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The balance between protective and pathogenic immune responses to pneumonia in the neonatal lung is enforced by gut microbiota

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SUPPLEMENTARY MATERIALS

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Materials and Methods

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MDAR Reproducibility Checklist

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Abstract

Although modern clinical practices such as cesarean sections and perinatal antibiotics have improved infant survival, treatment with broad-spectrum antibiotics alters intestinal microbiota and causes dysbiosis. Infants exposed to perinatal antibiotics have an increased likelihood of life-threatening infections, including pneumonia. Here, we investigated how the gut microbiota sculpt pulmonary immune responses, promoting recovery and resolution of infection in newborn rhesus macaques. Early-life antibiotic exposure interrupted the maturation of intestinal commensal bacteria and disrupted the developmental trajectory of the pulmonary immune system, as assessed by single-cell proteomic and transcriptomic analyses. Early-life antibiotic exposure rendered newborn macaques more susceptible to bacterial pneumonia, concurrent with increases in neutrophil senescence and hyperinflammation, broad inflammatory cytokine signaling, and macrophage dysfunction. This pathogenic reprogramming of pulmonary immunity was further reflected by a hyperinflammatory signature in all pulmonary immune cell subsets coupled with a global loss of tissue-protective, homeostatic pathways in the lungs of dysbiotic newborns. Fecal microbiota transfer was associated with partial correction of the broad immune maladaptations and protection against severe pneumonia. These data demonstrate the importance of intestinal microbiota in programming pulmonary immunity and support the idea that gut microbiota promote the balance between pathways driving tissue repair and inflammatory responses associated with clinical recovery from infection in infants. Our results highlight a potential role for microbial transfer for immune support in these at-risk infants.

INTRODUCTION

Distinct immune responses adapted for early postnatal life also render newborns more vulnerable to infections (1). Incomplete understanding of immune development early in life contributes to our inability to reduce neonatal morbidity caused by respiratory infections, which kill more newborns than any other cause (2). Whereas a series of coordinated events control the development of the newborn immune system (3), few are as important as the interactions of immune cells with successive waves of commensal bacteria that colonize the newborn's intestine after birth (4, 5). Evolving microbial signals play a critical role in the maturation of neutrophils, CD4⁺ T cells, monocytes, and innate lymphoid cells (ILCs) in the peripheral blood and the intestine (6, 7). Nevertheless, the role of intestinal commensal bacteria in functional programming of effector immune cells at extraintestinal mucosal sites, such as the lungs, is unclear (8).

Nearly half of vaginally delivered infants are exposed to antibiotics given to their mother to prevent group B *Streptococcus* sepsis (9–11). Some of these infants continue to receive empiric antibiotics, such as ampicillin and gentamicin, for neonatal sepsis (12). The use of vancomycin, although still uncommon, has increased because of the increased incidence of coagulase-negative *Staphylococcus* sepsis, especially in sick and preterm neonates (13). Although current clinical practices are essential for reducing infant mortality, they may extract a substantial biological cost in long-term disruption of the gut microbiota. Infants

treated with antibiotics experience greater morbidity in response to sepsis and pneumonia compared to milder disease and spontaneous resolution in infants not exposed to antibiotics (14, 15).

In this study, we defined how gut microbiota inform immune responses that mediate recovery and resolution rather than severe disease. We used a rhesus macaque model because it more closely resembles a human infant lung in structure, developmental stage, physiology, and mucosal immune mechanisms (16) compared to murine lungs (17). In addition, similar to humans, nonhuman primates develop lobar pneumonia and demonstrate a heterogeneous clinical response (18). Their larger size and similarity to human newborns enabled integration of clinical signs, longitudinal assessment of disease progression, and the ability to distinguish mild, self-resolving pneumonia from severe, often fatal pneumonia.

Here, we defined a tiered immunologic development program, anchored by stepwise engagement of effector cells to limit tissue damage and promote recovery, which was experimentally disrupted by early-life antibiotic exposure. Remodeling of the pulmonary immune response was anchored by the appearance of a population of senescent and hyperinflammatory neutrophils and dysregulated antigen presentation in macrophages. This was all coupled with global loss of tissue-protective, homeostatic pathways in dysbiotic newborns. Fecal microbiota transfer was associated with correction of broad immune maladaptations and protection against severe pneumonia in dysbiotic newborn macaques. Our data suggest that gut microbiota reinforce the balance between regenerative pathways driving tissue homeostasis and inflammatory responses, thereby limiting pathogenesis and promoting clinical recovery. Our results also highlight a potential role for microbiota transfer as immune support in these at-risk infants.

RESULTS

Antibiotic exposure during the first postnatal week delays microbiota maturation and reconfigures the peripheral immune system before infection

We treated a cohort of vaginally delivered, nursery raised rhesus macaques (table S1) with either a cocktail of antimicrobials or with saline ($n = 4$ in each experimental group) from postnatal day 1 (PN1) to PN7 (Fig. 1A) and profiled the fecal bacterial communities daily. Stools in saline-treated newborn macaques, referred to hereafter as control newborns, were dominated by facultative aerobic Enterobacteriaceae during the first week of life before changing to strict anaerobes, principally *Bacteroides* and *Clostridium* in the second week of life (table S2), similar to the pattern seen in human neonates (19, 20). Maternal microbiota, a critical driver of fecal microbial colonization (21–23) and immune development in the newborn (21, 24), were not different between the experimental groups (table S3).

Antibiotic use diminished the abundance of Enterobacteriaceae, reduced phylogenetic diversity, and delayed the maturation of intestinal microbiota (Fig. 1, B and C, and table S2). Antibiotic use also created a maladapted immune state, highlighted by reduced amounts of hematopoietic cytokines, for example, granulocyte colony-stimulating factor (G-CSF) and interleukin-2 (IL-2) (25–27), and cytokines critical for mobilization of monocytes from bone marrow, such as C-X3-C motif chemokine ligand 1 (CX3CL1) and C-X-C motif chemokine

ligand 8 (CXCL8) (28, 29), as assessed by a nonhuman primate multiplex cytokine kit (Fig. 1D and table S4). These immune changes occurred contemporaneously with changes in the stool microbiota (Fig. 1D and tables S2 and S4 to S6).

Peripheral blood neutrophils in antibiotic-exposed macaques, hereafter referred to as dysbiotic newborns, were phenotypically distinct, marked by increased expression of CD11b, a marker for neutrophil activation (30), and programmed cell death protein 1 (PD-1), a marker associated with senescence and exhaustion (Fig. 1E) (31, 32). The frequency and absolute number of neutrophils were decreased in the dysbiotic newborns (Fig. 1F; fig. S1, A to D; and table S5). Peripheral blood T cells in dysbiotic newborns were also phenotypically distinct with higher expression of CD45RA, CD38, and CD57, which are markers associated with T cell differentiation (33) and senescence (Fig. 1E) (34–36).

Gene networks associated with “cellular stress,” “ubiquitination,” and “apoptosis” were enriched in the transcriptomic analysis of peripheral blood from dysbiotic newborns (fig. S1E and table S7). Collectively, these data demonstrated that disruption of commensal microbes by early-life antibiotic use was associated with a maladapted state marked by both a robust, proinflammatory bias and neutrophil and T cell activation.

Early-life antibiotic exposure is associated with clinically severe pneumonia

After discontinuing antibiotics for 1 week, we challenged the newborn macaques in each experimental group with *Streptococcus pneumoniae* (serotype 19F), a common and often fatal respiratory pathogen in human infants (37). Vital signs, including heart rate, respiratory effort, blood pressure, urine output, oxygen saturation, and overall clinical condition of these newborn macaques, were monitored every 6 hours. In addition, chest radiographs were obtained daily. The pediatric early warning score (PEWS) (38), an extensively validated clinical decision-making and severity-scoring tool, was used to guide therapy, including intravenous fluids and supplemental oxygen (table S8).

The clinical status of dysbiotic newborns deteriorated, evidenced by higher PEWS and radiographic signs of lung consolidation (Fig. 2, A and B; fig. S2, A and B; and table S8). The progression of symptoms was also more rapid in the dysbiotic newborns (Fig. 2C), and all dysbiotic newborns received supportive treatment by 60 hours (Fig. 2D). At this time point, dysbiotic newborns had more severe disease (PEWS = 8 ± 1.5) than control newborns (PEWS = 5 ± 0.6) despite similar pathogen burden (fig. S2, C and D). These data recapitulate the rapid progression and increased morbidity seen in dysbiotic human infants (39).

Last, we sought to determine whether a unique immune signature in dysbiotic newborns before infections could predict disease severity and clinical outcomes after challenge with *S. pneumoniae*. We found that reduced serum concentrations of hematopoietic cytokines—G-CSF, IL-2, CX3CL1, and CXCL8—before infection were associated with lower PEWS and less severe illness after challenge with *S. pneumoniae* (fig. S2E). In contrast, increased serum levels of proinflammatory cytokines, such as IL-1 β , IL-6, IL-17, IL-21, and tumor necrosis factor- α (TNF- α), before infection were associated with higher PEWS and more severe illness after challenge with *S. pneumoniae* (fig. S2E). These data highlight the

relevance of our newborn macaque model in profiling immune responses in target tissues, such as the lungs, which is often impossible in human newborns.

Early-life antibiotic exposure remodels pulmonary immune responses to *S. pneumoniae* infection

The host inflammatory response to pneumonia is highly compartmentalized to the lungs (40), so systemic immune cell or cytokine responses alone may not fully explain the differential clinical response to pneumonia in dysbiotic newborns. We therefore used a combination of high-parameter (>20 markers) cytometry by time of flight (CyTOF) and single-cell RNA sequencing (scRNA-seq) to profile the innate and adaptive immune cell responses in the lungs of control or dysbiotic newborns at 60 hours after infection with *S. pneumoniae*. The frequency of neutrophils measured by CyTOF (fig. S3, A to E) decreased significantly ($P < 0.01$) in the lungs of dysbiotic newborns compared to control newborns after infection with *S. pneumoniae* (fig. S3F). In contrast, the frequency of CD4⁺ T cells (fig. S3, A to E) increased significantly ($P < 0.01$) in the lungs of dysbiotic newborns after infection with *S. pneumoniae* (fig. S3F).

Cell clusters identified by scRNA-seq of pulmonary immune cells after infection with *S. pneumoniae* in control or dysbiotic newborns were annotated using signature genes from published scRNA-seq atlases (Fig. 3, A and B, and table S9) (41, 42). In dysbiotic newborns, the proportion of neutrophils identified by scRNA-seq decreased, whereas the proportion of T cells increased, consistent with the CyTOF findings (table S9).

Immune pathways dominated by inflammation, such as IL-1 β activation, and dysfunction, such as apoptosis and cellular stress, were seen in all pulmonary immune cells of dysbiotic newborns (fig. S4A and table S10). Canonical immune programs anchored by cell migration, differentiation, activation, and tissue repair were also broadly disrupted in these same cells after infection (fig. S4A). Neutrophils and alveolar macrophages (AMs) were the principal sources of inflammatory gene programs after infection with *S. pneumoniae* (fig. S4B), consistent with their critical roles in lung defense against pneumonia. Neutrophils and AMs also exhibited transcriptomic remodeling (Fig. 3C) and differentially expressed transcripts related to inflammation and tissue repair (fig. S4A) in dysbiotic newborn macaques after infection.

Consistent with transcriptomic changes, cytokines associated with inflammation, such as CXCL8, CXCL10, and transforming growth factor- β (TGF β), and activation, such as IL-6, IL-8, and TNF- α , were increased in the bronchial lavage fluid of dysbiotic newborns, whereas cytokines associated with tissue repair, such as vascular endothelial growth factor (VEGF), IL-10, and platelet-derived growth factor (PDGF), were decreased (fig. S4C and table S11). A similar reconfiguration of immune responses was associated with rapid disease progression, worse symptoms, and increased mortality in humans in both bacterial and viral [influenza and severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)] pneumonia (43–45).

Paired scRNA-seq and CyTOF analyses of matched blood and lung samples reveal compartmentalized production of chemokines and activation of neutrophils in the lungs

Immune cells, including neutrophils in peripheral blood and lung, were phenotypically distinct (fig. S5, A and B, and table S12). Compared to neutrophils from the lungs, neutrophils in the peripheral blood expressed higher amounts of transcripts for chemokine and homing/egress receptors, such as *CX3CR1*, C-C motif chemokine receptor 2 (*CCR2*), and *SELL* (encoding CD62L) (fig. S5, C and D). In contrast, pulmonary neutrophils expressed higher amounts of proinflammatory mediators, including chemokines for recruitment of monocytes and other neutrophils, such as C-C motif chemokine ligand 2 (*CCL2*), *CCL3*, and *CCL4*. They also express higher amounts of complement components, such as complement C1q B chain (*C1QB*) and *C1QC*; antimicrobial peptides, such as *S100A8* and *S100A9*; and matrix metalloproteinases (MMPs), such as *MMP8* and *MMP9*, which are implicated in tissue damage in acute respiratory distress syndrome (ARDS) (fig. S5, C and D) (46–48).

Consistent with these scRNA-seq findings, cytokines associated with neutrophil and monocyte migration, such as *CCL2*, *CCL4*, and *CCL11*; activation, such as IL-1 β , IL-6, IL-8, IL-17, and TNF- α ; and inflammation, such as *CXCL8*, *CXCL10*, and TGF β , were uniformly abundant in the lungs (fig. S5E). The frequency of activated (*CXCR2*^{lo}, *CXCR4*^{hi}, and *CD62L*^{lo}) neutrophils was significantly higher in the lungs compared to the peripheral blood ($P < 0.05$; fig. S5F). These data support the idea of compartmentalized activation of neutrophils, which may further explain the increased morbidity of dysbiotic newborns.

Senescent and hyperinflammatory neutrophils dominate pulmonary immune responses to *S. pneumoniae* infection in dysbiotic newborns

We identified three transcriptionally distinct neutrophil populations in the lungs after infection with *S. pneumoniae* (Fig. 3, D to F; fig. S6A; and table S13). Neutrophil heterogeneity is influenced by developmentally encoded cell programs and by environment and pathogen-specific factors (49). To identify distinct neutrophil populations, we used module scores (42), reflecting the average expression of all genes related to neutrophil development, maturation, and activation using published gene signatures from granulocytes during homeostasis and in the setting of sepsis (Fig. 3F and table S14) (50). Cluster 1 represented mature neutrophils defined by the expression of genes associated with pathogen response and cytokine signaling, reflecting neutrophil heterogeneity caused by pathogen exposure (fig. S6, A and B). Cluster 2 represented hyperinflammatory, stressed neutrophils characterized by high expression of *CXCR4* and *CD63* and reduced expression of *CXCR2* and *SELL* (fig. S6, A and B) (51). These cells also had increased gene transcripts associated with glycolysis, a pathway associated with hyperinflammatory responses (fig. S6, B and C) (52). Cluster 3 represented immature neutrophils identified on the basis of the expression of gene transcripts related to neutrophil maturation (secretory vesicles, lysozymes, and phagocytosis) (fig. S6, A and B). The developmental relationship among these three clusters was predicted by the cellular trajectory analysis (Fig. 3, G and H) (53). Pseudo-time analysis of granule proteins, olfactomedin 4 (*OLFM4*) and *SELL*, and trafficking receptors, *CXCR2* and *CXCR4*, known to be involved in neutrophil maturation, suggests a continuum of

differentiation from immature neutrophils (cluster 3) to mature neutrophils (cluster 1) to senescent, hyperinflammatory neutrophils (cluster 2) (Fig. 3H).

Early-life antibiotic exposure strongly influenced the activation state of the lung neutrophil compartment after infection with *S. pneumoniae*. Pseudo-bulk RNA-seq analysis of these pulmonary neutrophils identified distinct signatures associated with dysbiosis (fig. S6C). Senescent, hyperinflammatory neutrophils were unique to the lungs of dysbiotic newborns, whereas mature neutrophils were absent in the lungs of dysbiotic newborns (Fig. 3I). CyTOF also showed consistent remodeling of the pulmonary neutrophil pool with the emergence of stressed, senescent (CXCR2^{lo}, CXCR4^{hi}, and CD62L^{lo}) neutrophils in the lungs of dysbiotic newborns after infection with *S. pneumoniae* (Fig. 3J).

There was broad induction of *NFκB*, enrichment of phagocytosis and degranulation gene sets, and increased expression of epigenetic regulators associated with inflammatory neutrophils, including protein-arginine deiminase 4 (*PADI4*), which is required for neutrophil extracellular trap formation (NETosis) (54, 55), and *CD274* [encoding for programmed cell death ligand 1 (PD-L1)], a marker of cell exhaustion in pulmonary neutrophils from dysbiotic newborns (table S15). We noted consistent changes in neutrophil-activating and chemotactic cytokines in the bronchial washings of dysbiotic newborns after infection with *S. pneumoniae* (fig. S6D). A similar reconfiguration of immune responses was associated with rapid disease progression, worse symptoms, and increased mortality in humans in both bacterial and viral (influenza and SARS-CoV-2) pneumonia (43–45).

Predicted regulatory networks active in pulmonary neutrophils from dysbiotic newborns after infection with *S. pneumoniae* include KDMA5 and NFIL3

Immature neutrophils (cluster 3) were enriched for regulatory networks anchored by the transcription factors (TFs) CCAAT/enhancer binding protein γ (C/EBP γ) and Kruppel-like factor 6 (KLF6), which are essential for neutrophil development (56–61), whereas mature neutrophils (cluster 1) were enriched for regulatory networks anchored by defense response-associated TFs (62–64), such as NFKB, interferon (IFN) regulatory factor 7 (IRF7), signal transducer and activator of transcription 5A (STAT5A), basic leucine zipper activating TF-like TF 3 (BATF3), and hypoxia-inducible factor 1A (HIF1A) (Fig. 3K, fig. S6E, and table S16). In contrast, the senescent, hyperinflammatory neutrophils (cluster 2) were enriched for regulatory networks anchored by epigenome-modifying enzymes, lysine demethylase 5A (KDM5A), histone deacetylase 2 (HDAC2), and nuclear factor IL-3 (NFIL3), a component of the circadian clock (Fig. 3K, fig. S6E, and table S16). NFIL3 and KDM5A are implicated in the microbiota-driven regulation of both nonhematopoietic cells, such as the intestinal epithelium, and hematopoietic cells, such as ILCs and natural killer (NK) cells, by chromatin histone modification (65, 66). Disruption of NFIL3 and KDM5A regulatory pathways may explain neutrophil dysfunction in dysbiotic newborns and highlights the need for experimental validation of their role in neutrophil homeostasis.

Gene scores of hyperinflammatory neutrophils from dysbiotic newborn macaques predict disease severity in an independent human cohort

Dysfunctional neutrophils contribute to pulmonary damage in experimental models of acute lung injury (67). We therefore hypothesized that the senescent, hyperinflammatory neutrophils from the dysbiotic newborn macaques would predict severe pneumonia outcomes. We identified the five most differentially expressed genes between senescent, hyperinflammatory neutrophils and all other cells in our dataset: *HIF1A*, *CXCR4*, *CD274*, lactotransferrin (*LTF*), and *S100A8*. We then used publicly available whole-blood bulk transcriptomic datasets of infants and children with severe sepsis and pneumonia as a reference for scoring with this identified neutrophil gene signature (68). We scored each sample in these datasets by the aggregated expression of these five genes. Our senescent, hyperinflammatory neutrophil gene score predicted sepsis diagnosis (area under the curve (AUC) = 0.79) in a cohort of at-risk infants (Fig. 3, L and M, and tables S17 and S18). Our data suggest that gene signatures from senescent, hyperinflammatory neutrophils might be potential prognostic indicators in human infants with sepsis and pneumonia.

Macrophages with dysfunctional features are a hallmark of the remodeled pulmonary immune response in the dysbiotic macaques

To interrogate macrophage plasticity, which is informed by developmental programs and tissue- and stress-specific signals, we used MacSpectrum (69), an analytical tool to stratify macrophage maturation and activation. The dominant population enriched for gene transcripts associated with antigen processing and presentation represents terminally differentiated mature AM (cluster 1); the population enriched for transcripts associated with inflammation, purinergic-inflammasome signaling, and IL-1 receptor activation represents polarized, inflammatory AMs (cluster 2); and immature AMs were represented by the expression of gene transcripts associated with DNA replication and cell division (cluster 3) (Fig. 4A; fig. S7, A to D; and table S19). Cellular trajectory analysis further identified a relationship between differentiation and activation as the AMs developed from immature (cluster 3) to mature (cluster 1) to polarized inflammatory AMs (cluster 2) (Fig. 4, B and C).

Early-life antibiotic exposure influenced the development and activation state of AMs after infection with *S. pneumoniae*. Mature AMs (cluster 1), known to maintain noninflammatory states by promoting tolerance and facilitating tissue repair (70), were decreased, whereas the frequency of polarized, inflammatory AMs (cluster 2) was increased in dysbiotic newborn macaques (Fig. 4D). In addition, the frequency of M1-activated AMs [identified as live major histocompatibility complex II-positive (MHC class II⁺), CD11C⁺, and CD86⁺ cells] was increased in dysbiotic newborns after infection with *S. pneumoniae* (Fig. 4E) and correlated with disease severity (fig. S7E).

We hypothesized that dysfunctional inflammatory macrophages were associated with severe pneumonia in dysbiotic macaques. Using published gene signatures from AMs in ARDS in humans (50), we found that gene transcripts predicting recovery were enriched in cluster 1, whereas gene transcripts predicting severe ARDS/death were enriched in cluster 2 (Fig. 4F and table S20). The expression of genes related to tissue damage and vascular inflammation was differentially enriched in dysbiotic macaques (Fig. 4G and table S21).

Adenosine triphosphate (ATP) released from damaged epithelium and endothelium activates an ATP-driven purinergic-inflammasome signaling pathway and is associated with fatal pneumonia and severe ARDS (71). We found increased ATP in the bronchial lavage fluid of dysbiotic macaques, although this did not reach statistical significance ($P=0.06$) (fig. S7F). Last, we noted decreased expression of tolerance-promoting MHC class II genes in AMs of dysbiotic newborns (Fig. 4G).

Conserved gene regulatory networks are associated with remodeled pulmonary neutrophil and macrophage compartments in dysbiotic newborns

Regulatory networks anchored by peroxisome proliferator-activated receptor γ (PPAR γ), C/EBP α , PU.1 (encoded by *SPI1*) (72), and upstream stimulatory factor 1 (USF1), TFs critical in macrophage development (73), were enriched in all mature AMs, whereas HIF1A and early growth response (EGR), critical drivers of transcriptional programs underlying macrophage activation and terminal polarization (74), were enriched in stressed and hyperinflammatory macrophages (Fig. 4H and table S22).

A regulatory network anchored by aryl hydrocarbon receptor nuclear translocator-like protein 1 (ARNTL) [encoded by brain and muscle ARNT-like 1 (*BMAL1*)], a component of the circadian clock, was overrepresented in the stressed and hyperinflammatory macrophages (cluster 2) that were expanded in dysbiotic newborns (Fig. 4H and fig. S7G). Diurnal oscillations of intestinal microbiota are thought to drive the programming of host immune responses via ARNTL (75) and other components of the circadian clock. In addition, retinoid X receptor alpha (RXR α) and NFIL3 were overrepresented in regulatory networks for cluster 2 (Fig. 4H). These data, coupled with similar observations in neutrophils (Fig. 3K), suggested a role for shared regulatory networks anchored by circadian clock components in the transcriptional remodeling of pulmonary neutrophil and AM compartments in dysbiotic newborns.

Antibiotic exposure also remodeled the interstitial macrophage (IM) pool after infection with *S. pneumoniae*. Analogous to AMs, we identified three unique populations of mature IMs marked by expression of genes related to macrophage migration, phagocytosis, tolerance promotion, and wound repair (cluster 1); activated IMs identified by their expression of genes related to antigen presentation, IL-1 receptor activation, and CD4⁺ T cell differentiation (cluster 2); and a population marked by expression of genes associated with endoplasmic reticulum stress, exhaustion, and apoptotic clearing, identified as exhausted and stressed IM (cluster 3) (fig. S7, H and I). In addition, antibiotic exposure was associated with contraction of pro-repair IMs and expansion of stressed, exhausted proinflammatory IMs (fig. S7, J and K, and table S23). These data collectively suggested the loss of phagocytic, tolerance-promoting, and antigen-presenting programs, which facilitate protective functions of lung macrophages and expansion of ATP-purinergic inflammasome signaling and proinflammatory programs, and are associated with lung damage and increased pneumonia-related morbidity in dysbiotic macaques.

Dysbiotic newborns display ineffective pulmonary T helper cell responses after infection with *S. pneumoniae*

Consistent with transcriptomic changes seen in other pulmonary cell types, gene transcripts associated with tissue repair and growth factor signaling, T cell fitness, and effector differentiation were decreased, whereas transcripts for genes in the aerobic glycolysis pathway, linked to T cell exhaustion (76), were increased in T cells of dysbiotic newborns (Fig. 4I and table S24). Pulmonary CD4⁺ T cells from dysbiotic newborns coexpressed T cell immunoglobulin and mucin-domain containing-3 (TIM-3) and lymphocyte activating gene-3 (LAG-3), which are markers linked to T cell exhaustion (fig. S8A) (31, 77). The frequency of dysfunctional CD4⁺ T cells (expressing LAG-3) correlated with disease severity (Fig. 4J). Furthermore, cytokines associated with T cell differentiation and effector responses, such as IL-6, IL-17, CXCL8, and CXCL10, were increased in the bronchial lavage fluid of dysbiotic macaques (Fig. 4K). Our data highlighted T cell responses marked by a failure to limit “cytotoxicity” and potentially increased tendency toward “exhaustion” in dysbiotic newborns, which is consistent with recent studies highlighting a critical role of exhausted T cells in severe coronavirus disease 2019 (COVID-19) (78).

Early-life antibiotic exposure increases stressed, inflammatory NK cells in lungs of dysbiotic newborns after infection with *S. pneumoniae*

Optimal NK cell functions promote infection control by serving as a rheostat in regulating T cell responses with excessive activation contributing to immunopathology (79). Consistent with these observations, we found that most NK cells in the lungs of dysbiotic newborns coexpressed activation marker CD69 and produced granzyme B and IFN- γ (fig. S9A).

Similar remodeling of the pulmonary NK pool was supported by scRNA-seq data. Expression of cytotoxic effector and activation markers, such as NK cell granule protein 7 (*NKG7*), *CD38*, and *CD52* (44, 80, 81), and proliferation markers, such as minichromosome maintenance (*MCM*), proliferating cell nuclear antigen (*PCNA*), and eukaryotic translation initiation factor 4A1 (*EIF4A1*), identified CD56⁺ NK cells (cluster 2) and proliferating NK cells (cluster 3), respectively, whereas a cluster expressing transcripts associated with cell survival, cellular stress, and inflammation represented cytotoxic and stressed NK cells (cluster 1) (fig. S9, B and C). The numbers of cytotoxic NK cells expanded in dysbiotic newborns (fig. S9D). The transcriptomic analysis further revealed an increased abundance of transcripts associated with T cell activation (82), inflammation, and exhaustion in dysbiotic newborns (fig. S9E and table S25). These data, coupled with reports implicating dysfunctional cytotoxic NK cell responses in severe COVID-19 (44), suggest that defects in NK cell cytotoxicity may be associated with adverse outcomes in dysbiotic newborns.

Miscommunication between innate and adaptive immune cells uncouples inflammatory and pro-repair pathways and contributes to increased pneumonia severity in dysbiotic newborns

To identify altered cell-cell communication in the lungs of dysbiotic newborns after infection, we used the analytical tool CellChat (83) to infer cell signaling interactions based on ligand-receptor transcriptional abundance. Globally, signaling pathways related to chemotactic localization, such as CXCL, CCL, and selectin P ligand (SELPLG), and

tissue homeostasis and repair (84, 85), such as NOTCH and semaphorin 4 (SEMA4), were dominant in control newborns, whereas signaling pathways related to inflammation, such as CD86 and RESISTIN (86, 87), and cell exhaustion, such as PD-L1 and PD-L2, were dominant in dysbiotic newborns (Fig. 5, A and B), potentially suggesting a global rewiring of immune cell-to-cell communication network in dysbiotic newborn macaques (Fig. 5C and fig. S10, A to C).

Dysfunctional macrophages were the central hub of misdirected cell-cell communications (Fig. 5, C to E, and fig. S10, A and B). In contrast, neutrophils and, to a lesser extent, T cells were predicted to be targets of such miscommunications, which included immune-inhibitory interactions, such as CD274(PD-L1)–programmed cell death protein 1 (PDCD1), macrophage migration inhibitory factor (MIF)–CD74/CD44/CXCR4, CD86–CD28, and protein tyrosine phosphatase receptor type C (PTPRC)–CD22 in dysbiotic newborns (Fig. 5, D and F, and fig. S10, C to F). In control newborns, communication circuits between macrophages and neutrophils or T cells were dominated by pathways related to interleukin and chemokine signaling, such as IL-1A/B–IL-1R2 and CXCL1/3/8–CXCR2; immune costimulation and complement activation, such as C3–C3AR1; and tissue repair, such as NOTCH, SEMA4, and THY1 (Fig. 5D and fig. S10, G to I) (88, 89). These data identified a potential mismatch between inflammatory and pro-repair pathways anchored by dysfunctional macrophages in dysbiotic newborns.

To identify the communication circuits informing the pathogenic remodeling of the pulmonary neutrophil pool, we interrogated specific signaling circuits between AMs and neutrophils. AM-derived paracrine signals related to neutrophil migration, such as CXCL–CXCR2 and thrombospondin 1 (THBS1)–CD47; neutrophil extravasation, such as integrin-subunit β 2 (ITGB2)–intracellular adhesion molecule 1 (ICAM1); and neutrophil activation, such as SELPLG–SELL, were enriched in control newborns. In contrast, pathways related to programmed cell death [CD80–CD274(PD-L1)] were dominant in dysbiotic newborns (Fig. 5, E to G, and fig. S10, C to E). PD-L1 negatively regulates the activation of T cell receptors and mediates lymphocyte apoptosis. Severe and often fatal sepsis is marked by increased neutrophil PD-L1 expression (88), suggesting a role for PD-L1 in pathogenic responses.

Collectively, these findings highlight the miscommunication between innate and adaptive immune cells in dysbiotic newborns, contributing to a hyperinflammatory signature in neutrophils and relative immune paralysis in the T cells (fig. S10, J and K). This pattern supports the concept that the uncoupling of inflammatory and pro-repair pathways contributes to the increased pneumonia severity in dysbiotic newborns.

Fecal transfer is associated with favorable changes in pulmonary immune cell responses and improved host resistance to pneumonia in dysbiotic newborns

Although no specific bacterial taxa have been consistently associated with pulmonary host resistance to pathogens, fecal microbiota transplantation, which transfers the entire gut microbiota from one host to another, has demonstrated improved clinical outcomes in immunotherapy trials (90, 91). We performed fecal transfer (FT) of the fecal contents of control newborns to dysbiotic newborns on PN8. Dysbiotic newborns who received the FT, referred hereafter as FT recipient newborns, were challenged with *S. pneumoniae* on PN14

(6 days after FT). FT recipients had lower PEWS after infection, less rapid progression of symptoms, and reduced need for supportive therapy (Fig. 6, A to D, and fig. S11A). FT recipients also showed a bacterial burden closer to control (fig. S11B). Although all recipients demonstrated clinical benefit, the benefit was variable (Fig. 6, D and E).

Gut microbiota composition of the FT recipients (after treatment) differed from their baseline (pre-FT) (Fig. 6F and tables S26 and S27). After FT, gut microbiota had a higher abundance of *Bifidobacterium bifidum*, a favorable modulator of immune responses in humans (Fig. 6G) (7, 20). Although our study lacked the power to establish a clear association between specific bacterial taxa and clinical response to pneumonia, our results suggest that FT could be feasible in dysbiotic newborns.

Next, we evaluated the immunological effects of FT after challenge with *S. pneumoniae*. The pulmonary neutrophil pool expanded after FT, although the numbers remained lower than control newborns (fig. S12, A and B). FT partially corrected the remodeling of the neutrophil pool observed in dysbiotic newborns (Fig. 6H). FT was associated with changes in the transcriptome of neutrophils compared to dysbiotic macaques. Transcripts of genes related to leukocyte apoptosis, such as cathepsin C (*CTSC*), TNF type 1-associated DEATH domain (*TRADD*), PYD and CARD domain containing (*PYCARD*), and *CIQBP*, respiratory burst, such as myeloperoxidase (*MPO*), neutrophil cytosolic factor (*NCF1*), and *NCF2*; and cellular stress and ubiquitination, such as *HIF1A*, *EGR1*, *NOP53*, and TRAFF family member-associated nuclear factor κ B (NF κ B) activator (*TANK*), which were previously increased in neutrophils of dysbiotic macaques, were decreased in FT recipient newborn macaques (Fig. 6I, fig. S12C, and table S28).

Expression of genes related to inflammasome or IL-1 signaling, such as nucleotide-binding oligomerization domain, leucine-rich repeat and pyrin-domain containing (*NLRP3*), *IL1B*, *IL10RA*, and *NFKB1*, decreased in AMs of FT recipient macaques compared to dysbiotic macaques, whereas gene transcripts related to macrophage migration, such as *CCR2*, *CCL3*, and *CX3CL1*; phagocytosis, such as C-type lectin domain containing (*CLEC7A*), *CD47*, and *C3*; molecules promoting tolerance, such as *CCR2*, adenosine deaminase (*ADA*), *IL10*, and Fc-receptor like 3 (FCRL3); and wound repair, such as angiogenin (*ANG*), *IL10*, and *VEGFA*, remained unchanged in FT recipients (Fig. 6I, fig. S12D, and table S29). Transcripts associated with T cell activation and differentiation, such as *IL7R*, *CCR7*, *CD3D*, and *CD3E*, and antigen processing and presentation, such as *CCR4*, inducible T cell co-stimulator (*ICOS*), and *LYN*, which were severely decreased in T cells of dysbiotic newborns, partially increased after FT (Fig. 6I, fig. S12E, and table S30). Transcripts associated with tissue repair and growth, such as VEGF, PDGFA, and EGF, did not recover (Fig. 6I and table S30).

Miscommunication between the innate and adaptive immune cells was reversed after FT, although signaling pathways associated with tissue repair, such as NOTCH, IL-10, and SEMA4, and chemotaxis, such as CXCL, remained dysregulated in FT recipients (Fig. 6J and fig. S12F). These data suggested that, although FT reversed cell stress and apoptosis and mitigated proinflammatory signatures in the pulmonary immune cells, it did not fully

reverse the loss of innate immune cell migration, differentiation, activation, and tissue repair signatures.

DISCUSSION

We leveraged a nonhuman primate model and systems immunology approach to examine the effect of antibiotic-induced dysbiosis on the clinical and immunologic progression of neonatal pneumonia, the most common infectious cause of infant mortality (2). Early-life antibiotic exposure was associated with an overrepresentation of *Enterococcus* and *Clostridium*, approximating the gut microbial communities in a large cohort of vaginally delivered infants exposed to perinatal antibiotics (92, 93) and increased morbidity to *S. pneumoniae*, recapitulating epidemiological observations (93). Early-life antibiotic exposure was associated with a remodeled pulmonary neutrophil compartment and the presence of senescent neutrophils with exhausted and hyperinflammatory signatures. We speculate that extended neutrophil life span and the resulting exhaustion, coupled with failure to remove senescent and exhausted neutrophils from the infected lungs, caused severe tissue damage because of the release of proteases, cationic peptides, and cytokines, which were increased in dysbiotic newborn macaques. An extended neutrophil life span was observed in patients with sepsis (94) and ARDS (95) and was associated with disease progression and poor prognosis.

Dysbiotic newborn macaques also displayed strong inflammatory patterns, terminally polarized macrophages, and a trend toward increased ATP amounts. Expanded ATP–purinergic inflammasome signaling is associated with ARDS, fibrosis, and worse clinical outcomes in viral pneumonia, including severe COVID-19 (96, 97). We speculate that, in the absence of tolerance-promoting and antigen-presenting function, the proinflammatory program in macrophages is pathogenically turned on in dysbiotic newborn macaques, contributing to excessive lung damage and increased infection-related morbidity. These data suggest that therapeutic approaches targeting hyperinflammation, neutrophil clearance from inflamed tissues, or induction of neutrophil apoptosis may potentially improve clinical outcomes in infected, dysbiotic infants.

Intestinal microbiota undergoes diurnal compositional and functional oscillations (98), which, in turn, contribute to the homeostatic programming of host immune cells via clock genes. Antibiotics disrupt these diurnal oscillations of the microbiota and lead to desynchronization of the circadian clock network both locally in the intestine and systemically (98). Circadian oscillations, anchored by BMAL1 and NFIL3 in the sympathetic neurons, regulate the egress of neutrophils from the bone marrow into the periphery via the CXCL12-dependent mechanism (99). Trafficking of neutrophils from blood to infected tissues is constrained by BMAL1-dependent regulation of CXCR4 and CCL2 expression (100, 101), and neutrophil aging and clearance of senescent neutrophils are regulated by circadian rhythms (51, 102). Last, NFIL3-driven circadian rhythms limit macrophage heterogeneity and amplify the pathogenic inflammatory responses in the macrophages (103). These observations, coupled with our present data identifying these TFs in regulatory networks controlling immune cells in dysbiotic newborns, suggest that

disrupted circadian oscillations underpin both the disrupted neutrophil maturation and macrophage dysfunction.

The present study identifies dysfunctional macrophages as a hub of misdirected cell-cell communications. Macrophages and dendritic cells sense respiratory pathogens and initiate immune responses, resulting in rapid recruitment of target cells, such as neutrophils, through the secretion of first-order cytokines, such as CXCR2/CXCL1, CXCL8, IL-1A, and IL-1B/IL-R2. Our study suggests that neutrophils transform the first-order cytokine signals into second-order cytokines that enhance the trafficking and extravasation [oncostatin M (OSM)– leukemia inhibitory factor receptor (LIFR)], immune coactivation (complement C3-C3AR1), and effector function, such as phagocytosis [PTPRC–mannose receptor C-type 1 (MRC1)], to eliminate pathogens. At the same time, reciprocal interactions limit macrophage activation (MIF-CD74/CXCR2 and CD83–platelet and endothelial adhesion molecule 1 (PECAM1)) and promote tissue repair factors (CSF1-CSFR1, TGFBR3-TGFB1, SEMA4–neuropilin (NRP)). This stepwise, tiered response was disrupted in dysbiotic macaques because of dysfunctional macrophages. Immune-inhibitory interactions, such as CD274(PD-L1)-PDCD1, MIF-CD74/CD44/CXCR4, CD86-CD28, and PTPRC-CD22, which promote neutrophil and lymphocyte dysfunction, dominated immune responses in dysbiotic newborns. Similarly, macrophage-anchored signaling pathways related to neutrophil migration, such as CXCL-CXCR2 and THBS1-CD47; neutrophil extravasation, such as ITGB2-ICAM1; and neutrophil activation, such as SELPLG-SELL, were disrupted in dysbiotic newborns. These changes in the communication circuits may explain the shared hyperinflammatory signature in all immune cells and global loss of tissue-protective, homeostatic pathways.

We show that FT was associated with restored immune responses and protection of the dysbiotic newborn against pneumonia. Our findings show that FT shifted the composition of the recipient's microbiota toward Gammaproteobacteria and *B. bifidum*, taxa favoring beneficial immune responses in humans. FT mitigated certain aspects of immune dysfunction seen in dysbiotic newborns. Although broad dysregulation in signaling pathways related to inflammation, complement activation, and cell exhaustion was corrected in FT-recipients, miscommunication in pathways related to tissue repair and chemotaxis persisted.

Several scenarios may explain this partial restoration. Distinct taxa drive the maturation of individual immune cells, so a lack of response may reflect the absence of favorable taxa in FT or failure of such taxa to engraft. A more logical explanation is that resolution of pneumonia requires a tiered response with the sequential engagement of innate immune cells and first-order cytokines, which activate other innate and adaptive immune cells via second-order cytokines. FT may not completely mitigate the disruption of such first- and second-order effectors, most likely because of persistent macrophage dysfunction, contributing to suboptimal benefit. Nevertheless, our study provides proof-of-concept evidence for the ability of FT to improve clinical outcomes in at-risk dysbiotic newborns.

Our study has some limitations. The necessity of frequent clinical examination, sample collection, and invasive procedures precluded us from using dam-reared infants. Therefore,

infants in our study received a diet consisting of formula, not breast milk. As infants grow, feeding practices play an increasing role in determining the composition of the infant gut microbiota (104–106); however, delivery mode and perinatal antibiotic use have a stronger influence on the composition of the microbial community immediately after birth (107). Nevertheless, further studies are needed to delineate the relative contribution of infant diet and microbial metabolites (108) on pulmonary immune maturation during infancy.

In conclusion, these findings reinforce the concept that the gut microbiota play a crucial role in programming the balance between tissue inflammatory and regenerative pathways in response to infection and highlight the role of a healthy gut microbiome in limiting pathogens and promoting clinical recovery.

MATERIALS AND METHODS

Study design

The overall goal of this study was to use a nonhuman primate model to investigate the immune response to respiratory pathogens within the lungs. Our overall hypothesis was that disruption of early-life microbiota would lead to dysfunctional pulmonary immune responses and worsened outcomes to bacterial pneumonia. The Institutional Animal Care and Use Committees (IACUCs) at Cincinnati Children’s Hospital Medical Center and University of California at Davis approved all the animal studies (IACUC2016–19222), which were carried out in Association for the Assessment and Accreditation of Laboratory Animal Care–accredited facilities at the California National Primate Research Center (CNPRC). Because there were no previously published studies in newborn primates, published data in newborn mice were used to estimate that four animals in each group would be sufficient to detect a 20% difference in morbidity with 80% power and an α of 0.05. Twelve vaginally delivered Indian-origin rhesus macaque infants (*Macaca mulatta*) (table S1) were used in this study, which was conducted per National Institutes of Health (NIH)’s *Guide for the Care and Use of Laboratory Animals*. Infant macaques were separated from their dams (for demographic characteristics of macaque dams, see table S31), raised in a nursery from the day of birth, and exposed to a normal light cycle (lights on for 12 hours starting at 08:00). The infant macaques had a diet consisting of Enfamil Lipil with iron for the study duration. The necessity of frequent clinical examination, sample collection, and invasive procedures precluded us from using dam-reared infants in our study. For PN1 to PN7, infant macaques were randomized to one of three treatment groups: saline ($n = 4$; control); a cocktail of vancomycin, gentamicin, and ampicillin ($n = 4$; dysbiotic); or the same cocktail of antibiotics and a FT at PN8 ($n = 4$; FT; see the “Fecal transplant” section below). This antibiotic regimen targeted both Gram-positive and Gram-negative intestinal bacteria and mimicked antibiotic exposure in human infants.

At PN7, antibiotics were discontinued to allow 1 week of washout before subsequent infection on PN14. Clinical assessments (both parameters for PEWS and other clinical parameters) were made by the veterinary staff caring for the rhesus macaque neonates, and these veterinary staff were blinded to the intervention. PEWS values were calculated as previously described (38). Briefly, clinical parameters, such as heart rate and respiratory rate, are given scores on the basis of predetermined ranges and are combined for an overall

PEWS, with a higher PEWS reflecting higher clinical severity of the disease. The veterinary staff provided supportive care to the rhesus macaque newborns based on a predetermined PEWS threshold, which consisted of intravenous fluids if PEWS = 8 and intravenous fluids plus oxygen supplementation if PEWS = 9.

Infectious studies

We grew *S. pneumoniae* serotype 19A (American Type Culture Collection, 700674) with gentle aeration (37°C, 200 rpm) in tryptic soy broth to log-phase growth. Neonatal macaques were inoculated with *S. pneumoniae* (10^6 colony-forming units) via the intratracheal route on PN14. See Supplementary Materials and Methods for full description.

Fecal transplant

FTs were performed in four newborn macaques who had received the cocktail of vancomycin, gentamicin, and ampicillin from PN1 to PN7. This group was separate from the other group of newborn macaques that just received the cocktail of antibiotics (the dysbiotic newborns). See Supplementary Materials and Methods for full description.

Microbiota analysis

Fecal samples were collected daily (PN1 to PN14) from saline-treated (control) or antibiotic-treated (dysbiotic infants) infants and from dams at the time of delivery. Infant fecal samples were collected daily (PN10 to PN14) in FT recipient infants. Fecal samples were frozen (-80°C) immediately after collection. All samples were analyzed together to prevent batch effects. See Supplementary Materials and Methods for full description.

Cytometry by time of flight

Frozen whole-blood leukocytes or CD326⁻CD31⁻CD45⁺ lung cells were thawed and plated at 1×10^6 cells per well in a 96-well U-bottom plate. All samples were analyzed together to prevent batch effects. See Supplementary Materials and Methods for full description.

Flow cytometry

Pulmonary CD4⁺ T cells, AMs, and pulmonary NK cells were thawed and further characterized using markers for functional status, as seen and noted in Fig. 4 and fig. S9. See table S32 for the specific antibodies used. Stained cells were acquired on a custom four-laser BD LSRFortessa flow cytometer (Becton Dickinson) with FACSDiva software and analyzed with FlowJo Software (TreeStar).

Single-cell RNA sequencing

Frozen CD326⁻CD31⁻CD45⁺ cells from digested lungs or frozen peripheral blood mononuclear cells were thawed and resuspended in 50 μl at a concentration of 1000 cells/ μl . After excluding dead cells, ~20,000 cells (~3 to 4000 cells per animal, barcoded to identify individual animals) were then loaded into one channel of the Chromium system using the v3 Single Cell Reagent Kit (10x Genomics) at the Cincinnati Children's Hospital Medical Center DNA Sequencing and Genotyping Core. After capture and lysis, complementary DNA (cDNA) was synthesized and amplified as per the manufacturer's protocol (10x

Genomics). The amplified cDNA was used to construct Illumina sequencing libraries that were each sequenced using an Illumina HiSeq 4000. See Supplementary Materials and Methods for full details on the analyses done.

Alignment and quality control

Raw sequencing data were aligned to the rhesus macaque reference Mmul_10 with Cell Ranger 1.3 (10x Genomics), generating expression count matrix files (see table S33 for dataset metrics). Cells that had fewer than 750 unique molecular identifiers (UMIs) or greater than 15,000 UMIs, as well as cells that contained greater than 20% of reads from mitochondrial genes or ribosomal RNA genes (RNA18S5 or RNA28S5) or hemoglobin genes, were considered low quality and were removed from further analysis. Putative multiplets were removed with DoubletFinder (version 2.0). Genes that were expressed in fewer than 10 cells were removed from the final count matrix.

Data analysis

The Seurat package (version 3.1.0; <https://satijalab.org/seurat/>) was used to identify common cell types across different experimental conditions, differential expression analysis, and most visualizations. Percentages of mitochondrial genes, ribosomal genes, and hemoglobin genes were regressed during data scaling to remove unwanted variation due to cell quality using the SCTransform() function in Seurat. Principal components analysis was performed using the 3000 most highly variable genes, and the first 20 principal components (PCs) were used to perform uniform manifold approximation and projection to embed the dataset into two dimensions. Next, the first 20 PCs were used to construct a shared nearest neighbor (SNN) graph [FindNeighbors()], and this SNN was used to cluster the dataset [FindClusters()]. Manual annotation of cellular identity was performed by finding differentially expressed genes for each cluster using Seurat's implementation of the Wilcoxon rank sum test [FindMarkers()] and comparing those markers to known cell type-specific genes from published studies (41, 42). Global differential gene expression profiles between all cell types were identified and organized with the software cellHarmony (109), using these Seurat clusters [fold change > 1.2, empirical Bayes *t* test $P < 0.05$, false discovery rate (FDR) corrected].

Sepsis diagnosis prediction using senescent neutrophil differentially expressed genes

To test whether the gene signature of senescent neutrophils could be used to predict sepsis diagnosis, we first developed a five-gene signature of the senescent neutrophils by identifying the most differentially enriched genes in cluster 2 neutrophils in our transcriptomic dataset relative to all other cells. Next, we downloaded normalized transcript counts from a publicly available whole-blood bulk transcriptomic dataset (GSE696686) (68). We then scored each sample in this dataset by the expression of the five genes enriched in our senescent neutrophil cluster (*HIF1A*, *CXCR4*, *CD274*, *LTF*, and *S100A8*). Last, we used these gene signature scores as a predictor variable and disease/diagnosis metadata reported by Wynn *et al.* (68) as the response variable to construct a receiver operating characteristic curve to quantify and visualize the sensitivity and specificity of the prediction.

Statistical analysis

All data met the assumptions of the statistical tests used. Statistical tests used for microbiome or single-cell analyses are described in relevant sections. For comparing the differences between groups, we used either unpaired two-tailed Student's *t* test, analysis of variance (ANOVA), or Wilcoxon signed-rank test. We used Pearson correlation coefficient to measure correlation between different variables. We used the Kaplan-Meier log-rank test to compare morbidity between groups (all in GraphPad Prism 8.0). *P* values are indicated as follows: **P* 0.05 or ***P* 0.01.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data and materials availability:

All data associated with this study are present in the paper or the Supplementary Materials. Data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus (accession number GSE176408). Processed data can be further explored at the Lung MAP atlas portal at https://research.cchmc.org/pbge/lunggens/lungExternal/Deshmukh_Macaque_2021.html. The reagents and metadata included in this manuscript are not covered by any material transfer agreement (MTA) and can be obtained from the corresponding author upon request. Scripts used for data analysis are available from GitHub at https://github.com/Deshmukh-Lab/2021_Stevens_Macaque (DOI: 10.5281/zenodo.6569295).

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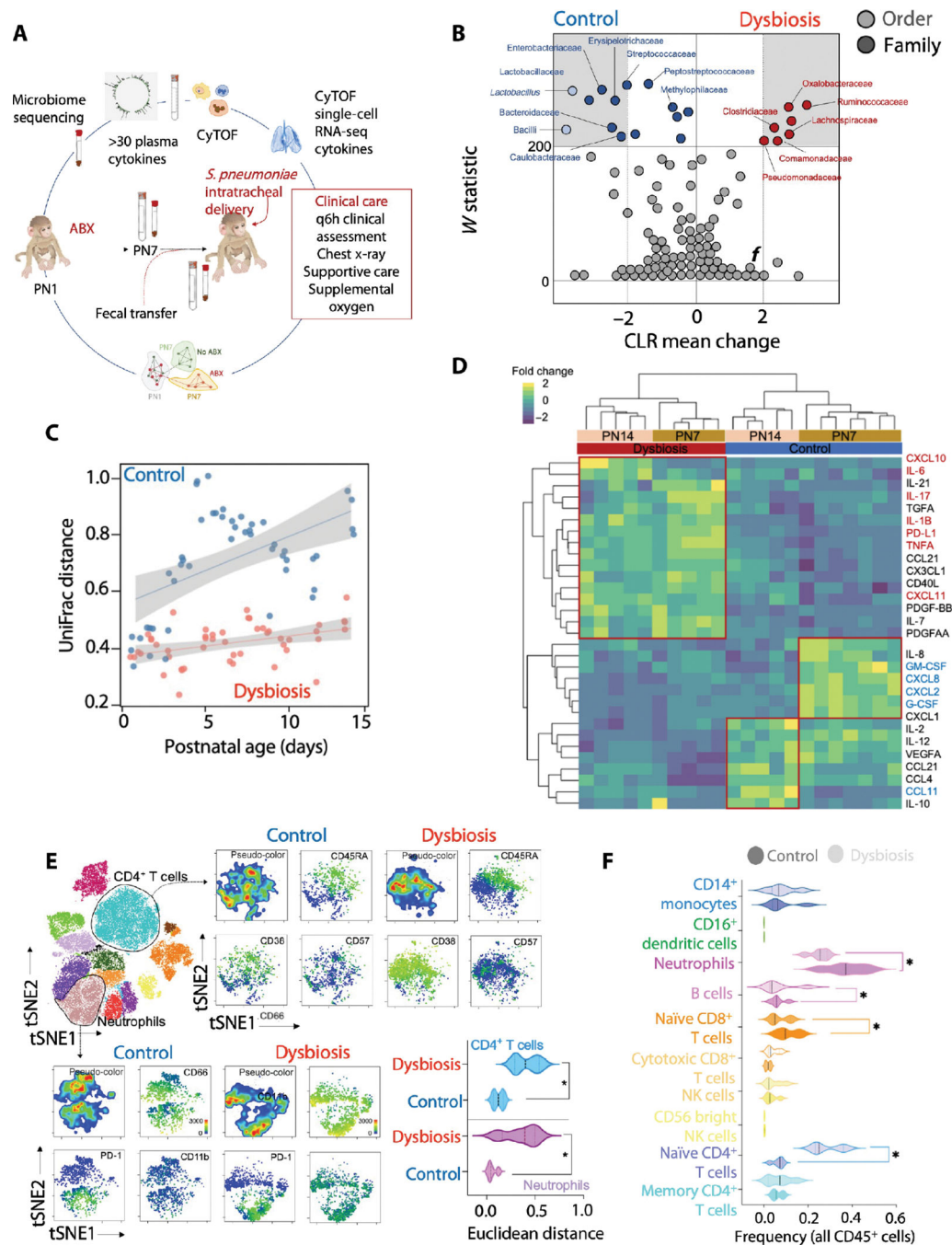


Fig. 1. Antibiotic exposure during the first week of life is associated with delayed microbiota maturation and reconfiguration of the peripheral immune system.

(A) Cohort of vaginally delivered, nursery-raised rhesus macaques was treated with a cocktail of antimicrobials (ABX) from postnatal day 1 (PN1) to PN7 (dysbiosis) or with saline (control) ($n = 4$ in each experimental group). (B) Differentially abundant taxa [FDR $q < 0.05$, center log transformation (CLR) > 2] between control and dysbiosis groups are presented in gray margins. The center log transformation mean difference represents compositional differences in microbial communities. (C) β -Diversity (unweighted UniFrac)

of fecal bacterial communities at the indicated day of life in control and dysbiotic newborn macaques. Linear fit is shown, and margins represent 95% confidence limits. **(D)** Abundance of plasma proteins in control and dysbiotic newborn macaques at 7 or 14 days of life, normalized against all subjects and scaled by row. *k*-means clustering was used to arrange subjects and plasma protein abundance. **(E)** Unsupervised analysis of cytometry by time of flight (CyTOF) data for CD4⁺ helper T cells or neutrophils. *t*-SNE (*t*-distributed stochastic neighbor embedding) projection of indicated functional markers in CD4⁺ helper T cells (top) or neutrophils (bottom) in peripheral blood of control and dysbiotic newborn macaques at 7 days of life. Pairwise Euclidean distances between CD4⁺ helper T cells (top) or neutrophils (bottom) in peripheral blood of control and dysbiotic newborn macaques at 7 days of life (*n* = 8; four in each experimental group; **P* < 0.05, Student's *t* test). Solid lines, median; dotted lines, quartiles. **(F)** Frequencies of the indicated immune cell types in peripheral blood of control and dysbiotic newborn macaques at 7 days of life (*n* = 8; four in each experimental group; *P* < 0.05, one-way ANOVA with Tukey's correction for multiple comparisons). Solid lines, median; dotted lines, quartiles.

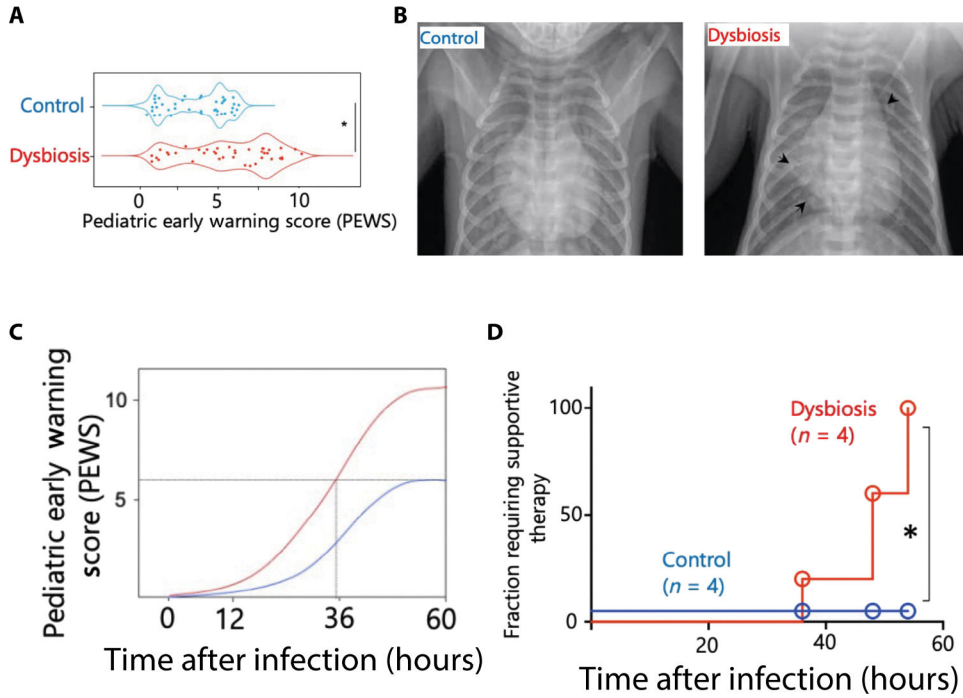


Fig. 2. Antibiotic exposure during the first week of life is associated with increased susceptibility to pneumonia.

(A) Pediatric early warning score (PEWS) after infection with *Streptococcus pneumoniae* in control (blue) or dysbiotic (red) newborn macaques. (B) Representative chest radiographs obtained at euthanasia in control and dysbiotic newborn macaques. Arrows indicate areas of consolidation. (C) Progression of PEWS after infection with *S. pneumoniae* in control (blue) or dysbiotic (red) newborn macaques. Lines represent the best-fit curve by the smoothed spline of the longitudinal distribution of PEWS from the start of infection. Broken lines represent time (after infection) to PEWS > 8, a predetermined threshold to initiate supportive therapy. (D) Kaplan-Meier plot of the fraction of control and dysbiotic newborn macaques requiring supportive therapy at indicated times after infection ($n = 8$; four in each experimental group; $*P < 0.05$, Mantel-Cox log-rank test).

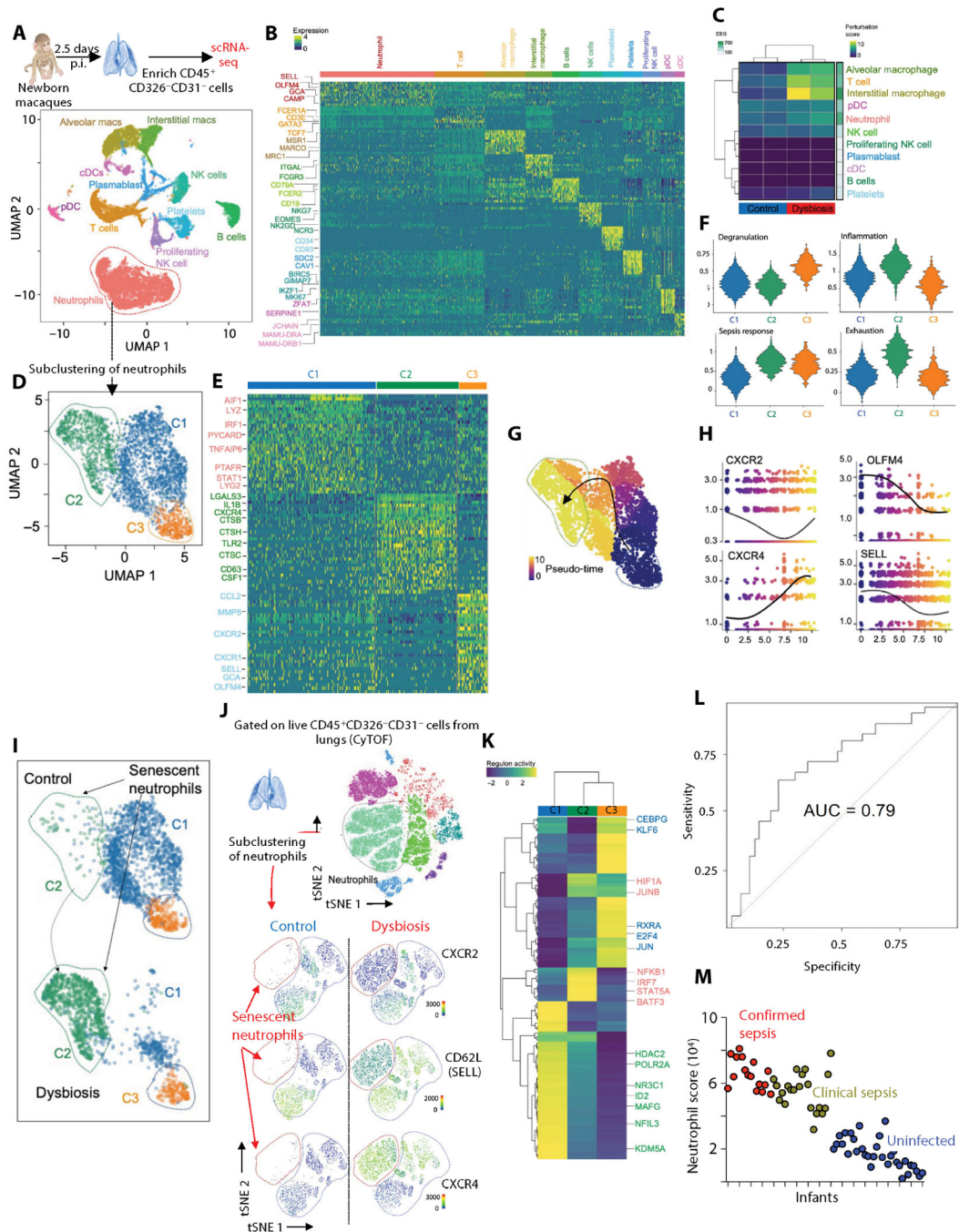


Fig. 3. Antibiotic exposure for the first week remodels the pulmonary immune transcriptome during response to respiratory pathogens.

(A) Lung from control and dysbiotic newborn macaques ($n = 4$; two in each group) was obtained at 60 hours after infection. p.i., post infection. Lung samples were dissociated into cell suspensions, enriched for immune cells (EPCAM⁻CD31⁻CD45⁺), and used for single-cell RNA sequencing (scRNA-seq). Uniform manifold approximation and projection (UMAP) embedding of all samples ($n = 13,377$ cells) colored by cell clusters was performed on scRNA-seq data of these pulmonary immune cells. pDC, plasmacytoid dendritic cells;

cDC, classical dendritic cells. **(B)** Row-scaled expression of the highest differentially expressed genes (DEGs) in each cluster (Bonferroni-adjusted $P < 0.05$). pDC, plasmacytoid dendritic cells; cDC, classical dendritic cells. **(C)** Cellular perturbation scores in indicated samples. The number of DEGs between dysbiotic and control newborn macaques for each cell type is indicated on the right. **(D)** UMAP embedding of neutrophils ($n = 4768$) extracted from a larger dataset of lung immune cells and reclustered into three distinct clusters. **(E)** Row-scaled expression of the highest DEGs in each cluster (Bonferroni-adjusted $P < 0.05$). **(F)** Module scores for each cluster of neutrophils showing enrichment of genes related to degranulation, inflammation, sepsis, and exhaustion. **(G)** UMAP embedding of neutrophils colored by pseudo-time with overlaid trajectory and **(H)** scatterplots showing expression of selected cluster-defining genes across pseudo-time. **(I)** UMAP embedding of neutrophils colored by cluster in control and dysbiotic newborn macaques indicating the emergence of the senescent C2 cluster in dysbiotic newborns. **(J)** CyTOF. *t*-SNE embedding of neutrophils extracted from a larger dataset of lung immune cells (top). Expression of key phenotypic markers (CXCR2, CD62L, and CXCR4) is shown (bottom). Neutrophil cluster coexpressing CXCR4 and CD62L is absent in control newborn macaques. **(K)** Row-scaled regulon activity for neutrophil clusters. *k*-means clustering was used to arrange clusters and regulons ($n = 4$; two in each experimental group; Benjamini and Hochberg-adjusted $P < 0.01$). **(L)** Receiver operating characteristic curve depicting sensitivity and specificity of sepsis diagnosis using the gene signature of senescent, hyperinflammatory neutrophils (*HIF1A*, *CXCR4*, *CD274*, *LTF*, and *S100A8*) in an independent cohort of 69 at-risk infants. AUC, area under the curve. **(M)** Scatterplot representing the ranked senescent neutrophil signature score (aggregated expression of *HIF1A*, *CXCR4*, *CD274*, *LTF*, and *S100A8*; see Materials and Methods) for each infant sample in bulk transcriptomic dataset (see Materials and Methods), colored by clinical diagnosis.

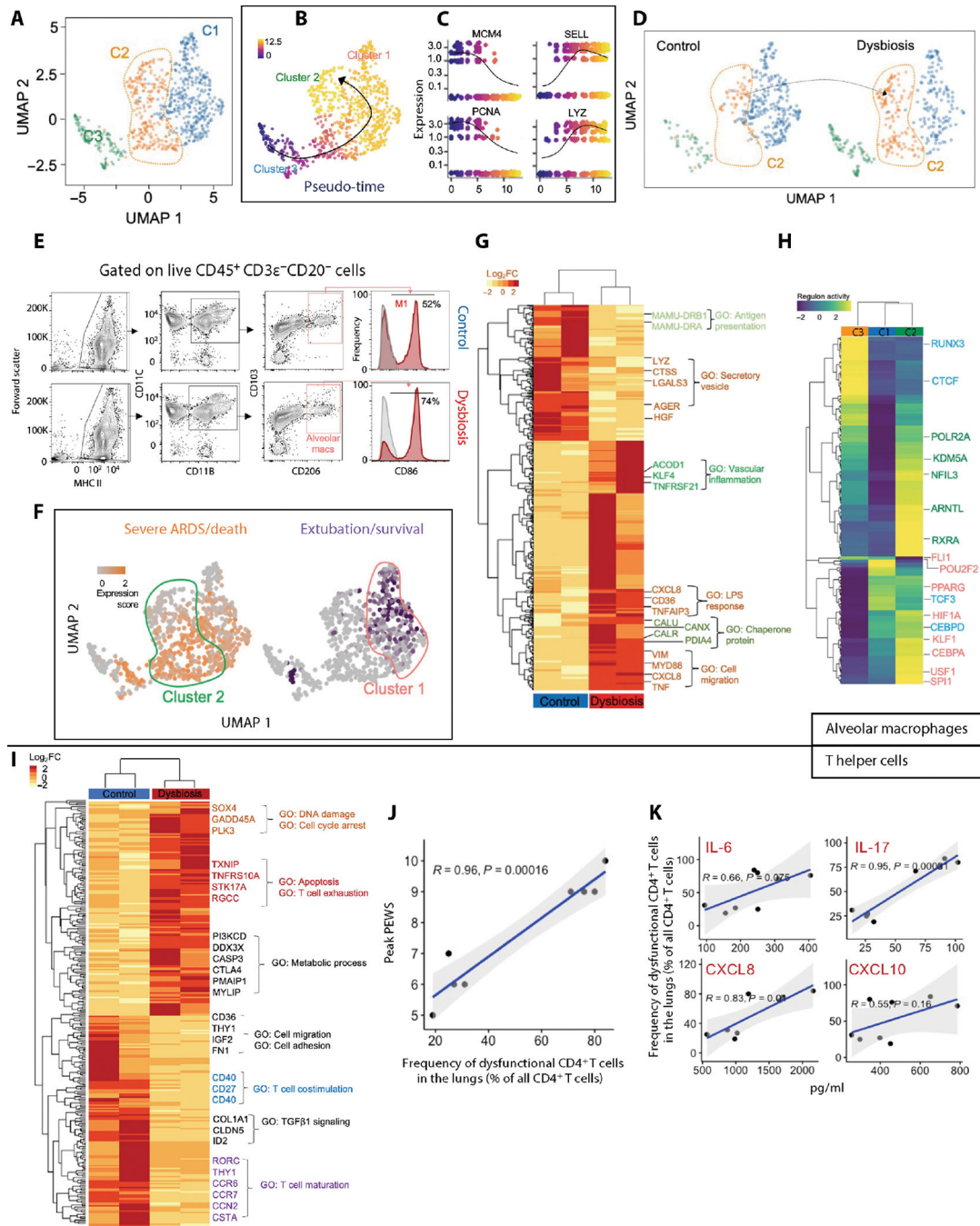


Fig. 4. Antibiotic exposure during the first week remodels the pulmonary macrophages and helper T cell compartments.

(A) UMAP embedding of AMs extracted from a larger dataset of lung immune cells colored by clusters. (B) UMAP embedding of AMs colored by pseudo-time with overlaid trajectory and (C) scatterplots showing expression of selected cluster-defining genes across pseudo-time. (D) UMAP embedding of AMs split by control and dysbiotic macaques showing an emergence of a unique dysbiotic cluster (cluster 2). (E) Flow cytometry. Bivariate contour plots showing gating strategy to identify macrophage subsets and histograms

showing coexpression of activation markers, CD86 and CD206, on AMs from control (top) and dysbiotic (bottom) newborn macaques. Numbers indicate the relative frequencies of M1-activated macrophage subset ($n = 8$; four in each experimental group). **(F)** UMAP embedding of AMs colored by average expression of genes associated with severe ARDS/death (orange) and survival/extubation (purple). These gene signatures were derived from public gene expression datasets from monocytes in pediatric bacterial sepsis subjects (see Materials and Methods). **(G)** Row-scaled expression of DEGs in AMs from control and dysbiotic newborn macaques ($n = 4$; two in each experimental group), normalized against all subjects. k -means clustering was used to arrange subjects and transcripts ($n = 4$; two in each experimental group; Benjamini and Hochberg-adjusted $P < 0.01$, \log_2 fold change (FC) > 2 , Wald's test]. GO, Gene Ontology. **(H)** Row-scaled regulon activity for AM clusters. k -means clustering was used to arrange clusters and regulons ($n = 4$; two in each experimental group; Benjamini and Hochberg-adjusted $P < 0.01$). **(I)** Row-scaled expression of DEGs in the pulmonary T cells from control and dysbiotic newborn macaques ($n = 4$; two in each experimental group), normalized against all subjects. k -means clustering was used to arrange subjects and transcripts ($n = 4$; two in each experimental group; Benjamini and Hochberg-adjusted $P < 0.01$, \log_2 fold change > 2 , Wald's test). **(J)** Pearson correlation between peak PEWS and the frequency of dysfunctional (LAG-3⁺) CD4⁺ T cells (percentage of all CD4⁺ T cells from lungs). Correlation coefficient (R) and significance with the associated P value is indicated. **(K)** Pearson correlation between indicated cytokines from bronchial washings (pg/ml) and the frequency of dysfunctional CD4⁺ T cells (% of all CD4⁺ T cells from lungs). Correlation coefficient (R^2) and significance with the associated P value is indicated.

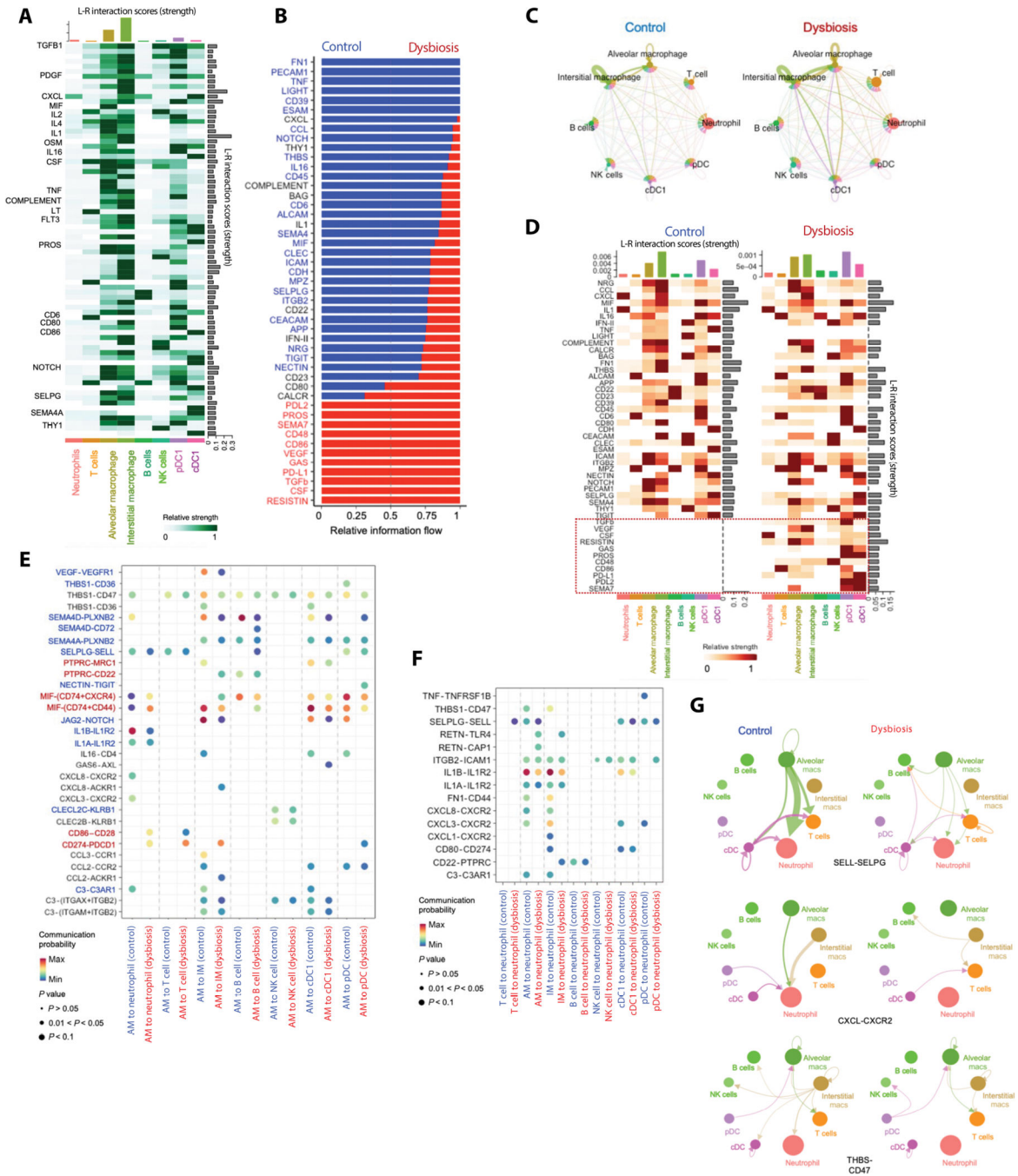


Fig. 5. Changes in communication circuits between neutrophils and macrophages underlie a dysfunctional remodeling of the pulmonary myeloid compartment. (A) Cellcell communication network between different pulmonary immune cells. Bar graphs at the top indicate ligand-receptor interaction scores (strength) for each indicated cell type. Bar graphs on the right show the ligand-receptor interaction scores (strength) of each ligand-receptor interaction. The network is dominated by pathways related to inflammation, chemotaxis, and tissue repair, as indicated by selected signal transcripts (on the left). (B) Cell-cell communication pathways ranked by overall information flow in control or dysbiotic newborn macaques. Cell-cell communications enriched in control exposed

newborn macaques (in blue text) are dominated by pathways related to tissue homeostasis. Pathways related to chemotaxis (in black text) are equally enriched in control or dysbiotic newborn macaques. Cell-cell communications increased in dysbiotic newborn macaques (in red text) are dominated by pathways related to inflammation. **(C)** Circle plot showing differential number of interactions in the cell-cell communication network between control (left) and dysbiotic (right) newborn macaques. Macrophages and dendritic cells are the hubs (senders), whereas neutrophils and T cells are targets (receivers) of cell-cell communication networks. **(D)** Communication pathways related to cell exhaustion and cell activation (boxed) are abundant in dysbiotic compared to control macaques. Bar graphs at the top indicate ligand-receptor interaction scores (strength) for each indicated cell type. Bar graphs on the right show the ligand-receptor interaction scores (strength) of each ligand-receptor interaction. **(E)** Dot plot of outgoing signaling patterns from AM (sender) to other immune cells in control (blue) or dysbiotic macaques (red). Dot color reflects communication probabilities and dot size represents computed *P* values. Empty space means that the communication probability is zero (*P* values calculated from one-sided permutation test). **(F)** Dot plot of incoming signaling patterns to neutrophils (receiver) from other immune cells in control (blue) or dysbiotic macaques (red) (*P* values computed from one-sided permutation test). **(G)** Autocrine and paracrine signaling pathways related to neutrophil migration (CXCL-CXCR2 and THBS1-CD47) and neutrophil activation (SELPLG-SELL) in control or dysbiotic macaques. Circle sizes are proportional to the number of cells in each cell group, and edge width represents the strength of cell-cell communication.

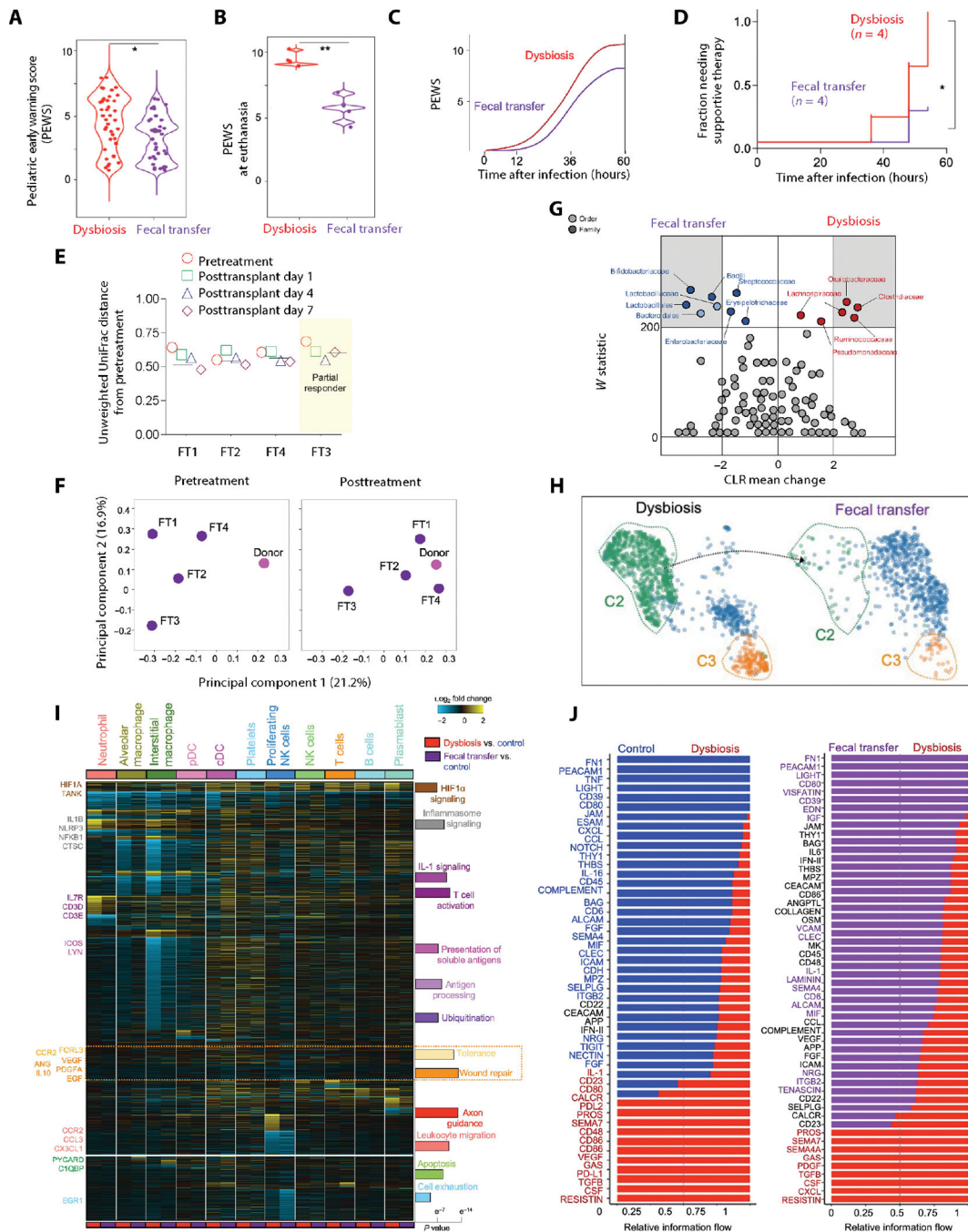


Fig. 6. Fecal transfer was associated with favorable changes in pulmonary immune cell responses and improved host resistance to pneumonia in dysbiotic macaques.

(A) ABX-exposed (dysbiosis) or fecal transfer (FT) recipient newborn macaques were challenged with *S. pneumoniae* (serotype 19F) on PN14. PEWS was determined every 6 hours ($n = 8$; four in each experimental group; the P value is indicated, Student's t test). (B) PEWS at euthanasia ($n = 8$; four in each experimental group; the P value is indicated, Student's t test). (C) Progression of PEWS after infection in dysbiotic or FT recipient newborn macaques. Broken lines (C) represent time (after infection) to PEWS

> 8, a predetermined threshold to initiate supportive therapy. **(D)** Kaplan-Meier plot of the fraction of dysbiotic or FT recipient newborn macaques requiring supportive therapy at indicated times after infection ($n = 8$; four in each experimental group; $*P < 0.05$, Mantel-Cox log-rank test). **(E)** Plot of unweighted UniFrac distances for each FT recipient from the corresponding pretreatment sample at 1, 4, or 7 days after transfer. 0 represents identical microbiota compositions, and 1 represents completely dissimilar compositions. The horizontal line represents the average posttreatment distance. **(F)** Principal component analysis (PCA) of fecal bacterial communities of the donor (pink) and the recipients (purple) before (pretreatment) or 7 days after FT (posttreatment), based on β -diversity (unweighted UniFrac). The distance between samples on the plot represents their dissimilarity. **(G)** Relative abundance of specific taxa in the recipients before (pretreatment) or 7 days after FT (posttreatment). Differentially abundant taxa (FDR $q = 0.05$, center log transformation > 2) are presented in gray margins. The center log transformation mean difference represents compositional differences in microbial communities. **(H)** UMAP embedding of pulmonary neutrophils colored by cluster in dysbiotic and FT recipient newborn macaques, showing the reappearance of cluster 1. **(I)** Heatmap of DEGs in all pairwise cell type comparisons (fold > 1.2 and eBayes t test $P < 0.05$, FDR corrected) in dysbiosis or FT versus control (cellHarmony). Bar plot denotes the Fisher's exact P values (FDR corrected) of GO terms adjacent to the enriched cellHarmony DEG cluster. **(J)** Cell-cell communication pathways ranked by overall information flow in dysbiotic (red text) or FT recipient newborn macaques (purple text). Miscommunication in pathways related to inflammation, immune costimulation, and cell exhaustion is reversed in FT recipient newborn macaques. Cell-cell communication associated with tissue repair and cell migration remains uncorrected after FT.