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# A human model of asthma exacerbation reveals transcriptional programs and cell circuits specific to allergic asthma

Jehan Alladina<sup>1,2,†</sup>, Neal P. Smith<sup>2,3,4,†</sup>, Tristan Kooistra<sup>1,2</sup>, Kamil Slowikowski<sup>2,3,4</sup>, Isabela J. Kernin<sup>2,3,4</sup>, Jacques Deguine<sup>3</sup>, Henry L. Keen<sup>5</sup>, Kasidet Manakongtreecheep<sup>2,3,4</sup>, Jessica Tantivit<sup>2,3,4</sup>, Rod A. Rahimi<sup>1,2</sup>, Susan L. Sheng<sup>1</sup>, Nhan D. Nguyen<sup>1,2</sup>, Alexis M. Haring<sup>1,2</sup>, Francesca L. Giacona<sup>1,2</sup>, Lida P. Hariri<sup>1,6</sup>, Ramnik J. Xavier<sup>3,7,8</sup>, Andrew D. Luster<sup>2,3,9</sup>, Alexandra-Chloé Villani<sup>2,3,4,‡,\*</sup>, Josalyn L. Cho<sup>10,‡,\*</sup>, Benjamin D. Medoff<sup>1,2,‡</sup>

<sup>1</sup>Division of Pulmonary and Critical Care Medicine, Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA.

<sup>2</sup>Center for Immunology and Inflammatory Diseases, Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA.

<sup>3</sup>Broad Institute of Massachusetts Institute of Technology and Harvard, Cambridge, MA, USA.

<sup>4</sup>Massachusetts General Hospital Cancer Center, Boston, MA, USA.

<sup>5</sup>Iowa Institute of Human Genetics, University of Iowa Carver College of Medicine, Iowa City, IA, USA.

<sup>6</sup>Department of Pathology, Massachusetts General Hospital, Boston, MA, USA.

<sup>7</sup>Center for Computational and Integrative Biology, Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA.

<sup>8</sup>Division of Gastroenterology, Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA.

<sup>9</sup>Division of Rheumatology, Allergy and Immunology, Massachusetts General Hospital and Harvard Medical School, Boston, MA, USA.

<sup>10</sup>Division of Pulmonary, Critical Care and Occupational Medicine, University of Iowa Carver College of Medicine, Iowa City, IA, USA.

# Abstract

Corresponding author. avillani@mgh.harvard.edu (A.-C.V.); josalyn-cho@uiowa.edu (J.L.C.).

<sup>&</sup>lt;sup>†</sup>These authors contributed equally to this work.

<sup>&</sup>lt;sup>‡</sup>These authors contributed equally to this work.

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Asthma is a chronic disease most commonly associated with allergy and type 2 inflammation. However, the mechanisms that link airway inflammation to the structural changes that define asthma are incompletely understood. Using a human model of allergen-induced asthma exacerbation, we compared the lower airway mucosa in allergic asthmatics and allergic nonasthmatic controls using single-cell RNA sequencing. In response to allergen, the asthmatic airway epithelium was highly dynamic and up-regulated genes involved in matrix degradation, mucus metaplasia, and glycolysis while failing to induce injury-repair and antioxidant pathways observed in controls. IL9-expressing pathogenic TH2 cells were specific to asthmatic airways and were only observed after allergen challenge. Additionally, conventional type 2 dendritic cells (DC2 that express CD1C) and CCR2-expressing monocyte-derived cells (MCs) were uniquely enriched in asthmatics after allergen, with up-regulation of genes that sustain type 2 inflammation and promote pathologic airway remodeling. In contrast, allergic controls were enriched for macrophage-like MCs that up-regulated tissue repair programs after allergen challenge, suggesting that these populations may protect against asthmatic airway remodeling. Cellular interaction analyses revealed a T<sub>H</sub>2-mononuclear phagocyte-basal cell interactome unique to asthmatics. These pathogenic cellular circuits were characterized by type 2 programming of immune and structural cells and additional pathways that may sustain and amplify type 2 signals, including TNF family signaling, altered cellular metabolism, failure to engage antioxidant responses, and loss of growth factor signaling. Our findings therefore suggest that pathogenic effector circuits and the absence of proresolution programs drive structural airway disease in response to type 2 inflammation.

### INTRODUCTION

Asthma affects 1 in 13 people in the United States and is increasing in prevalence (1). Most asthma is characterized by allergic inflammation to environmental antigens (2). However, not all allergic individuals sensitized to inhaled allergens develop asthma, a disease of the lower airways defined by airway inflammation, bronchial hyper-responsiveness, and mucus hypersecretion (2, 3). We therefore hypothesized that identifying key differences in the lower airway response to allergen between allergic asthmatics (AAs) and allergic individuals without asthma would provide fundamental insights into the mechanisms that drive asthma.

It is increasingly recognized that cellular cross-talk within the airway microenvironment is important in asthma pathogenesis (2–4). Consistent with this, the airway mucosa in asthma is characterized by innate and adaptive type 2 immune cells, including eosinophils, mast cells, and T helper 2 ( $T_H2$ ) cells, that localize in proximity to the airway epithelium. Moreover, murine and human studies have established that understanding epithelial cell activation and immune cell enrichment in the airway mucosa in response to allergen challenge is essential to elucidate asthma pathobiology (3–11). Single-cell transcriptional approaches have facilitated the discovery of tissue-specific cell subsets associated with allergic disease, including a recent steady-state analysis of human airways that showed that pathogenic effector  $T_H2$  cells, mast cells, and secretory airway epithelial cells (AECs) are enriched in asthmatics compared with healthy controls (12–16). However, high-dimensional analyses of the dynamic lower airway mucosal response to allergen challenge have not been reported.

Bronchoscopic segmental allergen challenge (SAC) is a powerful research tool that recapitulates the features of an allergen-induced asthma exacerbation in a single airway segment and has been used to identify asthma-relevant pathways and successfully predict the clinical efficacy of new therapeutics in asthma (9, 17, 18). We have previously shown that SAC induces type 2 airway inflammation in both AAs and allergic non-asthmatic controls (ACs) (9). However, asthmatics have increased measures of type 2 inflammation and uniquely demonstrate airway structural changes, including increases in mucin production, epithelial and subepithelial thickness, and airway smooth muscle thickness (9, 19). Thus, we hypothesize that asthmatics have distinct cellular and transcriptional programs in the airway microenvironment that amplify type 2 inflammation and link mucosal inflammation to the airway structural changes that define asthma.

We leveraged bronchoscopic SAC and single-cell RNA sequencing (scRNA-seq) to define the airway mucosal microenvironment and identified transcriptional programs that were only observed after allergen exposure. By comparing AAs with ACs, we distinguished cellular and molecular pathways that define asthma from those associated with allergy alone. These findings included a robust transcriptional response to allergen challenge in basal and secretory AECs from asthmatics that was greater than in ACs and characterized by up-regulation of genes involved in mucus metaplasia and matrix remodeling. In addition, an AEC glycolytic gene signature was associated with asthma, whereas antioxidant and growth factor signaling pathways were identified as potentially protective. *IL9*-expressing pathogenic  $T_H^2$  cells were only found in asthmatic airways after SAC, along with enrichment of conventional type 2 dendritic cells (DC2) and pathogenic monocyte-derived cells (MCs) that up-regulated inflammatory mediators and metalloproteinases. Cell-cell interaction analyses highlighted immune-epithelial interactions specific to asthma and were dominated by a  $T_H^2$ -mononuclear phagocyte (MNP)–basal cell interactome, suggesting that cross-talk between these key cell subsets is critical to the pathogenesis of allergic asthma.

# RESULTS

# A human model of allergen-induced asthma exacerbation transformed the airway mucosal cellular landscape

To test the hypothesis that differences in immune or structural cells in the airway mucosa distinguish asthma from allergy alone, we enrolled 61 adults with allergy to house dust mite (HDM) or cat with or without asthma, as well as 5 nonallergic healthy controls, to undergo SAC (Fig. 1A, data file S1, and Supplementary Methods) (9, 19). Allergy was defined as a clinical history of allergic rhinitis and/or conjunctivitis to HDM or cat, which was confirmed by skin prick testing (SPT). Quantitative SPT was used to standardize the dose of allergen administered according to each participant's level of allergy (fig. S1A and Supplementary Methods). ACs had no history of asthma, normal forced expiratory volume in 1 s (FEV<sub>1</sub>) and forced vital capacity (FVC), and a negative methacholine challenge test. AAs had a history of mild asthma, FEV<sub>1</sub> 75% predicted, and bronchial hyperresponsiveness confirmed by positive methacholine challenge testing. Bronchoalveolar lavage (BAL) and endobronchial brush samples of third- to fourth-generation airway segments were collected at baseline and 24 hours after administration of allergen and diluent, which controlled for

the effects of the bronchoscopy procedure (Fig. 1B). Similar to prior reports (9), SAC induced robust eosinophilic inflammation in both allergic groups that was restricted to the allergen-challenged segment and was not observed in healthy controls (Fig. 1C).

To characterize the lower airway mucosa during allergic inflammation, we performed flow cytometry of endobronchial brush samples and found enrichment of CD45<sup>+</sup> immune cells after SAC, including CD4<sup>+</sup> T cells, CD19<sup>+</sup> B cells, and HLA-DR<sup>+</sup> antigen-presenting cells (Fig. 1D). Given the limited number of cells recovered with this approach and difficulty identifying specific cell subsets due to overlapping expression of cell surface markers, we leveraged scRNA-seq to define cellular populations in the lower airway mucosa using an unbiased analytical framework. We generated high-quality scRNA-seq data from 21 lower airway brushings (totaling 52,152 cells) collected from four AAs (25,397 cells) and four ACs (26,755 cells) (Fig. 1E, fig. S1, B and C, and data files S2 and S3). Brushings were collected at baseline (Bln; 22,406 cells) and 24 hours after exposure to allergen (Ag; 17,462 cells) or diluent (Dil; 12,284 cells). Diluent samples could not be collected in two AAs and one AC and were thus excluded from downstream statistical comparisons with the other experimental conditions. We systematically performed iterative cell clustering to define distinct global and lineage-specific cellular populations in the lower airway mucosa. We subsequently used differential abundance, differential gene expression (DGE), and cell-cell interaction analyses to identify populations, transcriptional programs, and cellular networks that distinguish asthma from allergy alone (Fig. 1E and Materials and Methods).

Principal components analysis revealed variability driven by both disease group and experimental condition (fig. S1D). After data integration and unsupervised clustering analysis (Materials and Methods), we identified seven global cell lineages, including AECs (EPCAM), CD4 T cells (CD3D, CD4, and IL7R), CD8 T cells (CD3D and CD8A), MNPs (LYZ), B cells (MS4A1), mast cells (CPA3), and natural killer cells (GNLY) (Fig. 2, A to C, fig. S1E, and data file S4). All cell lineages were captured across all participants and experimental conditions (fig. S1E). At baseline, the airway mucosal landscape was dominated by AECs and CD8 T cells in both groups (Fig. 2D), with a similar cellular composition observed after diluent administration (fig. S1, F and G). Mixed-effects logistic regression was used to identify differences in cellular abundance that were associated with each disease group [ACs: odds ratio (OR) < 1, AAs: OR > 1; Fig. 2E, fig. S1H, and Materials and Methods] (20). Both B cells (OR 4.48, 95% confidence interval [1.73 to 11.61) and mast cells (OR 6.68, [3.20 to 13.94]) were associated with asthma at baseline, and mast cells expressed genes (CD38 and KIT) consistent with a subset that proliferates in situ during allergic disease (fig. S1I and data file S5) (13). We did not observe DGE [log<sub>2</sub>FC (fold change) > 0.5 and false discovery rate (FDR) < 0.1; Materials and Methods] in B cells or mast cells between groups, but mast cells in AAs responded to allergen challenge by up-regulating genes involved in mast cell growth and survival (IL3RA, IL2RA, BIRC3, and BCL2), chemotaxis (CXCR4 and C5AR1), and KIT expression (RUNX1), suggesting that these pathways may promote mast cell enrichment in asthmatic airways (fig. S1, J and K, and data file S6). After allergen challenge, immune cells dominated airway mucosal samples (Bln: 9658 cells, 43.1%; Ag: 13,459 cells, 77.1%) (Fig. 2F). The number of MNPs increased the most after allergen challenge in both groups (ACs: 6.7% versus 33.4%, P = 0.009; AAs:

5.9% versus 28.1%, P = 0.02), whereas CD4 T cells only increased in asthmatics (7.1% versus 20.6%, P = 0.03).

# Basal and secretory epithelial cells in asthmatics extensively altered their transcriptional profiles in response to allergen

Our prior work suggests that AECs in asthmatics are uniquely sensitive to type 2 inflammation (9). Subclustering of the 20,410 AECs enabled detailed characterization of the transcriptional programs characterizing AECs at baseline and in the setting of allergic inflammation. We identified 14 distinct AEC subsets [area under the curve (AUC) 0.75 and pseudo-bulk DGE FDR < 0.05; Materials and Methods], annotated by cross-referencing top marker genes with published transcriptional data (Fig. 3, A to C, fig. S2, A and B, and data file S4) (4, 16, 21–25). Three ciliated clusters (FOXJI and PIFO) were observed, including mucous-ciliated cells (SCGB1A1 and MUC5AC). A prior study identified mucous-ciliated cells in asthmatics but not healthy controls (16). However, we found similar numbers of mucous-ciliated cells in both asthmatics and ACs, indicating that these cells do not distinguish asthma from allergy alone. Four basal cell clusters (KRT5 and S100A2) were also identified. Suprabasal cells were distinguished from basal cells by lower TP63 and KRT5 expression, cycling basal cells expressed markers of proliferation (PCLAF and *MKI67*), and a cluster that was distinguished primarily by lower overall gene expression was termed quiescent (quiesBasal). Three secretory clusters had high expression of secretoglobin genes (SCGB1A1 and SCGB3A1). Goblet and quiescent goblet (quiesGoblet) cells had high expression of the tethered mucin genes (MUC5AC and MUC5B), whereas club cells were distinguished by lower expression of mucin and goblet cell differentiation genes (FOXA3 and SPDEF) (4, 16, 21, 23, 26). A small serous cluster expressed markers of both serous (LYZ, PRB3) and mucous cells (AZGP1, BPIFB2, and MUC5B), suggesting sampling of submucosal glands. Moreover, we identified the recently described deuterosomal (CCNO, CDC20B), hillock (KRT13 and SPRR3), and ionocyte (FOX11 and CFTR) populations (Fig. 3, A and B, and fig. S2C) (4, 16, 21–25). We observed AEC subset-specific expression of asthma-relevant mediators (Fig. 3C). Basal cells were enriched for IL33 and uniquely expressed TSLP. Periostin (POSTN) was expressed in both basal and ionocyte subsets, and ionocytes also expressed genes involved in prostaglandin signaling (PTGS2 and PTGER3).

Hillock cells expressed genes involved in squamous differentiation (*KRT13* and *KRT4*), barrier integrity (*SPRR3* and *ECM1*), and immune responses (*IL1A*, *S100A8*, and *ALOX15*) (Fig. 3, B and C, and fig. S2C). Hillock cells have been described as a highly proliferative transitional population between basal and secretory cells (4, 23, 25). In our dataset, hillock cells did not express proliferation markers, nor did they cluster between basal and secretory cells in Uniform Manifold Approximation and Projection (UMAP) space. However, we identified a discrete region within the club cell cluster that expressed hillock cell markers and aligned with transitional "hillock club" cells previously only described in mouse trachea (fig. S2, C and D, and data file S7) (23). In humans, hillock cells have only been reported in the nasal mucosa (25). To validate our scRNA-seq findings, we performed immunofluorescence staining of explanted asthmatic lung and observed contiguous groups of KRT13<sup>+</sup> cells lining the airway without overlying ciliated cells (fig. S2E), consistent with prior descriptions of hillock cells (23, 25). In atopic dermatitis and eosinophilic esophagitis,

keratinized squamous cells can sense and potentiate type 2 inflammation and induce barrier dysfunction (27). Consistent with this, hillock cells in asthmatics expressed interleukin-13 (IL-13)–responsive genes, including calpain 14 (*CAPN14*), which was highly enriched in this subset, has been linked to asthma, and induces epithelial barrier dysfunction (fig. S2F) (28, 29). These findings suggest that hillock cells may play a previously unappreciated role in directing the lower airway response to allergen.

Although quantification of AEC subsets did not reveal significant differences between groups (fig. S2G and data file S3), mixed-effects logistic regression identified an association between ACs and quiesGoblet (OR 0.39, [0.50 to 1.51]) at baseline and deuterosomal cells (OR 0.35, [0.17 to 0.75]) after allergen challenge (fig. S2, H and I). Ciliated cells (OR 2.10, [1.25 to 3.54]) and ionocytes (OR 2.59, [1.34 to 5.02]) were associated with asthmatics at baseline. To assess the response of AECs to allergen challenge, we quantified the number of differentially expressed genes (DEGs;  $log_2FC > 0.5$  and FDR < 0.1; Materials and Methods) after allergen compared with baseline (Fig. 3D and data file S6). AECs in asthmatics markedly altered their transcriptional profile after allergen challenge compared with ACs (total DEGs = 4286 in AAs versus 269 in ACs). Goblet cells in asthmatics had the greatest transcriptional response to allergen challenge (DEGs = 1454), followed by suprabasal (DEGs = 850), quiesGoblet (DEGs = 645), club (DEGs = 472), and basal cells (DEGs = 323), suggesting that these AEC subsets play a central role during allergic inflammation. At baseline, we found minimal transcriptional differences between groups (data file S6). Given the large number of DEGs induced by allergen challenge, we created an interaction term for disease group and experimental condition to identify DEGs uniquely associated with asthma after allergen challenge (FDR < 0.1; Materials and Methods and data file S6). Using the interaction term, we identified 65 shared DEGs across basal and secretory clusters specific to asthma (Fig. 3E). This core response was characterized by up-regulation of IL-13-induced genes, along with genes involved in inflammatory signaling, tissue remodeling, glycolysis, and down-regulation of antioxidant genes.

We also identified subset-specific DEGs between asthmatics and ACs, highlighting new aspects of AEC biology associated with asthma. Basal and suprabasal cells in allergic controls were characterized by an injury-repair response after allergen challenge, with increased expression of alarmins (*IL33* and *HMGB1*) and neutrophil chemoattractants (*CXCL6*) (Fig. 3F). In contrast, these cells in asthmatics up-regulated genes involved in type 2 inflammatory cell recruitment and signaling (*CCL26* and *IL13RA1*), mucus metaplasia (*SPDEF*), and genes that promote extracellular matrix (ECM) degradation (*MMP1* and *MMP7*) and allow for connective tissue regeneration and biogenesis (*POSTN* and *LOXL4*). We then performed Ingenuity Pathway Analysis (IPA) of DEGs identified by the interaction term to identify potential upstream regulators of transcriptional changes in suprabasal cells in each group, with |*z* score| > 2 considered significant (Materials and Methods and data file S8). In asthmatics, the strongest predicted upstream regulators included IL-13 and fibroblast growth factor 2 (FGF2) (|*z* scores| = 2.45), both involved in airway inflammation and remodeling (fig. S2J).

ACs also up-regulated stress response genes (*S100A8* and *HMGB2*) in goblet cells, along with genes that orchestrate tissue integrity and angiogenesis (*VEGFA*) and antioxidant

defense (*GSTA1*, *GSTA2*, and *GSTA4*) (Fig. 3F). Although all secretory clusters in asthmatics up-regulated *SPDEF*, a signal transducer and activator of transcription 6 (STAT6)–induced transcription factor that directs goblet cell differentiation (26), goblet cells in asthmatics additionally up-regulated genes that control mucin glycosylation and hydration (*GCNT4* and *GALNT10*) and regulate airway surface liquid pH (*CA2* and *SLC26A4*) (30). In asthmatics, the strongest predicted upstream regulator of goblet cells was the epidermal growth factor receptor (EGFR; |z score| = 2.0; fig. S2J), which can promote goblet cell hyperplasia and mucin production (21, 31, 32). Collectively, these results identify airway basal and secretory cells as highly dynamic cells during allergic inflammation and reveal mechanisms by which they may drive asthma pathogenesis.

# *IL9*-expressing pathogenic T<sub>H</sub>2 cells are specific to asthmatic airways after allergen challenge

 $T_{H2}$  cells play a central role in the inflammatory response to allergen in the airways (2, 6, 9). Subclustering of 18,714 T cells identified 10 distinct subsets annotated by cross-referencing top marker genes (AUC 0.75 and pseudo-bulk DGE FDR < 0.05; Materials and Methods and data file S4) with published transcriptional data (16, 33–39) and included four CD8 T cell subsets, five CD4 T cell subsets, and a small  $\gamma\delta$  T cell (TRDC) population (Fig. 4, A and B, and fig. S3, A and B). Quiescent CD8 T cells were distinguished by low overall gene expression. CD8 T (GZMK) cells were enriched for late effector genes (EOMES and KLRG1), whereas CD8 T (CLIC3) and CD8 T (EGR2) cells were both enriched for tissue-resident memory markers (ITGAE, ITGA1, and CD69). These findings were supported by gene set enrichment analysis (GSEA) (fig. S3C, Materials and Methods, and data file S7) (36). CD4 T cell subsets included CD4 regulatory T (T<sub>reg</sub>) (FOXP3), CD4 T<sub>H</sub>2 (GATA3), CD4 T (CD40LG), CD4 T<sub>H</sub>17 (RORA), and a recently described interferon (IFN)-responsive CD4 T<sub>H</sub>IFNR (ISG15) population (33, 37, 38). The identities of T<sub>H</sub>2, T<sub>H</sub>17, and T<sub>H</sub>IFNR clusters were confirmed by GSEA (Fig. 4C, fig. S3C, and data file S7) (33, 39). CD8 T cells represented the highest proportion of T cells in both groups at baseline and remained the largest proportion of T cells in ACs after allergen challenge (Fig. 4D, fig. S3D, and data file S3). In contrast, the proportion of CD4 T (*CD40LG*),  $T_{reg}$ , and  $T_{H2}$ cells increased in asthmatics after allergen challenge. TH2 abundance was associated with asthma both at baseline (OR 3.57, [1.71 to 7.45]) and after allergen challenge (OR 3.43, [1.87 to 6.29]) (Fig. 4E, fig. S3E, and data file S5). CD4 T (*CD40LG*) (OR 2.12, [1.23 to 3.68]) and Treg (OR 3.47, [2.20 to 5.47]) were also associated with asthma after allergen challenge. Further interrogation of the  $T_H^2$  population identified a subset of cells expressing genes associated with enhanced pathogenic function (PTGDR2 and HPGDS) and receptors for T<sub>H</sub>2-polarizing cytokines (IL1RL1 and IL17RB) (Fig. 4F and fig. S3F) (33-35).

To assess the response of T cells to allergen challenge, we quantified the number of DEGs  $(\log_2 FC > 0.5 \text{ and FDR} < 0.1; \text{Materials and Methods})$  after allergen compared with baseline and found the greatest number of DEGs in  $T_H2$  (DEGs = 76) and CD4 T (*CD40LG*) (DEGs = 54) in asthmatics (fig. S3G and data file S6). There were few DEGs identified between groups after allergen challenge (fig. S3, H and I, and data file S6). *IL9* was the most up-regulated gene in asthmatic  $T_H2$  cells in response to allergen (logFC 8.3, FDR =  $2.8 \times 10^{-8}$ ). *IL9*-expressing pathogenic  $T_H2$  cells have previously been described in

the blood of AAs (33), but they have not been reported in the airway. In our model, *IL9*expressing  $T_H^2$  cells were only observed after allergen challenge and were highly specific to asthmatic airways (Fig. 4G and fig. S3J). These cells were also enriched for inducers of IL-9 production (*PPARG* and *IRF4*) (fig. S3F) (33, 40, 41). Moreover, IL-9 protein levels in BAL samples increased in response to allergen challenge and were higher in asthmatics compared with ACs (median [interquartile range, IQR]: 489.9 [429.9 to 583.5] versus 33.5 [10.2 to 238.9] pg/ml, *P*< 0.0001) (Fig. 4H). IL-9 directly promotes mast cell activation and expression of profibrotic mediators, induces autocrine production of type 2 cytokines by  $T_H^2$  cells, and can indirectly increase AEC mucin production via induction of IL-13 (40, 42–44). *IL9R* expression was identified in mast cells and pathogenic  $T_H^2$ , similar to previous reports (figs. S1I and S3F) (16, 45). These results demonstrate that pathogenic  $T_H^2$ cells in the airway, and in particular their expression of *IL9*, distinguish asthma from allergy alone. Furthermore, they suggest that allergen-induced IL-9 expression may amplify type 2 inflammation and promote pathologic airway remodeling.

# DC2 and *CCR2*-expressing MCs are enriched in asthmatics after allergen challenge, whereas macrophage-like MCs are enriched in ACs

MNPs are necessary to sustain type 2 inflammation in murine models of asthma (46); however, the precise role of MNP subsets and their functional specialization in the context of human allergic airway inflammation are incompletely understood. MNPs are recruited to the lung during allergic inflammation (Figs. 1D and 2, D and F) and include dendritic cells (DCs), macrophages (Macs), and MCs (47-49). Subclustering of 8510 MNP cells identified 14 distinct subsets that were annotated by cross-referencing top marker genes (AUC 0.75 and pseudo-bulk DGE FDR < 0.05; Materials and Methods and data file S4) with published transcriptional data (Fig. 5, A to C, and fig. S4, A and B) (49-58). DC subsets included DC1 (CLEC9A), DC2 (CD1C), migratory DCs (migDCs; CCR7), plasmacytoid DCs (pDCs; TCF4), and Axl-Siglec6 DCs (AXL). Three clusters were annotated as Macs on the basis of their abundance in baseline samples, indicative of their tissue residency. Mac1 (FABP4) and quiesMacs (defined by overall lower gene expression) were enriched for canonical Macs (MARCO, MSR1, and VSIG4) and lipid metabolism genes (FABP4 and APOE), consistent with previously described luminal Macs (16, 25, 51). Mac2 (A2M) had high expression of complement family genes (CIQA, CIQB and CIQC) and lower relative expression of canonical Mac markers, reminiscent of airway-associated Macs (AAMs) recently characterized in mice (57, 58).

Five clusters were annotated as MCs on the basis of high expression of canonical monocyte genes (*CD14*, *VCAN*, and *FCN1*) and varying expression of Mac and DC markers, consistent with the well-described plasticity of MCs (47, 48, 59). Quiescent MCs were characterized by markers of monocyte plasticity and had lower overall gene expression levels than the remaining MC clusters. MC1 expressed high levels of IFN-responsive genes (*GBP1*, *STAT1*, and *WARS*) and uniquely expressed CXCR3 ligands (*CXCL9*, *CXCL10*, and *CXCL11*) (Fig. 5C and fig. S4C). MC2 were enriched for matrix-interacting genes (*SPP1*, *MERTK*, and *LGMN*), consistent with a previously described lung resident population (50–52). MC3 distinctively expressed genes involved in tissue repair (*AREG*, *VEGFA*, and *HBEGF*) and *NR4A1*, a transcription factor necessary for monocytes to

restrict inflammation and differentiate into regenerative Macs (60, 61). MC4, in contrast, had low expression of Mac markers, had high expression of genes associated with undifferentiated monocytes (*TMEM176B*, *AIF1*, and *CSF1R*), and were enriched for the monocyte chemotactic receptor *CCR2*, consistent with a recently recruited immature MC.

Tissue MCs and Macs play an important role in lung homeostasis, and imbalance in their proportions has been associated with pulmonary disease (50, 62). Mac1 (*FABP4*) was abundant in both groups at baseline, whereas DC subsets were rare (Fig. 5D, fig. S4D, and data file S3). Using mixed effects logistic regression, MC2 (*SPP1*) abundance was associated with ACs at baseline (OR 0.52, [0.31 to 0.89]), whereas quiescent MCs were associated with asthmatics (OR 2.05, [1.59 to 2.66]); Fig. 5, D and E, fig. S4E, and data file S5). Allergen challenge induced a dramatic shift in the MNP profile that was distinct between ACs and asthmatics. The abundance of Mac1 and Mac-like MCs (MC1, MC2, and MC3) was associated with ACs after allergen challenge, whereas the abundance of MC4 (OR 2.77, [1.29 to 5.92]) and DC2 (OR 2.11, [1.23 to 3.61]) was associated with asthmatics. Flow cytometry of endobronchial brush samples confirmed the enrichment of DC2 (CD45<sup>+</sup>HLA-DR<sup>hi</sup>CD1c<sup>+</sup>) in asthmatics compared with ACs after SAC (6.6% [2.6 to 9.2%] versus 2.9% [2.4 to 3.2%], P = 0.02; fig. S4F).

To identify genes associated with enrichment of MNP subsets in each group after allergen challenge, we used least absolute shrinkage and selection operation (LASSO) modeling (empirical P < 0.01 cutoff; fig. S4, G to I, Materials and Methods, and data file S9). MC2 (*SPP1*) and MC3 (*AREG*) enrichment in ACs was associated with EGFR signaling (*AREG* and *EGFR*). Lymphotoxin genes (*LTB* and *LTA*) were positively associated with DC2 (*CD1C*) enrichment in ACs. Lymphotoxin is a tumor necrosis factor (TNF) family member whose signaling orchestrates immune cell aggregation at mucosal surfaces and effector immune responses in the tissue, including DC maturation and DC–T cell interactions (63). Secreted LTa forms a heterotrimer with membrane-bound LT $\beta$  to signal via LT $\beta$ R, and we identified *LTBR* as primarily expressed in MNPs and AECs (fig. S4J). These data suggest that lymphotoxin signaling in the airway may play an important role in regulating MNPs in asthma.

#### CCR2-expressing MCs persist in an immature and pathogenic state in asthmatic airways

Monocytes are highly plastic cells that are recruited to the tissue during inflammation and are programmed by the tissue microenvironment (47, 48, 59). Murine models suggest that these cells are transiently pro-inflammatory but subsequently differentiate into Macs that resolve inflammation (48, 59, 62). To determine whether there is any transition between monocyte-derived subsets in the airway mucosa, cellular trajectory was inferred using RNA velocity and latent time analyses and predicted a transition from MC4 (*CCR2*) to MC2 (*SPP1*) and MC3 (*AREG*) with high confidence (Fig. 6A, and fig. S5, A and B, and Materials and Methods). Velocity analysis identified MC2 (*SPP1*) as the terminal population, suggesting that MC3 (*AREG*) is an intermediate state. Cells along this differentiation trajectory sequentially acquired expression of genes associated with Mac identity and survival (*MARCO, PPARG*, and *CSF1*), phagolysosomal function (*LAMP1*,

*CTSD*, and *CTSL*), and lipid metabolism (*APOC1*, *MGLL*, and *FABP5*) (Fig. 6B and data file S10).

We further interrogated this transitional relationship by performing scRNA-seq of human CD14<sup>+</sup> classical monocytes isolated from peripheral blood (9668 cells) and of cocultures of monocytes and primary AECs at air-liquid interface (ALI) after 4 days (8759 cells) and 21 days (15,139 cells) (Fig. 6C, fig. S5, C and D, Materials and Methods, and data files S2 and S4). We identified 10 distinct populations, including 3 MNP and 7 AEC clusters (AUC 0.75 and pseudo-bulk DGE FDR < 0.05; Materials and Methods). MNP clusters included classical monocytes (mono; CD14) and a small population of DC2 (CD1C) from day 0 peripheral blood, as well as monocyte-derived Macs (moMacs; CIQA) identified in both day 4 and 21 cocultures (Fig. 6C and fig. S5C). Using gene set scores derived from our airway mucosal MNP marker genes (Fig. 5B and data file S4), day 0 monocytes from blood aligned with MC4 (CCR2), whereas moMacs after 4 days in coculture aligned most closely with MC2 (SPPI), consistent with the differentiation continuum predicted by RNA velocity analyses (Fig. 6D and fig. S5E). moMacs after 21 days in coculture acquired a transcriptional profile similar to that of Mac2 (A2M) and were highly enriched for complement genes (C1QA, C1QB, and C1QC) and MERTK. Murine AAMs, which are transcriptionally similar to Mac2 (A2M), are replenished by CCR2<sup>+</sup> monocytes, integrate in the airway epithelium, and exhibit dendriform morphology (57, 58). We therefore performed immunofluorescence staining of day 21 cocultures and confirmed that these cells colocalize with the epithelial basal layer, acquire dendritic projections, and express C1Q and MERTK (Fig. 6E).

We subsequently quantified the number of DEGs ( $log_2FC > 0.5$  and FDR < 0.1; Materials and Methods) in MNP subsets after allergen challenge compared with baseline (fig. S5F and data file S6) and found that the transcriptional response was greater in asthmatics compared with ACs (total DEGs = 671 in AAs versus 89 in ACs). MC4 in asthmatics had the greatest transcriptional response to allergen (DEGs = 289), followed by MC2 (DEGs = 122), DC2 (DEG = 94), and Mac2 (DEGs = 48). To identify transcriptional changes in MNPs specific to allergic asthma, we next identified DEGs between groups after allergen challenge (fig. S5G and data file S6). MC2 (SPP1) and MC4 (CCR2) had the most DEGs between groups in response to allergen challenge. In ACs, MC2 (SPP1) up-regulated genes involved in monocyte and Mac chemotaxis and survival (CCL2 and CSFI), phagolysosomal function (MARCO, CD163, and CTSD), and lipid transport and metabolism (ACSL1 and ABCA1) (Fig. 6F). In contrast, MC2 in asthmatics up-regulated genes involved in chemotaxis of immune cells during type 2 inflammation (CCL17 and CCL26), antigen presentation (HLADRB1, HLADPA1, and CIITA), and eicosanoid synthesis (ALOX15 and GGT5). MC4 (CCR2) in ACs also up-regulated genes involved in monocyte and Mac chemotaxis and survival (OSM and CCL4), along with genes that orchestrate tissue repair (HBEGF and VEGFA) and reactive oxidative species clearance (NAMPT and SOD2), whereas asthmatics up-regulated genes implicated in ECM degradation and pathologic remodeling (MMP9, MMP12, and ADAM19) (Fig. 6G).

IPA and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analyses of the DEGs between groups after allergen challenge were used to predict signaling pathways

(pORA < 0.1) and upstream regulators that may be driving transcriptional differences between groups, with |z score| > 2 considered significant (Materials and Methods and data file S8). Distinct predicted signaling pathways were identified in MC2 (*SPP1*) in ACs [phagolysosome, mitogen-activated protein kinase (MAPK), and peroxisome proliferator– activated receptor (PPAR)] compared with asthmatics (antigen processing and presentation, C-type lectin receptor, and Wnt) (fig. S5H). The predicted upstream regulators of MC2 (*SPP1*) in ACs direct Mac survival and identity and included regulators of CSF1 (NCOA1) and PPAR $\gamma$  signaling (MED1), along with NCOA2, which encodes GRIP1 and promotes an anti-inflammatory Mac phenotype (|z score| = 2.0) (Fig. 6H) (64). In contrast, IL-4 (|z score|= 3.36) and IL-13 (|z score| = 2.45) were the strongest predicted drivers of transcriptional changes in MC2 (*SPP1*) in asthmatics.

Distinct predicted signaling pathways were also identified in MC4 (*CCR2*) in ACs [complement cascade, nuclear factor  $\kappa$ B (NF- $\kappa$ B), and PPAR] compared with asthmatics [arachidonic acid metabolism and Janus kinase (JAK)–STAT] (fig. S5H). STAT3 (|*z* score| = 2.12) was the strongest predicted upstream regulator of transcriptional changes in MC4 (*CCR2*) in ACs, whereas a nonsignificant trend was observed in asthmatics for STAT6 (|*z* score| = 1.73), which mediates IL-4/IL-13 signaling (Fig. 6I). Many STAT6-induced genes (*ALOX15*, *CCL17*, and *MMP12*) were enriched across MNP clusters in asthmatics after allergen challenge (fig. S5I). These data suggest that, in ACs, the MC phenotype is characterized by autocrine production of factors important in endocytic clearance, Mac differentiation and survival, and expression of trophic factors promoting angiogenesis and tissue repair. In contrast, IL-4/IL-13 signaling via STAT6 in asthmatics may prevent or arrest Mac differentiation and instead direct a pathogenic MC phenotype characterized by up-regulation of genes involved in inflammatory signaling, antigen presentation, and pathologic airway remodeling (summarized in Fig. 6J).

#### Asthmatic airways are characterized by a pathogenic T<sub>H</sub>2–MNP–basal cell interactome

Cellular cross-talk between airway epithelial and immune cells is critical to the initiation and resolution phases of allergic inflammation (2–4, 65, 66). We therefore used CellPhoneDB (67) to predict receptor-ligand pairs specific to ACs and asthmatics after allergen challenge (empirical P < 0.001 cutoff; Materials and Methods and data file S11). The greatest number of unique predicted interactions in both groups was between AEC and MNP subsets (Fig. 7A and fig. S6A). In asthmatics, the remaining interactions were dominated by T<sub>H</sub>2 cells.

In asthmatics, TNF family members (*TNF*, *LTA*, and *TNFSF14*) and type 2 cytokines (*IL4* and *IL13*) from  $T_H2$  were predicted to interact across AEC and MNP subsets (Fig. 7B and fig. S6B). Protein validation in BAL demonstrated an increase in LIGHT (*TNFSF14*; 229.1 [84.8 to 584.6] versus 119.6 [37.1 to 211.6] pg/ml, P = 0.006) and LTa (43.3 [12.8 to 124.3] versus 11.7 [7.2 to 26.8] pg/ml, P = 0.03) in asthmatics compared with ACs after allergen challenge (Fig. 7C).  $T_H2$  were also predicted to interact with AEC subsets via *IL33R/IL1RL1-IL33* and *LIFR-LIF*.  $T_H2$  and MNPs were predicted to interact via receptorligand pairs involved in T cell localization (*CCR4-CCL17*), immune synapse formation (*ICAM2*–integrin aLβ2 complex), costimulation (*CTLA4-CD86* and *ICOS-ICOSLG*), and T cell polarization (*TGFBR3-TGFB1* and *CCR2-CCL2*) (40, 68). We additionally observed

specific predicted interactions between  $T_H^2$  and DC2, including those important in T cell chemotaxis (*CCR4-CCL22*) and immunomodulation (*CD226-PVR* and *TIGIT-PVR*).

MNPs were predicted to interact with multiple AEC subsets in ACs, whereas basal cells were the primary interacting AEC subset in asthmatics (Fig. 7A). Interactions favoring IL-1 signaling were identified in ACs via CellPhoneDB analysis, as well as by performing linear modeling of the receptor-ligand pairs (FDR < 0.1; Fig. 7D, fig. S6, C to E, Materials and Methods, and data files S11 and S12). Whereas IL1B from MNPs was predicted to interact with its cognate receptor IL1R1 on basal cells in both groups, this cytokine was only predicted to interact with the decoy receptor IL1R2 in asthmatics. IL-1R2 is induced by IL-4/IL-13 and antagonizes the effects of IL-1 $\beta$  (69, 70). In asthmatics, predicted AEC-MNP interactions involved inflammatory leukocyte migration, along with noncanonical Wnt and transforming growth factor  $\beta$  (TGF $\beta$ ) signaling. TGF $\beta$  was predicted to interact with epithelial-specific integrin  $\alpha \nu \beta \delta$ , which mediates TGF $\beta$  activation and drives airway structural remodeling (71). TRAIL (TNFSF10) signaling was also enriched in asthmatics and, in addition to its inflammatory effects, can up-regulate metalloproteinase production and promote airway remodeling in allergic asthma (72). EGFR signaling was predicted to occur between basal cells and distinct MNP-derived ligands in asthmatics compared with ACs. Specifically, EGFR ligands known to promote tissue repair (AREG and HBEGF) characterized ACs (73, 74), whereas asthmatics uniquely interacted via TGFA, which promotes epithelial mucin production (31, 32).

To further characterize AEC-MNP cross-talk, we performed NicheNet analysis (75) to identify the regulatory networks between these cell types that distinguish allergy from asthma (Materials and Methods and data file S13). Thus, we defined ligands and their downstream targets as genes that were differentially expressed between asthmatics and ACs after allergen challenge (data file S6). Cellular communication pathways in ACs were characterized by growth factor signaling and the injury-repair response across subsets (fig. S6, F and G). In contrast, asthmatics were dominated by basal-MNP interactions that included STAT6-induced signaling and mediators of airway remodeling. To validate these findings, we quantified BAL protein levels in asthmatics compared with ACs (Fig. 7E and fig. S6H). Protein levels of STAT6-induced chemokines were higher in asthmatics compared with ACs after allergen challenge (CCL22: 19,080.6 [8876.2 to 22,692.5] versus 7927.6 [1301.3 to 13,504.2] pg/ml, P<0.0001; CCL26: 627.5 [176.1 to 1276.9] versus 76.9 [20.9 to 126.5] pg/ml, P < 0.0001), as were levels of proteins involved in mucin production and pathologic airway remodeling (TGFa: 59.7 [20.1 to 211.4] versus 32.2 [15.6 to 55.2] pg/ml, P = 0.003; matrix metalloproteinase-12 (MMP-12): 745.7 [176.7 to 1957.7] versus 228.6 [69.7 to 493.5] pg/ml, P = 0.02). Overall, these data support a dominant T<sub>H</sub>2–MNP–basal cell interactome that may override a protective injury-repair response observed in ACs, thereby driving asthma pathobiology (summarized in Fig. 7F).

### DISCUSSION

This study directly compared allergic asthmatics with allergic non-asthmatic controls to identify cell subsets and transcriptional effector programs in the lower airway relevant to asthma pathogenesis. In response to allergen challenge, basal and secretory epithelial cells

in asthmatics up-regulated genes involved in matrix degradation, mucus metaplasia, and glycolysis and failed to induce antioxidant pathways observed in ACs. Allergen challenge also induced *IL9*-expressing pathogenic  $T_H^2$  cells and led to enrichment of DC2 (*CD1C*) and immature pro-inflammatory MCs in asthmatic airways. Cross-talk between these cell types may be critical to the persistence and amplification of airway inflammation as well as the pathologic structural remodeling that defines allergic asthma.

Whereas SAC leads to increased eosinophils and type 2 cytokine levels in the airways of both AAs and ACs (9), our current study demonstrates that tissue reprogramming in response to type 2 inflammation is a defining feature of asthma (summarized in Fig. 8). This finding is most clearly observed in airway-resident basal and goblet cells. Basal and secretory epithelial cells in asthmatics up-regulated genes involved in type 2 immune cell chemotaxis, mucus metaplasia, and ECM remodeling after SAC. Asthmatic basal cells up-regulated *IL13RA1* after allergen challenge, suggesting a possible mechanism of heightened responsiveness of these cells to IL-13. In contrast, allergen challenge induced genes involved in epithelial repair and cytoprotection in ACs. Thus, our data suggest that type 2 programming overrides protective and repair pathways in the asthmatic airway epithelium. It remains unknown whether this programming is induced by persistent type 2 airway inflammation or results from an intrinsic susceptibility of AECs in allergic asthma.

Type 2 programming was also a defining feature of MCs recruited to asthmatic airways after SAC. Specifically, asthmatics were enriched for immature MC4 (*CCR2*) that up-regulated IL-13–induced inflammatory mediators and metalloproteinases. In marked contrast, macrophage-like MCs (MC1, MC2, and MC3) dominated the airway mucosal immune landscape in ACs 24 hours after allergen challenge. The proresolving phenotype observed in MCs from ACs included genes encoding scavenger receptors and phagocytic machinery, trophic mediators that promote epithelial repair, and regulators of lipid metabolism. Monocytes are highly plastic cells that are recruited to the tissue during inflammation and acquire functional roles based on environmental signals. Murine models suggest that these cells are transiently pro-inflammatory but subsequently differentiate into macrophages that resolve inflammation (48, 59, 62). Consistent with this, coculture of blood monocytes with AECs predicted a continuum of differentiation from monocytes to macrophage-like MCs. Together, these results suggest that IL-13 may arrest recently recruited MC4 (*CCR2*) in a pro-inflammatory state, thereby preventing differentiation into macrophage-like MCs that restore airway homeostasis.

Our study also identified signaling pathways that may amplify type 2 inflammation and critically link airway inflammation to the structural remodeling observed in asthma. *IL9*-expressing pathogenic  $T_H^2$  cells were highly specific to asthmatic airways after allergen challenge. IL-9 is known to amplify type 2 inflammation, promote mast cell activation and survival, and drive tissue pathology (40–44). Whereas three previous small trials targeting IL-9 in asthma had mixed results, they included heterogeneous asthma cohorts followed over short time periods (76, 77). Our findings, along with recent murine studies (78), suggest that IL-9 may be specific for the allergen recall response and thus may be an important disease mediator in a targeted subset of patients. Murine models of asthma demonstrate that T cell receptor activation by antigen-presenting DCs after allergen

challenge is required for *IL9* induction (41). Airway mucosal DC2 were enriched in AAs after SAC and highly expressed *ALOX15*, a lipoxygenase induced by type 2 cytokines that generates arachidonic acid metabolites. These metabolites can in turn bind to and activate PPAR $\gamma$ , a lipid-activated transcription factor that drives pathogenic T<sub>H</sub>2 effector responses in the airway (79). Cell-cell interaction analysis highlighted additional DC2-T<sub>H</sub>2 interactions comprising a feedforward loop of immunotaxis, DC maturation, and T<sub>H</sub>2 costimulation specific to asthmatic airways. This suggests that local DC2-T<sub>H</sub>2 cross-talk may establish T<sub>H</sub>2 residence in the airway, license the pathogenic T<sub>H</sub>2 phenotype, and promote production of IL-9 via PPAR $\gamma$  activation. A similar DC2-T<sub>H</sub>2 pathogenic circuit was recently described in atopic dermatitis and persisted despite IL-4R $\alpha$  blockade (12). Thus, targeting pathways that recruit, activate, and facilitate the persistence of airway mucosal DC2 and T<sub>H</sub>2 may be necessary to induce remission in allergic disease.

Our data additionally highlight a critical role for TGFa and TNF family members in linking airway inflammation to pathologic tissue remodeling in asthma. TGFa is an IL-13induced EGFR ligand that promotes epithelial mucin production (31, 32), and TGFA-EGFR signaling was predicted to be unique to asthmatics in our interaction analysis. Interaction analyses also predicted that TNF family members LIGHT and TRAIL interact with MC4 (CCR2) to induce MMP9 expression in asthma. Together with MMP-12, which is induced by type 2 cytokines, these MMPs increase collagen synthesis, degrade ECM, and activate TGF $\beta$  (80–83). In addition to being a potent elastase, MMP-12 has important immunoregulatory roles in the lung, including amplification of IL-13 signaling, promotion of eosinophil and MNP chemotaxis, and B cell follicle formation (80, 84). TNF family member lymphotoxin also induces immune cell aggregation at mucosal sites and promotes DC maturation, amplifying local inflammation (63). LTA was highly enriched in pathogenic T<sub>H</sub>2 cells and associated with DC2 enrichment in asthmatic airways. BAL protein levels of TGFa, LIGHT, MMP-12, and LTa increased in asthmatics but not controls after SAC, further supporting their relevance in asthma. These data reveal a T<sub>H</sub>2-MNP-epithelial cellular interactome that may drive asthma pathogenesis and suggest that targeting cross-talk between these cell types may be necessary to induce disease remission.

Our study focused on mild asthmatics in the setting of allergen-induced inflammation, and tissue sampling was restricted to the airway mucosa. Thus, our results may not be generalizable to the full spectrum of asthma, and additional studies that include other asthma endotypes and severities and alternative challenge models are warranted. Although the number of participants was limited, we observed highly significant differences in cluster abundances and DGE between groups that were represented across all participants, supporting the relevance of these findings to allergic asthma. We have not previously observed differences in the airway response to SAC due to participant sex or the allergen administered. However, we cannot rule out that some of our findings are attributable to these factors. Cellular origin and function cannot be definitively ascribed on the basis of gene expression alone, and follow-up studies are needed. Last, a single time point after allergen exposure cannot interrogate the full kinetics of airway inflammation and resolution, and analysis of later time points may identify additional pathologic processes in asthma.

Our data indicate that type 2 cytokines may override the "default" epithelial injury-repair program and proresolution macrophage maturation sequence observed in ACs, thereby programming pathogenic niche circuits that amplify and link airway inflammation to remodeling in asthma. Our findings also support the involvement of additional pathways that act in concert with type 2 inflammation to promote asthma, including TNF family signaling, altered cellular metabolism, failure to engage antioxidant responses, loss of growth factor signaling, or a complex interplay between these mechanisms. In sum, we present an integrated analysis of the airway cellular ecosystem in the context of allergen-induced inflammation and identify cell subsets and transcriptional effector programs specific to asthma, thereby advancing the current understanding of asthma pathogenesis and identifying potential targets to achieve sustained therapeutic responses.

## MATERIALS AND METHODS

#### Study design

Participant recruitment and study protocol-This study was a nonrandomized, nonblinded mechanistic study with the primary aim of comparing the airway mucosal response to allergen challenge in AAs and ACs. Volunteers were screened for eligibility with a full medical history, baseline spirometry and methacholine challenge, and allergen skin testing. Detailed inclusion and exclusion criteria can be found in Supplementary Methods, as previously published (9). Briefly, male and female participants between 18 and 50 years old with a clinical history of allergic rhinitis and/or conjunctivitis to HDM and/or cat hair and a positive skin prick test to the same allergen were enrolled. AAs additionally had a clinical history of mild asthma, baseline  $FEV_1$  75% of the predicted value, and positive methacholine challenge testing defined as a provocative concentration causing a 20% drop in FEV<sub>1</sub> (PC20) <16 mg/ml. Participants could not be on systemic steroids for 6 weeks before the study, and inhaled corticosteroids were held for 2 weeks before the study procedure. All participants provided written informed consent before testing and sample collection. The study protocol was approved by the Massachusetts General Brigham Institutional Review Board (IRB# 2007P001050) and registered at clinicaltrials.gov (NCT00595491). A full list of participants is compiled in data file S1.

**Allergen skin testing**—AAs and ACs had positive SPT to standardized HDM [*Dermatophagoides pteronyssinus*; Greer Laboratories, 10,000 allergy units (AU) per ml] and/or cat hair extract [*Felis catis*; Greer Laboratories, 10,000 bioequivalent allergy units (BAU) per ml]. The threshold level of allergen sensitivity was determined by quantitative SPT using serial threefold dilutions of extract. The lowest concentration of extract eliciting a positive skin prick test (3-mm wheal diameter) was used as the allergen dose administered during SAC (9). The same stock of allergen was used for both SPT and SAC.

**Segmental allergen challenge**—Briefly, baseline BAL samples were obtained by administering 120 ml of normal saline to the lingula. After clearing luminal contents, baseline endobronchial brushings were collected from the lingula using a 4-mm sterile nylon cytology brush under direct visualization (85). Diluent (2 ml) was then administered to the anterior segment of the right upper lobe (RUL) followed by allergen (2 ml) to the lateral

segment of the right middle lobe (RML). Twenty-four hours later, BAL and endobronchial brush samples were obtained in a similar manner from the diluent- and allergen-challenged segmental airways.

### Cytospins

Cell differential counts for BAL were determined by enumerating eosinophils on cytocentrifuge preparations (9).

#### Flow cytometry

Endobronchial brushings were obtained as above and stained with 4',6-diamidino-2phenylindole (DAPI) for dead cell exclusion and fluorescently conjugated antibodies against CD326, CD45, CD3, CD19, HLA-DR, and CD1c after Fc receptor blockade. Detailed information is provided regarding the antibodies used for flow cytometry staining in table S1. Samples were run on a BD FACSAria Fusion flow cytometer (BD Biosciences) and analyzed with FlowJo v10.7 (Tree Star).

#### Cytokine quantification

Cytokines were analyzed by MilliplexMAP kit (EMD Millipore) using 30-fold concentrated BAL supernatant (9). BAL was concentrated in Amicon Ultra-15 centrifugal filter units (EMD Millipore, catalog no. UFC9100) or Sartorius Vivaspin 15R centrifugal concentrators (Thermo Fisher Scientific, catalog no. 14-558-499) according to the manufacturer's instructions. Cytokines were measured on Luminex 200 and analyzed using xPONENT v3.1 (Luminex Corp.).

#### Human classical monocyte purification

Whole blood was collected from healthy volunteers (IRB# 2002P001658), and peripheral blood mononuclear cells (PBMCs) were separated using Ficoll gradient centrifugation. CD14<sup>+</sup> classical monocytes were isolated from PBMCs using magnetic bead isolation (Miltenyi Biotec, catalog no. 130-117-337) in accordance with the manufacturer's protocol. Excess cells were frozen in CryoStor CS10 (BioLife Solutions, catalog no. 210102) and later thawed and serially diluted in phosphate-buffered saline (PBS) + 2 mM EDTA + 0.5% bovine serum albumin (BSA) solution. Dead cells were magnetically removed (BioLegend, catalog no. 480159), and the remaining cells were resuspended at a concentration of 1000 cells per  $\mu$ l in preparation for loading on the 10X Chromium instrument (10X Genomics).

#### Human monocyte and airway epithelial coculture and scRNA-seq

Human primary small AECs (Lonza, catalog no. CC-2547) were cultured at ALI on laminincoated Transwell inserts (Corning, catalog no. 3470) using PneumaCult Ex-Plus followed by ALI medium (STEMCELL Technologies, catalog nos. 05040 and 05001) as previously described (86). CD14<sup>+</sup> classical monocytes were added to the apical chamber at a density of  $9 \times 10^4$  cells per cm<sup>2</sup> after an intact ALI was formed. Before dissociation of cocultures at days 4 and 21 for scRNA-seq, excess mucus was removed by adding 75 µl of 10 mM dithiothreitol (MilliporeSigma, catalog no. 43819) in PBS to the apical chamber for 5 min followed by PBS wash (×2). Cocultures were then dissociated with TrypLE Select (Gibco, catalog no. 12563011) for 15 min. Magnetic CD45<sup>+</sup> enrichment (STEMCELL Technologies, catalog no. 1000105) was used to increase the fraction of recovered MNPs from dissociated cocultures. All samples underwent magnetic dead cell removal (BioLegend, catalog no. 480159) and were resuspended at a target concentration of 1000 cells per  $\mu$ l in preparation for loading on the 10X Chromium instrument (10X Genomics).

#### Immunofluorescence staining

Detailed information is provided regarding the antibodies used for immunofluorescence staining in table S1. Staining for MERTK was performed on live cells by first adding Fc receptor blockade for 1 hour before adding anti-human CD45 and anti-human MERTK for 2 hours at 37°C. All other coculture staining was performed after fixation using 4% paraformaldehyde (PFA). Cells were permeabilized with 0.1% Triton X-100 and blocked with 1% BSA–0.3% Triton X-100. Cells were stained with mouse anti-human CD45 (2 hours at 37°C), rat anti-human C1q (overnight at 4°C), and rabbit anti-human p63 (overnight at 4°C). Secondary antibodies were added for 2 hours at 37°C along with Hoechst 33258. Fluorescent images were obtained using an inverted Zeiss LSM780 confocal laser-scanning microscope.

Lung tissue was collected from a lobectomy specimen of a patient with a clinical history of asthma (IRB# 2013P002332). Tissue was fixed overnight in PFA, washed three times, dehydrated in 30% sucrose, embedded in optimal cutting temperature (OCT) (Sakura), and cryosectioned. Slides were stained with antibodies against cytokeratin 13 (KRT13) and acetylated tubulin, and coverslips were mounted in antifade medium containing DAPI. Images were obtained on a ZEISS Axio Imager M2 widefield fluorescent microscope. All images were processed using Fiji or Zen Blue (Zeiss).

#### Single-cell RNA sequencing and computational data analysis

#### Preparation of single-cell suspensions from endobronchial brushings-

Endobronchial brush samples were collected in phenol-free RPMI 1640 (Thermo Fisher Scientific) with 2% human AB serum and ROCK inhibitor (Y-27632, Tocris Bioscience, catalog no. 1254) on ice and processed within 60 min of collection. Cells were gently removed from cytology brushes and resuspended in phenol-free RPMI 1640. Single-cell suspensions were depleted of dead cells using annexin V–conjugated beads (STEMCELL Technologies, catalog no. 17899) and of red blood cells using antibodies directed against glycophorin A (STEMCELL Technologies, catalog no. 18170). Cell viability was assessed via trypan blue (0.4%) staining and manual cell counting using a hemocytometer both before and after red blood cell and dead cell removal to achieve >95% viability after dead cell removal. Viable cells were resuspended at a concentration of 800 to 1000 cells per  $\mu$ l in preparation for loading on the 10X Chromium controller instrument (10X Genomics).

**Single-cell RNA sequencing**—About 12,000 single cells were loaded to each 10X channel with a recovery goal of 6000 single cells. Cell suspensions were loaded along with reverse transcriptase reagents, 3' gel beads, and emulsification oil onto separate channels of a 10X Single Cell B Chip, which was loaded into the 10X Chromium instrument to generate emulsions. Emulsions were transferred to polymerase chain reaction (PCR)

strip tubes for immediate processing and reverse transcription. Library preparation was performed according to the manufacturer's recommendations. Expression libraries were generated either using Chromium Single Cell 3'V3 chemistry (10X Genomics PN-120262; endobronchial brush samples) or Next GEM Single Cell 3'V3.1 chemistry (10X Genomics PN-1000268; coculture samples). DNA and library quality was evaluated using an Agilent 2100 Bioanalyzer, and concentration was quantified using the Qubit dsDNA high-sensitivity reagents (Thermo Fisher Scientific). Gene expression libraries were sequenced on an Illumina NextSeq instrument. Endobronchial brush samples were sequenced on the Illumina NextSeq 500/550 system using the following single-index sequencing configuration: read 1 = 28 base pairs (bp) [cell barcode, unique molecular identifier (UMI)], read 2 = 56bp (insert), index 1 = 8 bp (sample index), and index 2 = 0 bp. Coculture samples were sequenced on the Illumina NextSeq 2000 system using the following dual-index sequencing configuration: read 1 = 28 bp (cell barcode, UMI), read 2 = 96 bp (insert), index 1 = 10 bp (sample index), and index 2 = 10 bp (sample index). All sequencing statistics are compiled in data file S2. Detailed methods for read alignment and quantification, cell clustering, marker gene identification, DGE analysis, and all other downstream computational analyses can be found in Supplementary Methods.

**Statistical analysis**—Statistical approaches for computational analyses of scRNA-seq data are detailed in Supplementary Methods. Mixed-effects modeling with repeated measures was used to compare median differences in cellular proportions and protein levels between groups unless otherwise specified, where experimental condition and disease group were considered fixed effects and each participant was treated as a random effect. Adjustment for multiple comparisons was performed using Šidák's correction. A two-sided P < 0.05 was considered significant.

### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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### Data and materials availability:

scRNA-seq count matrices and related data are deposited in the GEO database (under accession number GSE193816), and raw human sequencing data are available in the controlled access repository dbGaP (www.ncbi.nlm.nih.gov/gap/) under accession number phs003101.v1.p1. Source code for data analysis in each main figure is available on GitHub (https://github.com/villani-lab/airway\_allergic\_asthma) and has been archived to Zenodo (https://zenodo.org/badge/latestdoi/412260708). A user-friendly portal is available to browse the single-cell data generated in this manuscript. Users can query the single-cell clustering solutions, visualize gene expression, and browse differential expression of genes. Users may navigate the portal by visiting https://villani.mgh.harvard.edu/allergy-asthma.

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#### Fig. 1. A human model of allergen-induced asthma exacerbation using SAC.

(A) Characteristics of AC and AA participants. (B) Design of the bronchoscopic SAC procedure. BAL and endobronchial brush samples were obtained at baseline (from the lingula) and 24 hours after administration of diluent to the right upper lobe (RUL) and allergen to the right middle lobe (RML). Representative images of RML airway segments from one AC participant (top row) and one AA participant (bottom row) at baseline (left) and after SAC (right). In the AA participant, the allergen-challenged segmental airway is narrowed, and mucus is present after SAC. (C) Representative images of cytospin preparations from BAL from one AA participant and quantification of BAL eosinophils in nonallergic healthy controls (HCs; n = 5), ACs (n = 20), and AAs (n = 22). Black

arrows indicate eosinophils; white arrows indicate lymphocytes. (**D**) Representative flow cytometry of endobronchial brush samples at baseline (Bln; top row) and after allergen challenge (Ag; bottom row) from one AA participant. Epithelial cells were identified as CD326<sup>+</sup>CD45<sup>-</sup>. After gating on CD45<sup>+</sup> cells, CD19<sup>+</sup> B cells and CD4<sup>+</sup> T cells were identified, as were CD19<sup>-</sup>CD4<sup>-</sup>HLA-DR<sup>+</sup> antigen-presenting cells. (**E**) scRNA-seq was performed on endobronchial brushing samples (n = 21) collected from segmental airways, and downstream analysis was performed to identify differences between ACs (n = 4) and AAs (n = 4). In (C), boxes represent the median (line) and IQR, with whiskers extending to the remainder of the distribution no more than  $1.5 \times$  IQR, with dots representing individual samples. *P* values were generated using a mixed-effects model with Šidák correction to adjust for multiple comparisons. \*\*\**P* < 0.001 and \*\*\*\**P* < 0.0001.





CD8 T

CD4 T

AEC

(A) UMAP embedding of 52,152 high-quality single cells color-coded by predicted cell lineage. NK, natural killer. (B) Heatmap showing the top discriminative gene sets for each cell lineage compared with every other lineage. Color scales denote the normalized gene expression (mean zero, unit variance) for each cluster and the mean number of genes per cluster (top bar). (C) Dot plot showing the percentage and expression level of marker genes defining each cell lineage, with the size of the dot representing the percentage of cells within each lineage expressing each marker and the color intensity indicating the scaled expression level. (D) UMAP embedding of cell density displaying the proportion of each cell lineage at baseline (left) and after allergen challenge (right) compared with every other cell lineage, faceted by disease group (ACs: top, AAs: bottom). (E) OR of disease association by cell lineage at baseline and after allergen challenge. Color-coding denotes

MNP

B cells

Mast cells

NK cells

significant associations with ACs (OR < 1, purple) or AAs (OR > 1, gold). (**F**) Contribution of each cell lineage defined in (B), shown as percentage (%) of total sample at baseline (Bln) and after allergen challenge (Ag). In (E), dots and whiskers represent OR with 95% confidence interval, calculated using a mixed-effects association logistic regression model (P < 0.05 corrected for multiple comparisons using the Tukey method). In (F), boxes represent the median (line) and IQR, with whiskers extending to the remainder of the distribution no more than  $1.5 \times$  IQR and dots representing individual samples. *P* values generated using a mixed-effects model with Šidák correction to adjust for multiple comparisons. \*P < 0.05 and \*\*P < 0.01.



Fig. 3. Dynamic transcriptional response in basal and secretory epithelial cells after allergen challenge.

(A) UMAP embedding derived from subclustering of 20,410 AECs. (B) Heatmap showing the top discriminative gene sets for each cluster compared with every other AEC cluster. Color scales denote the normalized gene expression (mean zero, unit variance) for each cluster and the mean number of genes per cluster (top bar). (C) Dot plot depicting gene expression levels and percentage of cells expressing genes across AECs. (D) UMAP plots with color intensity (top) indicating the number of DEGs induced by SAC in ACs and AAs compared with baseline. The number of DEGs induced by SAC, quantified by cluster for AAs and ACs (bottom). (E) Venn diagram depicting the top five AEC clusters with the most DEGs after SAC, with the number of genes up-regulated (up arrow) or down-regulated (down arrow) in AAs compared with ACs using an interaction term for disease state and experimental condition. The core transcriptional response is denoted in the table (right). Bolded genes are induced by IL-13. (F) Heatmap of selected DEGs. Color scale indicates FC differences between AAs and ACs after SAC. White dot indicates FDR < 0.1. (D) DEG based on FDR < 0.1 and  $\log_2 FC > 0.5$  using the Wald test on pseudo-bulk count matrix. (E and F) DEGs based on FDR < 0.1 and  $\log_2 FC > 0.5$  using a likelihood ratio test on pseudo-bulk count matrix.



#### Fig. 4. IL9-expressing pathogenic T<sub>H</sub>2 cells specific to asthmatic airways.

(A) UMAP embedding derived from subclustering of 18,714 T cells. (B) Heatmap showing the top discriminative gene sets for each cluster, compared with every other T cell cluster and the mean number of genes per cluster (top bar). (C) GSEA of CD4  $T_H2$ ,  $T_H17$ , and T<sub>H</sub>IFNR clusters comparing the marker genes for these clusters with published T cell gene sets from Seumois et al. (33) (data file S7). (D) UMAP embedding of cell density displaying the proportion of each T cell subset at baseline (left) and after allergen challenge (right) compared with every other T cell subset, faceted by disease group (ACs: top, AAs: bottom). (E) OR of disease association by cluster at baseline and after allergen challenge. (F) Feature plots of pathogenic T<sub>H</sub>2 genes using pseudo-coloring to indicate gene expression. (G) Feature plot of IL9 expression using pseudo-coloring to indicate gene expression, faceted by group. (H) BAL concentration of IL-9 (ACs, n = 14; AAs, n = 18). In (E), dots and whiskers represent OR with 95% confidence interval, calculated using a mixed-effects association logistic regression model (P < 0.05 corrected for multiple comparisons using the Tukey method). In (F) and (G), cell number and percentages (%) represent gene expression across all T cell subsets. Scaled gene expression in log(CPM). In (H), boxes represent the median (line) and IQR, with whiskers extending to the remainder of the distribution no more than  $1.5 \times$  IQR. P values were generated using a mixed-effects model with Šidák correction to adjust for multiple comparisons. \*P < 0.05, \*\*P < 0.01, and \*\*\*\*P < 0.0001.





(A) UMAP embedding derived from subclustering of 8510 MNPs. (B) Heatmap showing the top discriminative gene sets for each cluster compared with every other MNP cluster and the mean number of genes per cluster (top bar). (C) Dot plot depicting gene expression levels and percentage of cells expressing genes across MNP clusters. (D) UMAP embedding of cell density displaying the proportion of each MNP subset at baseline (left) and after allergen challenge (right) compared with every other MNP subset, faceted by disease group (ACs: top, AAs: bottom). (E) OR of disease association by cluster at baseline and after allergen challenge. In (E), dots and whiskers represent OR with 95% confidence interval, calculated using a mixed-effects association logistic regression model (P < 0.05 corrected for multiple comparisons using the Tukey method).



**Fig. 6. A pathogenic transcriptional program in airway MCs in asthmatics after SAC.** (**A**) RNAvelocities of a subset of MNP clusters visualized as streamlines projected on UMAP embedding. (**B**) Heatmap of gene expression pattern for the top 100 lineage driver genes correlated along the inferred differentiation trajectory. (**C**) Coculture model for blood CD14<sup>+</sup> monocytes and AECs at ALI. UMAP embedding derived from clustering of 33,566 cells consisting of baseline blood CD14<sup>+</sup> monocytes and cocultured cells collected at day 4 (d4) and day 21 (d21). Mono, monocyte; moMac, monocyte-derived Mac; mito, mitochondrial. (**D**) Alignment of blood CD14<sup>+</sup> monocytes (day 0; d0) and moMacs from d4 and d21 coculture, using gene set scores (*x* axis) of airway MNP subclusters (Fig. 5A). (**E**) Immunofluorescence staining with orthogonal reconstruction at d21 of coculture demonstrates CD45<sup>+</sup> immune cells (magenta) integrated into the p63<sup>+</sup> basal cell layer (yellow) with Hoechst nuclear counterstain (cyan). CD45<sup>+</sup> immune cells (magenta) exhibit

prominent cytoplasmic projections and expression of MERTK and C1q (yellow). (**F** and **G**) Volcano plots representing DEGs in MC2 (F) and MC4 (G), comparing ACs (purple) with AAs (gold) after SAC. Vertical dotted lines represent cutoff of  $|log_2FC| = 0.5$ , and horizontal dotted lines represent FDR cutoff = 0.1. (**H** and **I**) Predicted upstream regulators of MC2 (H) and MC4 (I) based on DEGs identified in (F) and (G) (ACs: purple, AAs: gold). Vertical solid lines represent *z*-score cutoff of |2|. (**J**) Summary figure of MC maturation sequence. (F and G) DEGs based on FDR < 0.1 and  $log_2FC > 0.5$  using the Wald test on pseudo-bulk count matrix.





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Fig. 7. A pathogenic cellular interactome in asthmatic airways after allergen challenge. (A) Top 25 cell-cell pairs with the most unique receptor-ligand interactions in ACs and AAs, restricted to interactions between distinct cell lineages. Tiers represent the number of interactions (tier 1: 50, tier 2: 40 to 49, tier 3: 30 to 39). (B) Dot plots showing predicted interactions between T<sub>H</sub>2-AECs and T<sub>H</sub>2-MNPs after SAC in AAs. (C) BAL concentration of LIGHT (*TNFSF14*; ACs: *n* = 13, AAs: *n* = 14) and LTa (ACs: *n* = 13, AAs: *n* = 16). (D) Dot plots showing predicted interactions between AEC-MNP after SAC in AAs and ACs. (E) BAL concentration of CCL22, CCL26, MMP-12, and TGFa (ACs: *n* = 13, AAs: n = 14). (F) Summary figure of a pathogenic T<sub>H</sub>2–MNP–basal cell interactome in allergic

asthma. In (B) and (D), dot size indicates significance (true: empirical P < 0.001), and color intensity indicates specificity of interaction to disease group  $[-\log_{10}(\text{rank})]$ . In (C) and (F), boxes represent the median (line) and IQR, with whiskers extending to the remainder of the distribution no more than  $1.5 \times$  IQR. In (C) and (F), *P* values were generated using a mixed-effects model with Šidák correction to adjust for multiple comparisons. \*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001, and \*\*\*\*P < 0.0001.



# Fig. 8. Transcriptional reprogramming in response to type 2 airway inflammation as a defining feature of allergic asthma.

After SAC, the airway landscape in allergic non-asthmatic controls was characterized by enrichment of macrophage-like MCs expressing growth factors associated with tissue repair as well as up-regulation of antioxidant genes in basal and goblet cells. In contrast, asthmatics failed to up-regulate these pathways and instead had enrichment of DC2 (*CD1C*), pro-inflammatory MC4 (*CCR2*), and *IL9*-expressing pathogenic T<sub>H</sub>2 cells. Type 2 cytokines and other mediators, including TNF family members, produced by T<sub>H</sub>2 cells may promote a pathogenic MC4 phenotype in asthmatics by interrupting the default MC maturation sequence to proresolution macrophages. These T<sub>H</sub>2 mediators may also override the epithelial injury response observed in ACs and instead promote ECM degradation, subepithelial fibrosis, and mucus metaplasia in asthma. Figure was created using biorender.com.