmented Th2-associated lung inflammation, we examined the frequency of innate lymphoid cells (ILCs; CD45<sup>+</sup>Lin<sup>-</sup>CD4<sup>-</sup>TCR- $\beta$ <sup>-</sup>Thy1.2<sup>+</sup>) and their subsets (by using the expression of the signature cytokines IFN- $\gamma$ , IL-5/IL-13, and IL-17A) in the lungs at 8 dpi (Figure 3E). As seen, in the naive mice, the majority of the ILCs were type 1 ILCs (ILC1s) (73%), with 26% being ILC2s and 1% ILC3s. In HDM<sup>-</sup>Ascaris<sup>+</sup> mice, the subset composition of the ILCs, although increased in frequency, was very similar to that of the naive mice, with 78% ILC1s, 22%



**Figure 2.** Allergic sensitization followed by *Ascaris* infection drives a marked influx of inflammatory cells to the lungs, increased mucus production, and a **robust type 2 immune profile.** Cellularity quantification (**A**), representative H&E perivascular and peribronchial histological analysis (original magnification, ×8) of the inflammatory influx of leukocytes at 8 dpi (**B**), AB/PAS staining (original magnification, ×8) for mucus production by the goblet cells in the lung epithelium (in blue; arrows) at 8 dpi (**C**), relative gene expression of Muc5ac (**D**), and pulmonary tissue levels of IL-4 (**E**), IL-13 (**F**), IL-33 (**G**), and IL-5 (**H**) in the lung's homogenates of naive HDM<sup>-</sup>*Ascaris*<sup>-</sup> (black circles) (a), HDM<sup>+</sup>*Ascaris*<sup>-</sup> (red squares) (b), HDM<sup>-</sup>*Ascaris*<sup>+</sup> (blue triangles) (c), and HDM<sup>+</sup>*Ascaris*<sup>+</sup> (green triangles) (d) BALB/c mice at day 5 (*n* = 9 mice per group), day 8 (*n* = 10 mice per group), and day 18 (*n* = 10 mice per group). Each symbol represents a single mouse, and the horizontal bars are the GMs. *P* value is indicated in each graph. Kruskal-Wallis test followed by Dunn's multiple-comparisons test was used for all comparisons; \*significantly different (*P* < 0.05) from naive (HDM<sup>-</sup>*Ascaris*<sup>-</sup>) group.

ILC2s, and 0% ILC3s. In contrast, the HDM allergic sensitization followed by *Ascaris* infection (HDM<sup>+</sup>*Ascaris*<sup>+</sup>) at 8 dpi induced a marked increase in the total number of ILCs with a marked shift in the proportion of ILC subsets with a slight predominance of ILC2s (50%) compared with ILC1s (49%) and ILC3s (1%). Furthermore, at 18 dpi, while the HDM-induced ILC2s decreased toward the baseline numbers in the lungs, there was a marked increase in the *Ascaris*-driven ILC2s (HDM<sup>-</sup>*Ascaris*<sup>+</sup>). Interestingly, the *Ascaris* infection in the allergic mice (HDM<sup>+</sup>*Ascaris*<sup>+</sup>) led to lower numbers of ILC2s than in the HDM<sup>-</sup>Ascaris<sup>+</sup> mice, suggesting either that HDM sensitization downregulated ILC2 expansion/recruitment or that the lower levels of migrating larvae contributed to a lack of ILC2s (Supplemental Figure 3).

The participation and function of granulocytes and myeloid cells in the context of helminth infection with or without a preallergic conditioning were also evaluated (Figure 4). The most marked alteration in the number of lung cells driven by the HDM sensitization was related to the eosinophils (Figure 4A). HDM



**Figure 3. HDM sensitization followed by** *Ascaris* **infection drives a strong innate and adaptive type 2 immunity in the lungs.** (**A**) Representative flow cytometry dot plot of lung homogenates, gated on Lin<sup>-</sup>CD45<sup>+</sup>TCR- $\beta$ <sup>+</sup>CD4<sup>+</sup> T cells, showing the frequencies of IL-5 and IL-13 (top) and IFN- $\gamma$  and IL-17 (bottom) in the different groups at day 8. (**B–D**) The net frequency of IL-5<sup>+</sup> or IL-13<sup>+</sup> CD4<sup>+</sup> Th 2 cells (**B**), IFN- $\gamma^{+}$  CD4<sup>+</sup> Th 1 cells (**C**), and IL-17A<sup>+</sup> CD4<sup>+</sup> Th17 cells (**D**) was assessed at different time points: day 5 (*n* = 9 mice per group), day 8 (*n* = 10 mice per group), and day 18 (*n* = 10 mice per group). Boolean analysis was performed to exclude multifunctional CD4<sup>+</sup> T cells expressing IL-5, IL-13, IFN- $\gamma$ , and IL-17 simultaneously. (**E**) Frequency of total innate lymphoid cells (ILCs) gated as CD45<sup>+</sup>Lin<sup>-</sup>CD4<sup>-</sup> TCR- $\beta^{-}$ Thy.1.2<sup>+</sup> in the different groups at day 8 of infection, as well as the proportion of ILC1s, ILC2s, and ILC3s, based on the expression of the signature cytokines IFN- $\gamma$ , IL-5/IL-13, and IL-17A, respectively, after PMA/ionomycin stimulation. Each symbol represents a single mouse, and the horizontal bars are the GMs. Net frequency of CD4<sup>+</sup> T cells was calculated by subtraction of the baseline frequency (nonstimulated) from the frequency following stimulation with PMA/ionomycin. *P* values are indicated in each graph. Differences were considered statistically significant at *P* < 0.05 by Kruskal-Wallis test followed by Dunn's multiple-comparisons test, used for all comparisons; \*significantly different (*P* < 0.05) from naive (HDM<sup>-</sup>Ascaris<sup>-</sup>) group.



allergic sensitization induced a marked influx of Siglec-F<sup>+</sup>C-D11c<sup>-</sup> eosinophils to the lungs. When compared with naive mice, HDM<sup>+</sup>Ascaris<sup>-</sup> mice at 5 dpi and at 8 dpi had markedly increased numbers of lung eosinophils. Notably, at 18 dpi (18 days after the last HDM sensitization), the numbers of eosinophils significantly decreased to almost baseline levels. Notwithstanding, in the context of HDM sensitization followed by Ascaris infection, HDM<sup>+</sup>Ascaris<sup>+</sup> mice showed a robust tissue eosinophilia when compared with HDM<sup>-</sup>Ascaris<sup>+</sup> mice at 5 dpi. This HDM-induced eosinophilic environment continued to be dominant during the peak of larval migration (8 dpi) in the lungs of HDM<sup>+</sup>Ascaris<sup>+</sup> mice compared with HDM<sup>-</sup>Ascaris<sup>+</sup> mice (Figure 4B).

Not only were there increased numbers of eosinophils in the lungs of HDM<sup>+</sup>Ascaris<sup>-</sup> and HDM<sup>+</sup>Ascaris<sup>+</sup> mice at 5 dpi, but these eosinophils released more of the eosinophil-specific granule protein eosinophil peroxidase (EPO), suggesting these eosinophils were activated and showed evidence of degranulation in the tissue. Interestingly, at 8 dpi, both HDM<sup>-</sup>Ascaris<sup>+</sup> mice and HDM<sup>+</sup>Ascaris<sup>+</sup> mice had increased EPO levels when compared with HDM<sup>+</sup>Ascaris<sup>-</sup> and naive mice. Finally, at 18 dpi, all groups had EPO levels similar to homeostatic levels seen in naive, HDM<sup>+</sup>Ascaris<sup>-</sup>, HDM<sup>-</sup>Ascaris<sup>+</sup>, and HDM<sup>+</sup>Ascaris<sup>+</sup> mice (Figure 4C). No relevant differences were observed among the groups and time points in the numbers of Ly6G<sup>+</sup> neutrophils.

We next evaluated the number of F4/80<sup>+</sup>CD11b<sup>+</sup> macrophages in the lung tissue of the animals, and we observed a pattern similar to that seen for the eosinophils (Figure 4, D and E). HDM allergic sensitization induced a marked increase of macrophages in the lungs of HDM<sup>+</sup>*Ascaris*<sup>-</sup> mice as well as HDM<sup>+</sup>*Ascaris*<sup>+</sup> mice at 5 dpi and 8 dpi, returning to baseline levels at 18 dpi. Interestingly, in parallel to what was seen for the eosinophils, in HDM<sup>-</sup>*Ascaris*<sup>+</sup> mice, there was a marked increase in the number of these macrophages from day 5 to day 18 post-infection, in comparison with naive mice.

Analysis in the lung tissue using markers of M1 (iNOS) and M2 (arginase-1 [Arg-1]) macrophages by quantitative PCR revealed that the HDM allergic sensitization in HDM<sup>+</sup>Ascaris<sup>-</sup> and HDM<sup>+</sup>Ascaris<sup>+</sup> mice drove the lung macrophages toward the M2 phenotype (17.6-fold change in Arg-1) in comparison with HDM<sup>-</sup>Ascaris<sup>+</sup> mice at 5 dpi. These data suggest that when the first group of Ascaris larvae reached the lung tissue of HDM-sensitized mice at 5 dpi, there was already present an M2 macrophage-rich environment not seen in HDM<sup>-</sup>Ascaris<sup>+</sup> mice. At 8 dpi, the Ascaris infection by itself also induced a significant increase of Arg-1 expression, suggesting that the primary migration of helminth larvae in the lungs is also associated with an increase in M2 macrophages (Figure 4F). No relevant differences were observed among the groups in macrophage differentiation at 18 dpi.

Ascaris migration and development occur normally in the absence of eosinophils in allergic animals. Eosinophils are critical to both allergic inflammation and host resistance to some helminth infections. Having identified an expansion of eosinophils in the lung tissue of HDM<sup>+</sup>Ascaris<sup>+</sup> mice, we hypothesized that the influx of eosinophils in mice with HDM sensitization was impairing the migration and development of Ascaris larvae. To test this, we sensitized eosinophil-deficient mice (\DeltadblGATA) with HDM before Ascaris infection (Figure 5A). As expected, HDM<sup>+</sup>Ascaris<sup>+</sup> WT mice had a marked increase in the frequency of Siglec-F<sup>+</sup>CD11c<sup>-</sup> eosinophils in the lung,

while eosinophils were absent in HDM<sup>+</sup>Ascaris<sup>+</sup> AdblGATA mice (Figure 5B). HDM sensitization in WT mice decreased the number of larvae in the lung tissue and BAL in comparison with HDMunsensitized Ascaris-infected WT mice, as seen previously (Figure 1). Strikingly, allergic sensitization in eosinophil-deficient mice had no impact on the larval migration (Figure 5C). In parallel with our migration findings, Ascaris development occurred normally in the absence of eosinophils, regardless of whether or not mice were sensitized with HDM (Figure 5D). These findings suggest that eosinophils are the cell type that impairs Ascaris development in allergic sensitized mice. Finally, the immune response of HDMinduced allergy and concomitant Ascaris infection in the absence of eosinophils was also evaluated. The quantification of type 2 cytokines in the lung's homogenate demonstrated that in the absence of eosinophils there was a marked reduction in the levels of IL-4 from HDM<sup>+</sup>Ascaris<sup>+</sup> eosinophil-deficient mice compared with HDM<sup>+</sup>Ascaris<sup>+</sup> WT mice, although there was no difference in the levels of IL-13 nor IL-33, which remained elevated in the 2 mouse strains (Figure 5F). The increase in the frequency of HDM-induced CD4+ Th2 cells in the lungs of HDM+Ascaris+ mice was not affected by the absence of eosinophils (Figure 5G), nor were there differences in the number of M2 macrophages (Figure 5H) or in mucus production (Figure 5I).

HDM-induced immunity to Ascaris parasites is dependent on eosinophils driven by CD4<sup>+</sup> T cells. After showing that HDMinduced larval arrest was related to the presence of eosinophils in the lungs, we sought to determine whether the HDM-induced eosinophil-mediated effect on Ascaris larval development was dependent on CD4<sup>+</sup> T cells. Thus, CD4<sup>+</sup> T cells were depleted in HDM<sup>+</sup>Ascaris<sup>+</sup> mice either before HDM allergic sensitization (day 0) or after HDM sensitization (day 18) (Figure 6). As can be seen, only the in vivo administration of anti-CD4 neutralizing antibodies prior to and during the HDM allergic sensitization (day 0) significantly impaired the HDM-induced reduction in larval numbers (and developmental arrest) (Figure 6, B–D), as neutralizing CD4<sup>+</sup> T cells following HDM allergic sensitization had little or no effect on larval numbers or development.

Moreover, the in vivo administration of CD4 neutralizing antibodies prior to the HDM allergic sensitization significantly reduced the frequency and numbers of eosinophils in the lungs (a reduction similar to that of the numbers found in nonallergic animals) and also diminished the HDM-specific IgE and IgG1 antibody levels. These data suggest that the type 2-dominant HDM-induced effect on *Ascaris* parasites is dependent on eosinophils driven by CD4<sup>+</sup> T cells (Figure 6E).

Delayed infection in HDM-sensitized mice prevents migration. While it is likely that humans would be sensitized to HDM early in life but infected with a helminth well after HDM exposure, we next investigated whether allergic inflammation still limited parasite infection after a delay. WT mice were sensitized with HDM and then, 20 days later, were challenged with HDM and then infected with *Ascaris* (Figure 7A). As noted previously, 20 days after the last intranasal sensitization with HDM, the frequency of eosinophils, M2 macrophages, and Th2 cells and cytokine and chemokine levels have already returned to baseline. Thus, we evaluated the parasite burden and lung-specific immune response in HDM<sup>-</sup>*Ascaris*<sup>+</sup> and HDM<sup>+</sup>*Ascaris*<sup>+</sup> groups. As seen previously, at 8 dpi the total burden



in the lungs (tissue + BAL) showed a marked approximately 68% reduction in the number of larvae in HDM+Ascaris+ mice compared with HDM-Ascaris+ mice. Surprisingly, the HDM+Ascaris+ (20 days) mice also showed a marked approximately 45% reduction in the parasite burden when compared with HDM-Ascaris+ mice (Figure 7B). The immunophenotyping of inflammatory cell subsets in the lungs demonstrated that, although of lower magnitude than observed in HDM<sup>+</sup>Ascaris<sup>+</sup> mice following the standard sensitization protocol, Ascaris infection 20 days after HDM sensitization induced an increase in the absolute number of eosinophils (Figure 7C), and the frequency of Th2 cells (Figure 7D), in comparison with HDM-Ascaris+ mice. Interestingly, no differences were observed in the IL-4 levels in the lung homogenates of HDM<sup>+</sup>Ascaris<sup>+</sup> (20 days) mice compared with HDM<sup>-</sup>Ascaris<sup>+</sup> mice (Figure 7E). There was, however, a marked increase in IL-13 and IL-13Ra2 levels in HDM+Ascaris+ mice compared with HDM<sup>-</sup>Ascaris<sup>+</sup> animals (Figure 7, F and G), highlighting a potential role for IL-13 signaling as a secondary mechanism of HDMinduced protective immunity to helminths.

#### Discussion

Allergic diseases and helminth infections are 2 distinct and relatively nonoverlapping geographical public health concerns (45) that together affect more than 3 billion people (46, 47). However, globalization and its extensive population movement have resulted in the intersection of these once separate public health problems.

Our study takes a novel look at the relationship between helminth infection and allergic sensitization/disease by examining the influence of preexisting allergic sensitization on the outcome of helminth infections, rather than the more traditional approach of examining the converse (48). As has been studied quite extensively, chronic helminth infection has been associated with diminished allergic reactivity. As reviewed by many (23, 49, 50), among the various hypotheses to explain the modulatory influence of helminth infection on allergy, the IL-10-induced suppression of Th2 effector responses (51, 52) and/or the expansion of natural and parasite-induced regulatory T cells (Tregs) have been the leading candidates (26, 53, 54). Nevertheless, "allergic-like symptoms" have been associated with relatively acute helminth infections (e.g., urticaria in many helminth infections, angioedema in filarial infections, asthma in slaughterhouse workers exposed to Ascaris, swimmer's itch in schistosomiasis). In this way, understanding whether allergic inflammation can regulate helminth parasite intensity and development in their early migration phase provides a new perspective for this interaction. The present study provides a window on the mechanisms underlying tissue-specific responses that can drive a protective response against the early stages of the helminth parasites prior to their establishing long-lasting infections in the host.

Here we used a murine model of HDM-induced allergic pulmonary inflammation followed by *Ascaris* infection to demonstrate that allergic sensitization to environmental aeroallergens (e.g., HDM) in the context of a subsequent helminth infection drives an eosinophil-rich pulmonary type 2 response that directly hinders larval development and markedly reduces parasite burden. This effect is dependent on the presence of eosinophils, as eosinophil-deficient mice were unable to limit parasite development or numbers. In the present study, we have carefully dissected the immunobiology of larval ascariasis in the lungs of naive mice and compared it with the course of the infection in mice with preexisting allergic asthma. By using different time points of investigation, we have demonstrated that during primary exposure to Ascaris, the larvae in the lungs elicit a strong innate tissue response characterized by increased levels of IL-4, IL-5, IL-6, and IL-33 cytokines as well as marked increases in the chemokines CCL-11 (eotaxin), CCL-2 (MCP-1), and CXCL-10 (IP-10). Moreover, there is a small but measurable accumulation of CD4+ effector Th2 cells, activated eosinophils, and alternatively activated macrophages (M2). Despite this type 2-associated inflammation, the environment still allows lungstage Ascaris larvae to develop in the tissue, increasing considerably in length and width, and to reach the airways while causing significant hemorrhage and tissue damage. If, however, there is HDM sensitization prior to infection with Ascaris, there is an even more profound type 2 inflammation characterized initially by even more elevated levels of CCL-11, CCL-5, IL-25, and IL-33 followed by an increase in both IL-5/IL-13-producing ILC2s and CD4+ Th2 cells.

It has been shown previously that during HDM sensitization airway epithelial cells can sense allergens (e.g., HDM) through Toll-like receptor 4 (TLR4) (55), which in turn leads to the release of chemokines, cytokines, and signals, including thymic stromal lymphopoietin (TSLP), granulocyte-macrophage colonystimulating factor (GM-CSF), IL-25, and IL-33, that can activate and recruit other immune effector cells to the site of inflammation (56). This process can trigger the differentiation of naive T cells into adaptive CD4<sup>+</sup> Th2 cells, induce eosinophilic inflammation, polarize toward M2 macrophages, and drive mucus production and airway hyperresponsiveness (57-59). In the present study, the HDM allergen-driven inflammation following Ascaris infection induces an IL-33-driven inflammatory milieu with dramatic increases of activated eosinophils, differentiation of lung macrophages toward the M2 phenotype, and an IL-4- and IL-13-rich environment associated with mucus production by goblet cells in the epithelium (Figure 8).

IL-13 is a pleiotropic cytokine predominantly produced by Th2 cells that mediates its effects through the IL-13Ra1/IL-4Ra heterodimer to induce genes specific to Th2 inflammation (60). This Th2-associated response has been shown to lead to a protective response to certain helminth parasites (60) but, at the same time, can play an important role in the pathogenesis of asthma (61). It has been shown that IL-13 signaling through IL-13Ra1 induces the upregulation of IL-13R $\alpha$ 2 decoy receptor (62), which serves to downregulate/limit the IL-13-mediated responses in vivo (63-65). In fact, HDM allergic sensitization followed by Ascaris infection markedly upregulated levels of both IL-13 and IL-13Ra2 at 8 dpi when compared with Ascaris infection alone. Moreover, the high expression of both IL-13 and IL-13Ra2 was also associated with the eosinophil-dominated type 2 response in the last protective phenotype in the allergic animals with a 20-day delay before infection with the helminth. These findings suggest a potential role for the regulation of the IL-13/IL-13Ra2 axis in the helminth-protective lung-specific responses induced by the allergic sensitization.

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**Figure 7. HDM allergic sensitization induces a persistent protective type 2 immunity followed by later** *Ascaris* **infection.** (**A**) Experimental design scheme for unsensitized mice followed by *Ascaris* infection, HDM<sup>-</sup>*Ascaris*<sup>+</sup> (blue triangles) (*n* = 7 mice); and HDM allergic sensitization followed by *Ascaris*<sup>+</sup> (green triangles) (*n* = 7 mice), infected right after the last sensitization with HDM, and HDM<sup>+</sup>*Ascaris*<sup>+</sup> (20 days) (purple triangles) (*n* = 7 mice), infected 20 days after the last sensitization with HDM. (**B**) Total parasite burden in the lung tissue and BAL at day 8 of infection. (**C** and **D**) Characterization of the lung-specific immune response by flow cytometry, showing eosinophils (**C**) and IL-5<sup>+</sup> or IL-13<sup>+</sup> CD4<sup>+</sup> Th2 cell frequency (**D**). (**E**-**G**) IL-4, IL-13, and IL-13Ra2 levels quantified in the lung homogenates of each group. Each symbol represents a single mouse, and the horizontal bars are the GMs. *P* values are indicated in each graph. Kruskal-Wallis test followed by Dunn's multiple-comparisons analysis was used for differences among the groups; \*significantly different (*P* < 0.05) from HDM<sup>-</sup>*Ascaris*<sup>+</sup> group by the same analyses.

Although the mechanisms of helminth killing in the tissue prior to adult worm expulsion remain largely unstudied, studies using the rodent helminth parasites *Nippostrongylus brasiliensis* and *Heligmosomoides polygyrus* have identified some important mediators associated with the control of the larval migration in the tissue. However, these have typically been seen in the context of secondary infections after the initial infection had been cleared. It may be, therefore, that the HDM sensitization mirrors a primary helminth infection such that the *Ascaris* infection (following HDM sensitization) initiates the kind of response that clearly controls larval tissue penetration and development. It is clear that the extraordinarily robust lung-specific response results in larval developmental arrest (and a 70% reduction in the parasite burden), a process that appears largely to be mediated by eosinophils.

Although eosinophilia is a hallmark of tissue-invasive helminth infections (37), the role of these cells in controlling the infection is unclear. The capacity of eosinophils to kill helminth nematodes has been extensively studied using in vitro assays and in vivo models. O'Connell et al. (66) suggested that eosinophils can kill *Strongyloides stercoralis* larvae through an eosinophil granule protein-dependent mechanism if other effector cells are absent. Moreover, it has been shown that eosinophil granule proteins are toxic to microfilariae of *Brugia* parasites in vitro (67). Recently, it has been shown that *H. polygyrus* intestinal infection led to a significant increase in immune-mediated killing of *N. brasiliensis* larvae within the lung of coinfected mice. This enhanced protection was associated with IL-33-mediated activation of CD4<sup>+</sup> Th2 cells, which orchestrated IL-5-mediated eosinophil-dependent larval killing (68). In addition, our results suggest that HDMinduced reduction in *Ascaris* numbers and arrested development is dependent on eosinophils in the lungs driven by CD4<sup>+</sup> T cells (see Figure 6). However, our data also demonstrate that once a type 2 allergic environment is established, the CD4<sup>+</sup> Th2 cells play little or no role in its maintenance.

When *Ascaris* larvae reached the lungs of allergic mice, they found a marked IL-33/CD4<sup>+</sup> type 2-dominated eosinophilic environment with elevated levels of EPO in the lung tissue, which was not observed in nonallergic but *Ascaris*-infected mice. In addition to eosinophil activation and degranulation, the release of eosinophil extracellular traps (EETs) may be an additional mechanism of killing or immune modulation, as already demonstrated for neutrophils (69) in a process termed NETosis (70) that can contribute to antimicrobial defense (71). Indeed, it has been shown that eosinophils from patients with allergic bronchopulmonary aspergillosis induce the release of EETs upon stimulation with the fungus *Aspergillus fumigatus* (72). Moreover, extracellular DNA traps generated by eosinophils have also been demonstrated in human atopic asthmatic airways (73, 74).

Finally, in the absence of HDM stimulation, *Ascaris*-infected mice demonstrated type 2-dominated lung inflammation, but only after 18 dpi. It has been suggested that this response in a primary helminth infection plays a key role in tissue remodeling and prevention of reinfection (75). Huang et al. (76) suggested that a primary infection with the helminth *Trichinella spiralis* induces



**Figure 8. Working hypothesis model.** Left: In a primary exposure to *Ascaris* parasites, after penetrating the intestine at the level of the cecum or proximal colon, the infective L3-stage larvae migrate through the portal vessels to the liver and subsequently to the lungs in their quest to reach the airways. Once in the lungs they migrate through the parenchyma, leading to a marked influx of neutrophils and correlated levels of IL-6, and penetrate into the alveolar spaces, causing bleeding and mechanical damage in the organ. These migrating *Ascaris* larvae, while growing in size toward the molting to L4 stages of biological development, induce a local inflammatory response in the lungs with a signature type 2 response, initially by early IL-5 levels, and later by IL-4 and IL-13 levels, culminating in the differentiation of M2 macrophages and eosinophilia in the tissue. However, this robust type 2 response is only fully mounted when most of the larvae have reached the airways, allowing the establishment of a chronic and long-standing infection. Right: HDM allergic sensitization prior to infection with *Ascaris* induces a robust Th2-dominated inflammatory response in the lung characterized by an increase in both IL-5/ IL-13-producing type 2 innate lymphoid cells (ILC2s) and eosinophils. This allergen-driven inflammation in the lungs leads to an IL-4- and IL-13-rich environment that drives the differentiation of lung macrophages toward the M2 phenotype expressing arginase-1. When the *Ascaris* larvae migrate from the circulation to the lung tissue and airways in this HDM-sensitized environment, they encounter an eosinophil-dominated type 2 response, which leads to a 70% reduction in the number of lung-stage *Ascaris* larvae and a profound larval developmental arrest, allowing the reduction of intensity and burden of disease, inducing lower bleeding and mechanical damage in the tissue during their migration.

an eosinophil-dominant protective immunity in tissue of the host against reinfection. Moreover, it has been recently demonstrated that multiple exposures to *Ascaris* parasites in mice induced a remarkably protective lung-specific mixed type 2 and type 17 immune response associated with elevated number of eosinophils in the lung tissue and airways (77).

Indeed, we have previously shown by using cells from *Loa loa*-infected travelers, who present clinically with allergic-type symptoms much more frequently than those born and bred in *Loa loa*-endemic regions of the world (78, 79) and who had evidence of HDM allergic sensitization prior to the acquisition of their filarial infection, that there was a marked expansion of parasite antigen-specific CD4<sup>+</sup> Th2 cells that were in turn associated with marked elevation of IgE and eosinophilia, in comparison with nonallergic but *Loa loa*-infected patients (39). Like these travelers, our data from this murine system suggest that allergic sensitization coincident with helminth infection drives an exaggerated eosinophil-rich pulmonary type 2 immune response in the lungs, mediated predominantly by the allergic sensitization that occurs

prior to the development of the *Ascaris*-specific response in the lung. This in turn limits helminth parasite numbers. It is very likely that HDM allergic sensitization acts not unlike a primary *Ascaris* infection, such that a lung-specific type 2-mediated eosinophildependent helminth larval killing in the tissue occurs and limits (like a secondary infection) parasite numbers. Studies to increase understanding of the mechanisms by which HDM-driven eosinophils are activated and drive helminth larval killing are under way.

#### Methods

*Mice*. Wild-type (WT) BALB/c mice (male, 8 weeks old; Taconic Farm) were used throughout the study.  $\Delta$ dblGATA mice (male, 8 weeks old) were provided by Helene Rosenberg, Laboratory of Allergic Diseases, National Institute of Allergy and Infectious Diseases, NIH, and bred in-house at the NIH Animal Facility.

HDM-induced allergic inflammation. Mice were anesthetized with isoflurane and sensitized intranasally with 200  $\mu$ g in 30  $\mu$ L of *Dermatophagoides pteronyssinus* HDM extract (Greer Laboratories) at day 0 and day 7. At days 14, 16, and 18, mice were given 50  $\mu$ g HDM extract

#### **Table 1. Key antibodies**

Antibody	Source	Catalog number
APC–eFluor 780 anti–mouse TCR- $\beta$ (clone H57-597)	eBioscience	47-5961-82
PerCP-Cy5.5–anti–mouse CD4 (clone RM4-5)	BD Pharmingen	550954
FITC–anti–mouse CD45 (clone 30-F11)	BioLegend	103108
BV605–anti–mouse Thy1.2 (clone 53-2.1)	BioLegend	140318
BV421-streptavidin	BioLegend	405225
PE-Cy7–anti–mouse IFN-γ (clone XMG1.2)	eBioscience	25-7311-82
PE-anti-mouse IL-13 (clone eBio13A)	eBioscience	12-7133-82
AF 700–anti–mouse IL-17A (clone TC11-18H10.1)	BioLegend	506914
APC–anti–mouse IL-5 (clone TRFK5)	BD Pharmingen	554396
Biotin-TER119 (clone TER-119)	eBioscience	13-5921-82
Biotin-CD19 (clone MB19-1)	eBioscience	13-0191-85
Biotin- $\gamma\delta$ TCR (clone eBiogI3)	eBioscience	13-5711-85
Biotin-Gr1 (clone RB6-8C5)	eBioscience	13-5931-85
Biotin-CD11c (clone N418)	eBioscience	13-0114-85
Biotin-CD11b (clone M1/70)	eBioscience	13-0112-85
Biotin-CD49b (clone DX5)	eBioscience	13-5971-85
Biotin–TCR-β (clone H57-597)	eBioscience	13-5961-82
BV605–anti–mouse CD11b (clone M1/70)	BioLegend	101257
PE-Cy7–anti–mouse F4/80 (clone BM8)	BioLegend	123114
FITC-anti-mouse CD11c (clone HL3)	BD Pharmingen	553801
V500–anti–mouse MHC-II (clone M5/114.15.2)	BD Horizon	562366
APC-Cy7-anti-mouse Ly6C (clone AL-21)	BD Pharmingen	560596
PE-anti-mouse Siglec-F (clone E50-2440)	BD Pharmingen	552126
PE-Texas red-anti-mouse Ly6G (clone 1A8)	BD Horizon	562700
FITC–anti–mouse CD8b (clone eBioH35-17.2)	Invitrogen	11-0083-85

in a volume of 30  $\mu$ L intranasally. Mice were infected with *Ascaris* at day 18, immediately after the last sensitization with HDM by oral gavage. Tissues were examined immunologically at day 23 (5 dpi; liver and lungs), day 26 (8 dpi; lungs and BAL), and day 36 (18 dpi; lungs).

Parasite infection and parasitological analysis. Adult Ascaris suum worms were harvested from pigs at a Brazilian slaughterhouse (Belo Horizonte). Parasite eggs were isolated from female uteri through gentle mechanical maceration, purified by straining, and cultured to embryonation in 0.2 M  $H_2SO_4$  as described by Boes et al. (80). On day 100 of culture, which is the peak of larva infectivity (81), the fully embryonated eggs were used for experimental infections. Mice were inoculated through the intragastric route with 2500 fully embryonated eggs. The parasite burden was evaluated based on the total number of larvae recovered after 5 days in the liver, after 5 and 8 days in the lungs, and after 8 days in the BAL fluid postinfection. The tissues were collected, sliced finely with scissors, and placed in a Baermann apparatus for 4 hours in PBS buffer at 37°C. The larvae were recovered, fixed with formalin, and counted by microscopy.

*Flow cytometry immunophenotyping.* Lungs were digested with collagenase IV (Sigma-Aldrich) in RPMI culture media for 60 minutes at 37°C, and single-cell suspensions were made by filtering through 100- $\mu$ m gauze (BD Biosciences). After addition of Percoll 40% and further ACK buffer for red blood cell lysis, leukocyte suspensions were counted (Countess II, Life Technologies) and incubated for 3 hours at 37°C, 5% CO<sub>2</sub>, in the absence of (media) or upon stimulation with 0.5/0.05 nM of PMA/ionomycin (Sigma-Aldrich) and 10  $\mu$ g/mL of brefeldin A. Cells

were then stained using 2 panels of antibodies for flow cytometry (Table 1): CD4+ T cell polarization and innate lymphoid cell (ILC) subset panel, and myeloid lineage panel. Briefly, staining using the T cell polarization panel required 3 steps: The cells were stained with a LIVE/ DEAD marker (Fixable Blue, UV450, Thermo Fisher Scientific) and the biotinylated antibodies for lineage exclusion: TER119, CD19, γδTCR, Gr1, CD11c, CD49b, CD11b. Secondly, after washing with FACS buffer (PBS, 2% FBS), antibodies against the extracellular markers CD45, TCR-β, Thy1.2, and streptavidin were used. After washing, cells were then fixed with 2% paraformaldehyde and permeabilized (Perm buffer, eBioscience), and finally stained with the following antibodies: CD4, IL-5, IL-13, IL-17A, and IFN-y. The myeloid lineage panel was performed using a 2-step staining protocol: cells were stained with a LIVE/DEAD marker (Fixable Blue, UV450, Thermo Fisher Scientific) and the biotinylated antibodies for lineage exclusion: TCR-β, γδTCR, CD49b, CD19. Secondly, after washing with FACS buffer, CD11c, Ly6C, CD11b, MHC-II, Siglec-F, Ly6G, and F4/80 were used. After washing, cells were then fixed with 2% paraformaldehyde. CD4+T cell depletion was evaluated in the peripheral whole blood as well as in the lung tissue by flow cytometry analysis using TCR-β, CD4, and CD8 surface staining. A description of the antibodies used, including the source and catalog/clone numbers, is included in Table 1. Data were acquired on an LSR Fortessa flow cytometer (BD Biosciences) and analyzed with FlowJo software (Tree Star).

Lung tissue chemokine, cytokine, cytokine receptor, and eosinophil peroxidase measurement. To evaluate the inflammatory mediators in the lung tissue homogenate, the bottom lobe of the right lung was homogenized for 2 cycles of 20 seconds at 5000 rpm each using Precellys tubes with metal beads in 500 µL of RIPA lysis buffer supplemented with protease inhibitor cocktail (ChemCruz) and 200 nM PMSF. Following centrifugation at 8000 g for 10 minutes at 4°C, the supernatant was used to measure cytokines in the lung's homogenate. The cytokines IL-1β, IL-3, IL-4, IL-5, IL-6, IL-13, IL-10, IL-33, IFN- $\gamma$ , and TNF- $\alpha$  as well as the chemokines CCL-2 (MCP-1), CCL-5 (RANTES), CCL-11 (eotaxin), CXCL-5 (LIX), and CXCL-10 (IP-10) were measured using a commercial kit (Millipore). Lung IL-13Ra2 protein concentration was determined by ELISA as previously described. High-protein-binding 96-well plates were coated with anti-IL-13Rα2 (1 µg/mL) (R&D Systems) in PBS overnight, and a biotinylated antimouse IL-13 (2 µg/mL; Centocor) was used for detection. Mouse eosinophil peroxidase (EPO) levels were measured in the lung homogenates using a sandwich ELISA kit (LSBio).

*Histological examinations*. The lobes of the left lung were fixed in 10% buffered formalin (pH 7.2) and embedded in paraffin. Sections (5 mm thick) were mounted on glass slides and stained with H&E for inflammatory infiltration analysis, periodic acid–Schiff (PAS) for mucus production, or Masson's trichrome or Picrosirius for quantification of fibrotic process and collagen deposition. All images of the tissue sections were scanned using Scan Aperio (Leica), and the analyses were performed with specific algorithms for score quantification using Aperio e-Slide Manager (Leica).

Morphometric analysis on the Ascaris larval stages. The development of different Ascaris larval stages was evaluated in BALB/c and  $\Delta$ dblGATA preallergic and nonallergic mice, by morphometric measurement of the length and width of larvae recovered from the lung tissue at 5 dpi and 8 dpi as well as in the BAL at 8 dpi. After individual counting, larvae recovered from HDM<sup>+</sup>Ascaris<sup>-</sup> and HDM<sup>+</sup>Ascaris<sup>+</sup> mice were pooled in separate wells, stretched by heat-killing, and fixed in 10% buffered formalin. In vitro egg-hatched L3 larvae were also evaluated as previously described (82). Images of Ascaris larval stages were collected using a high-resolution bright-field confocal SP8 microscope (Leica). The morphometric analysis was performed using Imaris 9.0 software (Biplane).

RNA purification and quantitative PCR gene expression analysis. Lung tissue was homogenized in Trizol (Life Technologies) in a Precellys 24 homogenizer (Bertin Technologies). Total RNA was extracted with chloroform using a MagMax-96 Total RNA Isolation Kit (Qiagen) and reverse-transcribed to cDNA using SuperScript II Reverse Transcriptase (Life Technologies). Gene expression was quantified using Power SYBR Green PCR Master Mix (Applied Biosystems) by reverse transcriptase PCR on an ABI Prism 7900HT Sequence Detection System (Applied Biosystems). Expression of the target genes was described relative to RPLP2 mRNA levels. The specific primer pairs for each gene were: arginase-1, 5'-GGAAAGCCAATGAAGAGCTG-3' and 5'-GCTTCCAACTGCCAGACTGT-3'; Fizz-1, 5'-CCCTCCACT-GTAACGAAGACTC-3' and 5'-CACACCCAGTAGCAGTCATCC-3'; Nos2, 5'-TGCCCCTTCAATGGTTGGTA-3' and 5'-ACTGGAGGGAC-CAGCCAAAT-3'; Rplp2, 5'-TACGTCGCCTCTTACCTGCT-3' and 5'-GACCTTGTTGAGCCGATCAT-3'; Muc5ac, 5'-CAGGACTCTCT-GAAATCGTACCA-3' and 5'-AAGGCTCGTACCACAGGGA-3'.

In vivo  $CD4^+$  T cell depletion. Two hundred microliters of sterile PBS with 500 µg of anti-CD4 (GK1.5) or 500 µg of rat IgG2b isotype control (both from BioXCell) was given i.p. using 2 different protocols. The first protocol, "group day 0," received the i.p. injections on day 0 and days 7, 14, and 21 after HDM sensitization. The second protocol, "group day 18," received the injections at days 18 and 25 after the HDM sensitization (Figure 6A). The efficacy of the in vivo CD4<sup>+</sup> T cell depletion was evaluated in the peripheral whole blood as well as in the lung tissue by flow cytometry analysis using TCR- $\beta$  and CD4 surface staining (Supplemental Figure 4).

HDM-specific antibody response. Antibody responses to HDM were evaluated by ELISA. Immunolon 4HBX plates were coated with 5  $\mu$ g/ well of HDM extract (Greer Laboratories). Plates were incubated overnight at 4°C and, after washing, blocked with 200  $\mu$ L of 5% PBS-BSA for 1 hour at room temperature. Plates were washed 6 times with Tween-PBS. Fifty microliters of mouse serum (1:500 for IgG1 and 1:1 for IgE in 1% PBS-BSA) was added in duplicate and incubated for 1 hour at room temperature. Plates were washed and then incubated separately with biotinylated rat anti-mouse IgG1, IgE, and IgA (BD Pharmingen) (1:250 in 1% PBS-BSA) for 1 hour at room temperature. Plates were washed and incubated with 50  $\mu$ L of streptavidin conjugated with HRP (1:200 in 1% PBS-BSA for IgG1 and 1:100 in 1% PBS-BSA for IgE) for 30 minutes at room temperature. After washing, TMB substrate TMB (3,3',5,5'-tetramethylbenzidine) was added (50  $\mu$ L/well) and incubated for 5 minutes in the dark for IgG1 and 15 minutes for IgE. Twenty-five microliters of 2N H<sub>2</sub>SO<sub>4</sub> was used to stop the reaction, and plates were read at absorbances of 450 nm and 550 nm on a multiplate reader (Molecular Devices).

Statistics. Unless stated otherwise, geometric means and 95% confidence intervals were used as the measure of central tendency. The Kolmogorov-Smirnov test was used to determine whether data followed a normal distribution pattern. For nonparametric data, the Kruskal-Wallis test followed by Dunn's multiple-comparisons test was used for the analysis of tissue cytokine and chemokine profiles, the frequencies of ILCs and myeloid and T cells, and the comparison of parasite burden and larval development between different mouse strains or antibody treatments. The Mann-Whitney U test was used to determine the differences between the variables associated with parasite burden in BALB/c mice and \(\Delta\)dblGATA separately. For the differences in gene expression, the nonparametric Wilcoxon matched-pairs test or the Mann-Whitney U test was used for the analysis between HDM+Ascaris- mice and HDM+Ascaris+ mice. Correlation analysis was performed using the Spearman rank test. Graph generation and statistical analysis were performed using Prism 7.0 for Windows (GraphPad Software Inc.), and differences were considered statistically significant when P was less than 0.05.

*Study approval.* The maintenance and use of animals were in accordance with the recommendations of the NIH *Guide for the Care and Use of Laboratory Animals* (National Academies Press, 2011). All animal experiments were approved by NIH Animal Care and Use Committee (Animal Study Protocol LPD-6). All efforts were made to minimize suffering and pain, as described in these approved protocols and guidelines.

#### Author contributions

PHGG, RQP, AR, SBA, JS, and EPK performed the experiments. PHGG, RQP, and TBN designed the experiments. PHGG, RQP, AR, SBA, JS, EPK, RTF, and TBN carried out analyses. PHGG wrote the paper; all authors edited the paper. RTF and TBN provided funding and reagents.

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