It is now well accepted that the cells lining the airways constitute more than just a barrier between the external environment and the underlying mesenchyme. This collection of specialized epithelial cells responds to microbes and noxious stimuli that overcome the mucociliary barrier and are a vital component of host defence, interacting with cells of the immune system to maintain homeostasis while facilitating immune reactions when necessary. The respiratory epithelium must also manage responses to the diverse toxins contained within the inhaled environment and there is emerging evidence that epithelial dysfunction is a driver of numerous chronic diseases affecting the lungs. The traditional view of the epithelial layer incorporates basal cells in close proximity to secretory and ciliated cells, forming a tight unit that maintains a physical barrier but is also responsive to the inhaled environment via cells and molecules from the immune system. However, with the advent of advanced sequencing techniques, this view has changed to one of a dynamic cellular structure encompassing a wide range of highly specialized cells that are able to respond to environmental change, interact with resident microbial communities and cooperate with multiple other specialized cellular systems such as the immune and neural systems.

The respiratory tract is a complex organ system divided into the upper respiratory tract, that includes the nasal cavity, pharynx and larynx, and the lower respiratory tract comprising the conducting airways (trachea, bronchi and bronchioles) and the respiratory zone (respiratory bronchioles and alveoli) (Fig. 1). Each area has a specific function and the regional differences in cellular composition reflect this. Specialized epithelial cell populations line the entire respiratory tract from the nasal cavity to the alveoli. Elegant electron microscopy studies provided early insight into the morphology and ultrastructure of the principle epithelial cell types residing in the human airways. In addition to using classical electron microscopy-defined morphological features, standard immunohistochemical staining for cell type-specific markers has been used to characterize and quantify epithelial cell populations throughout the human respiratory tract, determining the influence of anatomical location on cellular composition. The airways are lined by ciliated and secretory cells primarily adapted to facilitate mucociliary clearance of particulate matter and infectious pathogens in the air we breathe. Like other mucosal surfaces, the airway epithelium is at an interface with the environment and is therefore critically important to host defence. Despite sharing the same embryological origin as the gut mucosa, immunological activity at the airway mucosal surface is necessarily distinct and shaped by differences in environmental conditions (temperature gradient, bidirectional airflow), resident microbial communities and airborne antigens. Although not the focus of this Review, the epithelial cells lining the distal alveolar region of the lung are phenotypically and functionally distinct; alveolar epithelial type 1 (AT1) cells provide a specialized surface for gas exchange and type 2 (AT2) cells secrete pulmonary surfactant to prevent alveolar collapse during expiration. In each anatomical niche, there are progenitor cell populations, for example, basal cells in the airways and AT2 cells in the alveoli, that ensure robust epithelial regeneration under homeostatic conditions and following injury.

Regulation of immune responses by the airway epithelial cell landscape

Richard J. Hewitt and Clare M. Lloyd

Abstract | The community of cells lining our airways plays a collaborative role in the preservation of immune homeostasis in the lung and provides protection from the pathogens and pollutants in the air we breathe. In addition to its structural attributes that provide effective mucociliary clearance of the lower airspace, the airway epithelium is an immunologically active barrier surface that senses changes in the airway environment and interacts with resident and recruited immune cells. Single-cell RNA-sequencing is illuminating the cellular heterogeneity that exists in the airway wall and has identified novel cell populations with unique molecular signatures, trajectories of differentiation and diverse functions in health and disease. In this Review, we discuss how our view of the airway epithelial landscape has evolved with the advent of transcriptomic approaches to cellular phenotyping, with a focus on epithelial interactions with the local neuronal and immune systems.
The advent of novel sequencing techniques has facilitated not only the identification of novel cell types but reveals potential functions of previously named but poorly understood cell types, for example, tuft/brush cells. There is also an indication that heterogenous cell types and states exist and that subtle changes in these will be influenced by the changing environment, for example, by smoking or exposure to allergens or pollutants. It is now clear that the airway wall represents a dynamic ‘community’ of epithelial cells existing in close association with resident immune and neuronal cells to generate an integrated unit that plays a critical role in maintaining mucosal immune homeostasis as well as facilitating host defence against inhaled pathogens. In this Review, we outline how single-cell transcriptomic techniques have been used to map the airway epithelial landscape and how this has highlighted the function of novel specialized epithelial cells and how they interact with cells of the immune and neuronal systems to regulate airway immunity.

Mapping the airway epithelial landscape

Single-cell RNA-sequencing (scRNA-seq) has transformed our view of the airway epithelium, unveiling a level of cellular diversity that had not been documented using microscopy for phenotyping\(^1\). This technology profiles the transcriptome of individual cells and therefore facilitates an unbiased characterization of different cell subsets in a heterogenous cell population, driving the discovery of unidentified cell types and states\(^2-5\). The Human Cell Atlas Consortium aims to comprehensively chart — at single-cell resolution with integration of spatial data — the changes that occur in lung cell composition and molecular phenotype in health and disease\(^6\). Added to this, it is also possible to computationally interrogate epithelial–immune interactions in the airway niche using known ligand–receptor interactions\(^16,15-17\).

Studying human epithelial biology using single-cell transcriptomics relies on the availability of lung tissue. One of the earliest studies in humans analysed diseased explant parenchymal lung tissue from patients with idiopathic pulmonary fibrosis (IPF) and lung lobes from donors unsuitable for transplant as control tissue\(^18\). For studies of the human airway epithelium, the acquisition of bronchial brushings and endobronchial biopsy samples in the setting of disease can be achieved through fibre-optic bronchoscopy. Samples in the setting of disease can be achieved through fibre-optic bronchoscopy and endobronchial biopsy samples in the context of routine clinical care\(^19,20\). It is important to note that the cellular content differs by the sampling method used and this will influence the cell clusters obtained by scRNA-seq (FIG. 2). Acquiring age-matched, healthy control samples is more challenging as it necessitates research volunteers undergoing an invasive procedure. The procurement of appropriate human lung tissue is only feasible through close collaboration between clinicians and scientists and through human tissue biobanks. Importantly, logistical
Bronchial brushings
A procedure in which a brush is used through a bronchoscope to obtain cells lining the airway under direct visualization.

Endobronchial biopsy
A procedure in which a small biopsy is performed using forceps through a bronchoscope to obtain a sample of airway wall cells under direct visualization.

Air–liquid interface cultures
Epithelial cells are cultured on a microporous membrane in a Transwell plate and exposed to air to allow mucociliary differentiation.

practicalities require careful consideration because the hypothermic preservation of tissue and the minimization of time delays between fresh sample collection in the clinic and processing in the laboratory are critical to ensuring high-quality scRNA-seq data. A cold ischaemic time of 72h results in an increase in the presence of mitochondrial reads due to cellular stress or death in multiple lung cell types and a cell-specific decrease in the proportion of CD4+ and CD8+ cytotoxic T cells. The enzymatic dissociation of tissue with collagenase is widely used to isolate cells for scRNA-seq studies but, as a caveat, it can induce the expression of immediate early genes, including Fos and Jun, and heat shock protein (HSP) genes in a subpopulation of cells, thus also potentially introducing an important artefact and potential bias in single-cell data sets.

During scRNA-seq analysis, cells sharing similar gene expression profiles are clustered together and can be assigned an annotation based on known marker genes (Table 1). In the first single-cell studies of the mammalian airway, a new cluster of cells was identified that did not map to any of the previously known epithelial cell types and, based on its gene expression signature, was termed the ‘pulmonary ionocyte’. This sentinel discovery of a novel cell type highlights the power of scRNA-seq.

Sub-clustering can be used to identify diverse cell states within a specific cell-type cluster. A single-cell analysis of human nasal epithelial cells differentiated in air-liquid interface cultures demonstrated a precursor subset of FOXJ1-expressing multiciliated cells termed ‘deuterosomal cells’, which express specific genes, including DEUP1 (also known as CCDC67), FOXN4 and CDC20B, all of which are critical for centriole amplification and ciliogenesis. Deuterosomal cells are identifiable in the healthy human airways by scRNA-seq.

Subtle but distinct molecular states of basal, goblet and ciliated cells denoted by differential marker gene expression have now been described in the human airway. Goblet cells could be divided into two subsets according to their uniquely expressed genes; for example, ‘goblet 2’ cells in the nasal epithelium have higher expression of genes involved in immune cell recruitment such as CXCL10, IL19 and CSE3 (REF. 19). The anatomical origin of the cell population within the respiratory tract plays an important role in governing its transcriptional signature and cell state. A single-cell study of brushings and biopsies taken from four distinct anatomical

**Sample types obtained for scRNA-seq airway studies**

**Fig. 2 | Single-cell transcriptomic studies of the airway.** For human single-cell RNA sequencing (scRNA-seq) studies of the lower respiratory tract, a flexible bronchoscopy is performed to obtain bronchoalveolar lavage (BAL) fluid, airway wall brushings and endobronchial biopsies. The starting material determines the relative contributions of airway immune cells, epithelial cells and underlying stromal cells that are sequenced. A viable, single-cell suspension is obtained by enzymatic dissociation of brushings and biopsies. Bioinformatic pipelines generate distinct cell clusters by dimensional reduction and visualization techniques such as uniform manifold approximation and projection (UMAP) and facilitate assignment of cluster annotations based on cell-marker genes. Novel cell clusters and cell states can be determined in health and disease and trajectory analysis can be used to study dynamic, differentiating cell types.
Asthma

Lung disease characterized by chronic airway inflammation and remodeling, causing variable expiratory airflow obstruction and airway hyper-responsiveness to a range of stimuli. Often associated with allergen sensitization.

locations in healthy subjects, highlighted region-specific sub-clusters of suprabasal, secretory and ciliated cells between nasal and tracheobronchial compartments14–19. Spatial variations in gene expression profiles between cells of the same type relate, in part, to the biological function required at a specific airway location. Secretory cells in the nasal epithelium are enriched in gene sets associated with cell motility, differentiation and sensory perception pathways, whereas in the bronchial tree, they are enriched in genes related to innate immunity and wound healing response pathways19. Noteworthy is the inconsistent annotation of secretory cell types, including club cells and goblet cells, between single-cell studies due to overlapping gene expression signatures19,20,22–24,30,67.

A further determinant of airway epithelial cell state is revealed by the comparison of cells from healthy volunteers versus those collected from patients with respiratory disease. An analysis of airway wall biopsies from mild-to-moderate asthmatics exposed several cell states not found in the healthy airway19. Cells co-expressing gene markers of ciliated cells, including FOXJ1, and goblet cell genes such as MUC5AC, were termed ‘mucous ciliated cells’ and are postulated to represent a novel transitional cell state driven by IL-4/IL-13 signalling in asthma.

Single-cell analyses of parenchymal lung tissue from patients with IPF, a progressive scarring lung disease, have reported a previously unidentified, transcriptionally distinct cluster of cells in the fibrotic distal lung that express some but not all markers of airway basal cells27,28. These intriguing ‘aberrant basaloid’ cells, which are KRT5–KRT17+, co-express genes encoding mesenchymal markers such as collagen type 1 a1 chain (COL1A1), the transcription factor SOX9 (which plays a role in distal airway development) and genes linked to IPF pathogenesis, including MMP7, which encodes matrilysin.

The pulmonary epithelium represents a dynamic cellular community and single-cell transcriptomics captures a unique snapshot of cells at different stages in the continuous process of differentiation — a feature that was first harnessed to elucidate epithelial lineage hierarchies at four distinct stages of alveolar development in mice29. With the refinement of scRNA-seq bioinformatic approaches it has been possible to study cell fate in human samples and to unveil the subtleties of human airway epithelial cell differentiation15,20,23,30. The transcriptional changes that accompany the differentiation of each cell can be exploited computationally by trajectory inference algorithms that order cells according to progress along their individual trajectories, which is quantified by ‘pseudotime’31,32. A pseudotime analysis of nasal airway epithelial cells in air–liquid interface cultures demonstrated that goblet cells could act as precursors to ciliated cells and that transitory FOXJ1+ MUC5AC+ cells could be found in the context of this differentiation trajectory31. Furthermore, leveraging the relative abundance of unspliced and spliced mRNA per gene, it is possible to estimate the rate of gene expression change, termed ‘RNA velocity’, and to predict future cell states15,24.

Pseudotemporal ordering of cells using a diffusion map approach31 identified an intriguing population of Krt13-expressing cells in the mouse airway as an intermediary in basal to club cell differentiation31. This transitional cell was arranged in discrete, stratified, high-turnover structures termed ‘hillocks’ and expressed
Many unanswered questions for the research community.

The comprehensive transcriptomic information charted at a cellular level by scRNA-seq has heightened our knowledge of the airway epithelial landscape at steady state. This technology has also been harnessed to study airway samples from critically ill patients with COVID-19 in order to provide finer grain insight into the pathogenesis of severe acute respiratory system coronavirus 2 (SARS-CoV-2) infection.6,36,37. Studying single-cell gene expression in multiple tissue types from data sets collated from the Human Cell Atlas Consortium, genes associated with SARS-CoV-2 cell entry, namely ACE2, which encodes a viral entry receptor, and TMPRSS2, were significantly enriched in nasal secretory and ciliated epithelial cells. Noteworthy was the high ACE2 expression in cells at other mucosal surfaces, including the cornea, conjunctiva, small intestine and colon, which may, in combination with high nasal expression, explain the higher transmissibility of SARS-CoV-2.6,36 This demonstrates how existing data sets can be exploited to further our understanding of the pathogenesis of a novel emerging disease.

Functions of rare epithelial cell types

Basal cells are multipotent stem cells that account for a third of all airway epithelial cells and give rise to other major subpopulations, such as secretory and ciliated cells, during homeostatic maintenance of the epithelial barrier and regeneration in response to injury.28. Basal cells are anchored to the basement membrane of the conducting airways and express keratin 5 (KRT5) and transcription factor tumour protein p63 (TP63). Lineage tracing and single-cell transcriptomic studies have demonstrated that functionally distinct basal cell subsets exist in the airway.24,26. Two discrete basal cell states are present in the upper and lower airways, but their frequency is lower in the upper airways. These cell states reflect distinct differentiation stages in basal cells, with mature basal cells (located at a more apical region) expressing lower levels of TP63 and NPPC than the less mature basal cells.20. Basal cells with a lower TP63 expression and higher KRT19 and NOTCH3 expression were annotated as ‘suprabasal’ cells by Deprez et al.15.

In addition to these major cell types, rarer cell populations comprising less than 1% of all human airway epithelial cells have been identified via detailed single-cell transcriptomic analyses.16,22 Although uncommon, these cells seem to have very distinct phenotypes and perform specific tasks that are vital for the efficient functioning and maintenance of the epithelial landscape.

**Hillock cells**. The advent of scRNA-seq has enabled investigators to examine cell development trajectories. In addition to the connection between basal, club and ciliated cells, Montoro et al. defined a distinct trajectory, which links tracheal basal and club cells via a novel transitional cell that uniquely expresses Krt13 and Krt4 (Ref. 3). By looking at the location of these cells within the lungs coupled with lineage tracing in mice, the authors noted that Krt13+ cells lie within contiguous groups of stratified cells that do not contain luminal ciliated cells. They termed these groupings ‘hillocks’, a structure composed of luminal Scgb1a1+Krt13+ cells lying on top of Trp63+Krt13+ basal cells, with the expression of Trp63 occurring along a gradient from basal to suprabasal divisions. Critically, hillocks contain a particularly high number of cycling cells and expressed markers of cellular adhesion and epithelial differentiation as well as genes associated with barrier function and immunomodulation.41. A parallel study by Plasschaert et al. confirmed the presence of a distinct population of Krt13+Krt4+ cells in the murine trachea and in ex vivo differentiated human basal epithelial cells, although they concluded that this population represented an intermediate population between basal stem cells and differentiated luminal secretory cells.22. Furthermore, Krt13+Krt4+ expressing cells emerged in the tracheal epithelium 1 day after polidocanol-induced injury in mice.42. In humans, KRT13 is expressed in the airway epithelium by a subpopulation of ‘cycling’ basal cells that are characterized by the expression of genes associated with cell proliferation, including MKI67 (which encodes Ki67)10. Further analyses are needed to confirm whether hillocks represent a truly distinct niche rather than a metaplastic zone in transition and to determine their origin and purpose. Indeed, it is not clear whether these structures are present throughout the airways or are restricted to the trachea. In support of their existence, Deprez et al. describe a population of KRT13+ cells, a high proportion of which were cycling, in nasal

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**Box 1 | Unanswered questions**

The field of single-cell transcriptomics has provided a transformative, high-definition view of the cellular communities that exist in the human airway but has also generated many unanswered questions for the research community.

- **How do cells in the contemporary single-cell landscape reflect those detected by traditional microscopy methods?** Data from single-cell RNA sequencing has refined our characterization of historically described airway cell types, such as tuft cells, and has led to the discovery of a previously undescribed cell — the ‘ionocyte’.6,22,23. Electron microscopy studies in 1994 reported ‘cells of indeterminate type’5 and there remain our characterization of historically described airway cell types, such as tuft cells4, and traditional microscopy methods? Data from single-cell RNA sequencing has refined how existing data sets can be exploited to further our understanding of the pathogenesis of a novel emerging disease.

- **How much significance should we place on the discovery of clusters with low cell abundance or sub-clusters of a specific cell type?** In addition to confirming the existence of a specific cell type through immunofluorescence microscopy, to truly bring the high expression in cells at other mucosal surfaces, including the cornea, conjunctiva, small intestine and colon, which may, in combination with high nasal expression, explain the higher transmissibility of SARS-CoV-2.6,36 This demonstrates how existing data sets can be exploited to further our understanding of the pathogenesis of a novel emerging disease.

- **How many unanswered questions for the research community?**

- **How do we account for the inherent biological variability between human samples and cell clusters that arise in a minority of study subjects?** One approach to provide greater transparency is to show cellular composition and clusters for individual subjects as well as those integrated by disease.6,39,40.

- **How do we reconcile differences in the assignment of cell identification?** Consensus must be reached on specific positive and negative marker genes that define a particular cell type because there are discrepancies in the annotation of cell types and subsets6,39,36,40. Consistency and clarity on the anatomical origin of the airway cells should also be factored into any new classification system.

- **Cell fate mapping has traditionally relied on the use of reporter mice, but now various computational tools allow the inference of cell trajectory from human single-cell data.41. Differences have been reported in lineage relationships between basal cells and rare cell types (tuft cells, pulmonary neuroendocrine cells and ionocytes), which may reflect the species studied (mouse versus human), model and/or trajectory algorithm used41,42 but also highlights the need for further validation in human samples.**
The emphasis of this Review is the human airway epithelial landscape. However, much of our understanding of the function of epithelial cells and their interaction with immune cells comes from experimental studies in laboratory mice. It is for this reason that we highlight salient proximal–distal variations in cellular composition and structural differences between the murine and human airway.

In mice, basal cells are largely restricted to the trachea and proximal bronchi yet, in humans, basal cells extend throughout the conducting airways. In humans, mucus-producing goblet cells are the most abundant secretory cell type in the proximal airways yet, in adult mice, these are comparatively rare and instead club cells predominate. Submucosal glands lie below the luminal surface and secrete mucus and other antimicrobial factors into the airway but also act as a stem cell niche containing myoepithelial basal cells. In mice, submucosal glands are localized to the larynx and the proximal trachea but, in humans, they are distributed throughout the cartilaginous airways.

The pseudostratified ciliated epithelium of the murine trachea is akin to that found throughout the human airways. Therefore, it has been suggested that this may represent a more relevant region to study for translatable human airway epithelial cell biology. In support of this concept, some of the single-cell findings from murine tracheal epithelial cells, including the discovery of the pulmonary ionocyte, have been recapitulated in studies of the human airway.

Beyond the trachea, the murine airway epithelium transitions to a simple columnar epithelium comprising ciliated, secretory (club) and clusters of pulmonary neuroendocrine cells, thus modelling the composition of only the most distal human airways.

Mouse models are an indispensable tool for biomedical research and have revolutionized our understanding of immunology. However, it is noteworthy that important differences exist between the mouse and human immune system. Differences in the living environment of mice, specifically microbial exposure history, have a profound influence on their immune cell repertoire. For example, in contrast to adult humans or pet-store mice, laboratory mice kept in specific-pathogen free facilities lack differentiated memory CD8+ T cell subsets. These critical environmental differences profoundly affect the development of the pulmonary immune system and thus the interactions with the local epithelial landscape.

Tuft cells. The respiratory tract contains several groups of chemosensory epithelial cells that coordinate interactions with the external environment — the enteroendocrine cells and the tuft cells (also known as ‘brush’ cells). These chemosensory epithelial cells share similarities with taste cells and are predicted to evoke both positive and negative responses from immune and neuronal cells. They have a distinctive morphology, being bottle shaped with apical microvilli, and are expressed in a range of organs, including the gut and airways. Their existence is well documented in the gut, where they are thought to be triggered by dietary metabolites such as succinate and bacterial products (for example, quorum-sensing lactones or D-amino acids). By contrast, their role in the lung is less certain, but they have been detected in the nose, trachea and proximal airways and exist in close contact with nerve fibres, mediating communication between neuronal and immune pathways. However, they are not universally identifiable; indeed, Vieira Braga et al. could not identify a unique population of tuft cells in a scRNA-seq analysis of the human airways using both bronchial brushing and endobronchial biopsy. This likely reflects the rarity of these cells and perhaps the methods used for the collection of airway material.

Tuft cells express a range of indicative markers, including POU domain, class 2, transcription factor 3 (POU2F3), which is considered the lineage-defining transcription factor for tuft cell specification, as well as transient receptor potential cation channel subfamily M member 5 (TRPM5). scRNA-seq has now identified two terminally differentiated Trpm5+ tuft cell populations; one is positive for Gng13 and is likely to be responsible for ‘taste’ sensing, and the other is positive for Alox5ap, suggesting that it contributes to leukotriene synthesis. Tuft cells are thought to generate cysteinyl leukotrienes via the ATP sensor P2Y2. Although generally present in low numbers, the expansion of tuft cells in the airways occurs following the inhalation of common Aeroallergens via activation of the cysteinyl leukotriene pathway. Endogenously generated leukotriene E4 is sensed via CysLT1R, which regulates the number and function of murine airway brush cells.

Thus, tuft cells are able to respond to a diverse range of signals via the combinatorial expression of different receptors. Tuft cells are thought to promote protective respiratory reflexes, such as sneezing, but can also contribute to apnoea as well as to local neurogenic inflammation of the mucosa. Given their distribution and function, tuft cells have also been referred to as ‘solitary chemosensory cells’, enhancing the chemoresponsive protection provided by the local neuronal system. scRNA-seq analysis shows that mouse tuft cells express high levels of cholinergic and bitter taste signalling transcripts (Tas2r108, Gnat3, Trpm5); Acylcholine is released from tracheal tuft cells following stimulation with either the bitter taste receptor type 2 (TAS2R) ligand denatonium or Pseudomonas quinolone quorum-sensing molecules, increasing mucociliary clearance mediated by TRPM5 and M3 muscarinic acetylcholine receptor (M3R), together with calcium release from ciliated cells. It is postulated that the detection of these quorum-sensing molecules from Gram-negative pathogenic bacteria offers a mechanism by which the epithelium triggers capsaicin-sensitive nerve fibres, releasing calcitonin gene-related peptide (CGRP) and substance P, which promote innate immune cascades and microvascular leak and combat bacterial invasion, thus limiting the population densities capable of forming destructive biofilms.

Solitary chemosensory cells, with tuft cell properties, are found to be the primary epithelial source of IL-25 in patients with chronic rhinosinusitis with nasal polyps, facilitating the activation of group 2 innate lymphoid cells (ILC2s) and the production of IL-13, thereby maintaining the type 2 environment in the upper airways. Once activated, tuft cells can release neurotransmitters (such as acetylcholine), eicosanoids involved in leukotriene and prostaglandin biosynthesis and ATP, and cytokines such as IL-25 and thymic stromal lymphopoietin (TSLP). All of these mediators are implicated in allergic inflammation, being able to elicit ILC2 activation and accumulation via the expression of specific receptors.

Ionocytes. Single-cell transcriptomic studies of the mouse tracheal epithelium and differentiated human tracheal epithelial cells in culture discovered a novel cluster of cells, termed ‘pulmonary ionocytes’ due to the similarity of their gene expression profiles to those of the ionocytes, indicating that hillock cells may exist in other areas of the human respiratory tract.

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transport cell of the mucociliary epithelium of *Xenopus* larval skin;[25,26] As in *Xenopus*, ionocytes express genes encoding subunits of a V-ATPase proton pump, which regulates ion transport and pH. Crucial in determining the ionocyte lineage in mouse and human epithelial cells is FOXI1, which belongs to the forkhead family of transcription factors. Knockout studies in mice and lentiviral studies in human cell cultures show that FOXI1 regulates the expression of the *CFTR* gene that encodes a critical chloride-ion transporter that is defective or absent in cystic fibrosis. Cystic fibrosis is a life-limiting disease characterized by increased mucus viscosity, impaired mucociliary clearance, chronic infection and airway inflammation, leading to loss of lung function[27]. Ionocytes account for only 1–2% of human airway epithelial cells but are more enriched in *CFTR* mRNA than in any other airway cell type. Combining scRNA-seq with conventional in vivo lineage tracing over three different time points in a technique called ‘pulse-seq’, indicated that basal cells were the principal source of ionocytes and other rare cell types[28]. Given the clustering of rare cell types in scRNA-seq data sets, Goldfarbmuren et al. further investigated rare cell lineage relationships using a CRISPR–Cas9 knockout system in human tracheal basal cells to determine that, during differentiation, ionocytes and pulmonary neuroendocrine cells (PNECs) arise from POU2F3 tuft-like cells.[29]

**Pulmonary neuroendocrine cells.** PNECs are solitary cells resident within the surface epithelium of the trachea, bronchi and bronchioles. They can also exist in formations called neuroendocrine bodies in the intrapulmonary airways. PNECs function as chemosensors of the airway and respond to changes in oxygen, stretch and chemical stimuli.[30] They are a rich source of neuropeptides and neurotransmitters that elicit immune and physiological effects. PNECs are the only innervated airway epithelial cell type, are conserved across species and represent just 1% of the total lung epithelial cell population. They are reported to be the earliest specialized cell type that forms in the lung epithelium during development[31–33] and are thought to be particularly important functionally in early life since neonatal mice lacking PNECs are protected from allergic inflammation[34]. In vitro analyses predicted roles for PNECs in a range of activities, including oxygen sensing, the maintenance of bronchial and vascular smooth muscle tone, and in the coordination of immune responses. However, an elegant series of experiments with mutant mice has confirmed the functional characteristics of PNECs that influence a range of pulmonary diseases. A defining characteristic marker of PNECs is the gene *Ascl1*, a transcription factor that is essential for their formation. Mice that lack *Ascl1* die at birth, but a conditional knockout designed to inactivate *Ascl1* in PNEC precursors led to viable mice that completely lack PNECs, as marked by a lack of CGRP+ cells in the airway epithelium.[35] These *Ascl1*-mutant mice were normal at baseline but were protected from developing severe goblet cell hyperplasia and type 2 inflammation when exposed to allergens during the perinatal period. This absence of PNECs was accompanied by a reduction in key neuropeptides, including the neurotransmitter GABA.[36]

The expression of the roundabout (ROBO) genes is vital for the clustering of PNECs into neuroendocrine bodies, limiting immune cell infiltration, and in preventing alveolar simplification during postnatal lung development. Although PNEC defects were apparent in *Rboh*-deficient mice at E15.5, the physiological effects only occurred after birth, indicating that the effects of PNEC are dependent on the physical exposure to air that occurs with the first breath, thus reinforcing the idea that PNECs are sensors of the changing inhaled environment. By contrast, the genetic ablation of PNECs in adult mice had no effect on airway homeostasis or repair following chemical injury.

In early life murine models, PNECs were shown to facilitate mucus hypersecretion during allergen exposure via neurotrophin 4 regulation of PNEC innervation and the secretion of GABA, which promotes Muc5ac expression. PNECs represent the only source of GABA in primate lungs and in ex vivo cultures of human epithelial cells[37,38]. PNECs also reside in close association with ILC2s in the airways and communicate via CGRP to maximize the ILC expression of IL-5 and GABA to elicit mucus production.[39] GABA production was found to be absolutely required for goblet cell hyperplasia but not in type 2 inflammation in both neonatal and adult models of allergic inflammation[40]. Importantly, these findings are reflected in humans, with increased numbers of the CGRP+ subset of PNECs in patients with allergic asthma.

The fact that abnormalities in PNEC numbers are associated with a wide range of pulmonary diseases, including rare genetic disorders such as congenital diaphragmatic hernia and small cell lung cancer as well as more common diseases such as asthma, indicate a key role for these cells in physiological and immune pathways critical for effective functioning of the lung at homeostasis.

**Immune properties of airway epithelial cells**
Antimicrobial mediators and airway mucins, including MUC5AC and MUC5B, are produced by secretory epithelial cells lining the airways and submucosal glands and contribute to the first layer of host defence at the airway epithelial surface.[41] Secretory IgA (SIgA) — produced by sub-epithelial plasma cells and transported to the apical surface of airway epithelial cells via the polymeric immunoglobulin receptor (pIgR) — prevents the adherence of airborne microorganisms in a process called ‘immune exclusion’.[42,43] Epithelial cells are equipped with pattern recognition receptors, such as Toll-like receptors, which rapidly sense and initiate an immune response to microbial threats, and cytokine receptors, including TNFRI, which allow them to respond to signals produced by immune cells such as airway macrophages[44]. Adjacent airway epithelial cells are linked by intracellular tight junctions and adherens junctions, which selectively regulate the paracellular diffusion of ions and molecules and maintain barrier integrity.[45] These tight junctions separate ligands present at the apical epithelial surface, for example, growth factor heregulin, from its ErbB family receptors located at the basolateral surface, so that activation occurs following

GABA γ-aminobutyric acid, an inhibitory neurotransmitter.

**Pattern recognition receptors**
Specialized immune receptors that recognize conserved molecular structures on bacteria and viruses. These include Toll-like receptors.
microenvironment, but they also have an intrinsic capacity for inflammatory memory — a discovery that was made in the context of chronic allergic inflammatory disease. Chronic rhinosinusitis (CRS) is a type 2 immune-mediated disease characterized by inflammation and epithelial dysfunction in the nose and paranasal sinuses, with or without the formation of abnormal tissue outgrowths called polyps. Basal cell hyperplasia is a feature of tissue remodelling in this condition. Single-cell transcriptomics was used to study human tissue samples collected during ethmoid sinus surgery from subjects with CRS with and without polyps as well as nasal scrapings from the inferior turbinate of healthy subjects and from those with CRS and polyps. The most striking disease-related changes to the transcriptomic state were seen in basal cell, differentiating/secreting cells in inflammatory CRS tissue compared to basal cells in healthy nasal mucosa. This suggests that basal cells in CRS are a key player in the development and progression of the disease.
cell and glandular cell populations. In the polyp samples, there was notable basal cell expansion at the expense of epithelial cell diversity. Basal cells upregulated IL-4-responsive and IL-13-responsive genes and displayed a failure to differentiate by pseudotime analysis. A combination of bulk RNA sequencing and transposase-accessible chromatin sequencing (ATAC-seq) showed the upregulation of transcription factors such as KLF5 and AT2F3, known to maintain undifferentiated cell states, corresponding to enriched motifs in sorted polyp basal cells. These findings suggested that intrinsic changes at an epigenetic level may be responsible for the differences observed in polyp basal cell state and may be governed by the local type 2 inflammatory milieu. Bulk RNA sequencing of ex vivo-stimulated basal cells in submersed cultures revealed that IL-4 and IL-13 induced more than 10 times the number of genes in non-polyp basal cells than in polyp basal cells. Additionally, at baseline, the Wnt pathway activator CTNNB1 was expressed in polyp basal cells at levels only achievable in non-polyp basal cells through IL-4 and IL-13 stimulation, suggesting a ‘memory’ of in vivo exposure to type 2 cytokines by polyp basal cells. These data have profound implications for our understanding of allergic diseases given that efforts to treat these diseases are generally focused on manipulating cells of the immune system.

_Circadian rhythm._ For over a decade, it has been recognized that, in addition to neuro-hormonal control of the circadian rhythm by the suprachiasmatic nucleus in the brain, peripheral tissues and their cells, including airway epithelial cells, possess an intrinsic circadian clock responsible for rhythmic immune oscillations with the time of day. In mouse and human lung tissue, the clock gene products CLOCK and PER2 were expressed in CCSP+ club cells. Circadian oscillations, measured by the bioluminescence of PER2 in ex vivo mouse lung slices from transgenic mice expressing a Per2-luciferase fusion gene (Per2-Luc mice), could be ‘reset’ in response to glucocorticoids and were lost with the selective ablation of club cells. A notable time-of-day variation in inflammatory response, in particular the BAL neutrophil count, was observed in response to an aerosolized lipopolysaccharide (LPS) challenge in C57BL/6 mice. The targeted deletion of the clock gene _Bmal1_ in club cells using _Cesp-Bmal1^−/−_ mice augmented the duration of the neutrophil inflammatory response to LPS challenge as well as to _Streptococcus pneumoniae_ across the circadian cycle but without an increase in myeloperoxidase activity or a reduction in bacterial load. Driving this pulmonary inflammatory response was the neutrophil chemoattractant CXCL5, which lost circadian regulation in the _Bmal1_ knockout.

REV-ERBα is a nuclear receptor that sits in a negative feedback loop downstream of core transcriptional activators of the molecular clock, namely CLOCK and BMAL1 (REF 7). REV-ERBα plays a key role in the circadian modulation of innate immune responses in myeloid cells and airway epithelial cells. The targeted deletion of the DNA-binding domain of REV-ERBα in CCSP+ bronchial epithelial cells in mice led to the amplification of airway neutrophilic inflammation upon an aerosolized LPS challenge. Again, this inflammatory response was mediated through the increased expression of CXCL5 at the transcript and protein level. Interestingly, the dual deletion of REV-ERBα and its paralog REV-ERBβ resulted in a further increase in neutrophilic inflammation upon LPS challenge in these mice as well as in unchallenged mice at steady state, suggesting that REV-ERB proteins play a homeostatic anti-inflammatory function. Inflammatory responses to LPS in vivo were not impacted by the deletion of REV-ERBβ alone, emphasizing the leading role of REV-ERBα in this pathway.

A further understanding of the local circadian control of cell function may reveal why some patients experience ‘night-time’ asthma or cough.

**Immune cell interactions at the epithelial surface** scRNA-seq facilitates the mapping of the diverse innate and adaptive immune cell communities of the airway niche at high resolution. Airway epithelial cells act in concert with resident and recruited immune cells to regulate pulmonary immunity (FIG. 4). Whilst transcriptomic studies of human sputum and BAL facilitate the detailed characterization of immune populations within the airway, bronchial brushing samples and biopsies capture both the epithelial and immune cell components. Computational advances are now being leveraged to unveil the complex epithelial–immune crosstalk that occurs in the airway and alveolar space in health and disease.

Macrophages are the most abundant immune cells located in the airway lumen and are found readily in human sputum samples, airway brushings and BAL. Murine studies show that resident airway macrophages are perinatally derived from fetal monocytes and self-maintained locally throughout life. This paradigm was challenged by recent human work revealing, through scRNA-seq, that following lung transplant, most tissue-resident airway macrophages in the donor lung were replaced by airway macrophages from the circulating monocyte pool of the recipient. A single-cell transcriptomic study of sputum samples from patients with cystic fibrosis indicated a disease-related shift in dominant airway immune populations from resident airway macrophages to recruited monocytes and neutrophils, displaying pro-inflammatory phenotypes and gene expression changes suggestive of impaired phagocytosis. Bi-directional interactions between resident airway macrophages and epithelial cells ensure the maintenance of a homeostatic state of immune tolerance to harmless stimuli and appropriate protective responses to inhaled pathogens with effective tissue repair when required. A mouse model utilizing live confocal microscopy showed that CD11c-expressing alveolar macrophages formed direct connections with alveolar epithelial cells by connexin 43-containing gap junctions to limit the inflammatory response to LPS. The exploration of cell–cell signalling networks using scRNA-seq data from lung tissue across four mammalian species (including humans) showed crosstalk between epithelial cells and macrophages in the alveolar cell niche at homeostasis.
There are few studies that investigate epithelial–macrophage crosstalk in the human airways, yet this is a critically important area if we are to fully understand how dysregulated immunoregulatory interactions at this environmental interface contribute to human disease.

To explore the epithelial–immune axis in an inflammatory airways disease, BAL was studied in human subjects with atopic asthma before and after sub-segmental broncho-provocation with an identified allergen. Colony-stimulating factor 1 (CSF1) was elevated in BAL samples after allergen challenge and was shown to be secreted by airway epithelial cells. Epithelial cell-specific deletion of Csf1 in transgenic mice eliminated allergic airway inflammation. Furthermore, epithelial cell-derived CSF1 increased the numbers of a subset of alveolar dendritic cells expressing the CSF1 receptor in BAL following allergen challenge and promoted the migration to regional lymph nodes. Taken together, these findings suggest that epithelial cell–dendritic cell interactions in the airways enhance antigen presentation and augment adaptive allergic airway responses.

A robust immune response to respiratory pathogens at the airway epithelial surface is also contingent on an effective adaptive T cell response. Noteworthy is the induction of a subset of T cells — tissue-resident memory T (T<sub>RM</sub>) cells — at the site of antigen entry in a spatially restricted niche around the airways, poised to generate a rapid immune response to subsequently
encountered pathogens\textsuperscript{41,44}. CD4\(^+\) and CD8\(^+\) T\(_{RM}\) cells expressing CD69 were transcriptionally distinct to circulating CD69\(^+\) effector memory T cells and expressed adhesion molecules that promote retention within the lung mucosal tissue\textsuperscript{36}. Moreover, \textit{parabiosis} experiments in influenza-immune mice found that lung airway CD8\(^+\) T\(_{RM}\) cells were continually replenished by interstitial CD8\(^+\) T\(_{RM}\) cells rather than by circulating memory T cells\textsuperscript{36}. This was mediated by chemokine receptor CXCR6 binding to CXCL16 secreted by airway epithelial cells and macrophages\textsuperscript{46}. In human lung transplant recipients, donor T\(_{RM}\) cells persisted in BAL samples from the lungs but not in peripheral blood for over 1 year\textsuperscript{46}. A significant proportion of airway CD8\(^+\) T\(_{RM}\) cells co-expressed the α\(_5\) integrin CD103, which combines with β\(_7\) integrin to form the heterodimer α\(_5\)β\(_7\), and interacts with E-cadherin on epithelial cells\textsuperscript{46}, promoting retention at this mucosal site. Donor and recipient T cells were localized around the airways in biopsy samples and infiltrating T cells from the recipient acquired a T\(_{RM}\) phenotype over time in BAL samples\textsuperscript{46}.

Crosstalk between T\(_{RM}\) cells and airway epithelial cells can shape the immune response to environmental antigens. Pneumococcal pneumonia generates CD4\(^+\) T\(_{RM}\) cells, which were discovered to fine-tune lung epithelial cells to increase CXCL5 expression and enhance neutrophil recruitment during heterotypic recall infections\textsuperscript{46}. In the context of a house dust mite allergic airways mouse model, the persistence of CD4\(^+\) T\(_{RM}\) cells around the airways was associated with a rapid response to allergen re-challenge, resulting in airway hyperresponsiveness, a cardinal feature of asthma\textsuperscript{46}. Single-cell transcriptomics of airway wall biopsies from asthmatic patients and healthy controls defined two CD4\(^+\) T cell subsets — classic CD4\(^+\) T\(_{RM}\) cells and a novel cluster called tissue migratory CD4\(^+\) T cells — that expressed genes associated with cell egression into tissues such as SIPRI and \textit{SELL}\textsuperscript{46}. A bioinformatic tool was used in this study to interrogate cell–cell interactions in the asthmatic airway wall and unveiled increased predicted crosstalk between T helper 2 cells and epithelial cells, other immune cells, fibroblasts and smooth muscle cells known to play an important role in the pathogenesis of the disease.

The specific properties of the airway microenvironment can drive adaptive gene expression changes in immune cells by epigenetic programming, which shape cell phenotype and function\textsuperscript{46}. Compared with T\(_{RM}\) cells in the spleen and lung interstitium, following influenza infection, CD8\(^+\) airway T\(_{RM}\) cells displayed a distinct transcriptomic and epigenetic profile, enriched for genes associated with the integrated stress response and with amino acid starvation, which ultimately led to a gradual loss of these cells over time by apoptosis\textsuperscript{46}.

Spanning the innate and adaptive arms of the immune system, a subset of innate-like T lymphocytes called mucosal-associated invariant T (MAIT) cells, contribute approximately 4% of the total T cells in human airway wall biopsies\textsuperscript{46}. The expression of a semi-invariant αβ T cell receptor (TCR) allows MAIT cells to recognize metabolites of riboflavin (vitamin B\(_2\)) biosynthesis derived from bacteria and yeasts and presented by MHC-related protein 1 (MR1)\textsuperscript{46}. When activated, MAIT cells generate a rapid pro-inflammatory cytokine response to provide protection against respiratory pathogens\textsuperscript{46} but also display a tissue repair transcriptomic signature, suggesting other important functions in barrier integrity and healing postinfection\textsuperscript{46}.

Mediators produced by resident immune cells can hinder the epithelial cell response to pathogens at the airway mucosal surface. Type I (IFNα and IFNβ) and type III (IFNλ) interferons are key mediators in the host anti-viral response but have recently been shown to compromise epithelial repair following infection\textsuperscript{106,107}. Sustained IFNα produced by lung-resident dendritic cells in response to polynosine-polycytidylic acid (poly(I:C)), a synthetic viral RNA analogue, impaired epithelial barrier integrity, leading to severe pathology with \textit{Staphylococcus aureus} superinfection in a mouse model\textsuperscript{106}. During the recovery phase of murine influenza infection, airway epithelial proliferation and regeneration was reduced in a TP53-dependent manner by prolonged exposure to IFNα\textsuperscript{107}. The nature of the epithelial interferon response is also directly influenced by the type of invading pathogen. A delayed type I interferon response drives the accumulation of inflammatory monocytes and macrophages and leads to severe immunopathology in mice infected with SARS-CoV\textsuperscript{108}. Interestingly, the transcriptional response of human bronchial epithelial cells to SARS-CoV-2 was found to be distinct from that induced by other respiratory viruses — genes associated with type I and type III interferon pathways were suppressed, yet there was a robust expression of genes encoding chemokines and pro-inflammatory cytokines, which is likely to be responsible for many of the clinical manifestations of the disease\textsuperscript{109}. Defects in the type I interferon response due to rare genetic variants or IgG neutralizing autoantibodies were also discovered in a proportion of patients with severe COVID-19 pneumonia\textsuperscript{108,110}.

Complex interactions exist between epithelial cells and innate and adaptive immune cells in the airway to maintain host defence. Signals from epithelial cells maintain tissue resident immune cells and modulate their response to antigen while, conversely, immune cells can directly alter the function and phenotype of airway epithelial cells.

\textbf{Airway neuroimmune interactions}

The general location of airway epithelial cells enables them to sense and react to environmental change via the secretion of a range of mediators that facilitate interactions with immune and stromal cells in the underlying parenchyma. In addition to the specialized epithelial cells that monitor the airways and trigger avoidance reflexes, such as coughing and sneezing, it is well recognized that the airways are richly innervated with sensory neurons, facilitating the execution of these avoidance reflexes\textsuperscript{112}. Recent evidence indicates that extensive crosstalk between neuronal and immune systems and specialized chemosensory pulmonary epithelial cells is vital for the regulation of tissue homeostasis\textsuperscript{113,114} (Fig. 5). This integrated landscape is also able to coordinate responses to invading pathogens or noxious particles, to monitor environmental changes such as fluctuations in temperature or oxygen, or even to react to physical...
events such as stretch or constriction that occur during gasping. For example, PIEZO1, a mechanosensory ion channel expressed by myeloid cells in mice, was shown to sense environmental cyclical hydrostatic pressure changes and generate a pro-inflammatory response. The lungs contain a range of different neural cells, including nociceptor sensory neurons that react to noxious or potentially harmful stimuli, while cholinergic neurons regulate airway tone, airway smooth muscle contraction, mucus secretion and vasodilation via the interaction with muscarinic acetylcholine receptors (mACHRs) found on airway smooth muscle, glands and pulmonary vasculature. Given that many basic respiratory responses, such as cough, are mediated by sensory neurons, it is perhaps not surprising that they are predicted to play an important role in lung inflammatory responses. In addition to relaying sensory changes to the brain, activated lung neurons can themselves release stimulatory peptides that contribute to the ensuing inflammatory reaction. Direct evidence of their contribution to pulmonary inflammation stems from experiments in which the ablation or chemical suppression of Nav1.8 or transient receptor potential vanilloid type 1 (TRPV1) sensory neurons resulted in reduced allergic inflammation and bronchial hyperresponsiveness.

Similarly, targeted lung denervation, a bronchoscopic radiofrequency ablation therapy, is used in patients to durably disrupt parasympathetic pulmonary nerves because it has been found to decrease airway resistance, mucus hypersecretion and inflammation in patients with COPD.

The combination of specialized chemosensory cells and direct neuronal interaction may also represent an alternative way in which the airways respond to inhaled respiratory pathogens, airborne pollutants and allergens. The activation of epithelial TRPV4 triggers protective responses to bacterial LPS, an increasing ciliary...
beat frequency and the production of bactericidal nitric oxide. Mice lacking TRPV4 display exacerbated airway hyperreactivity and enhanced neutrophil recruitment into the airways. Conversely, the specific deletion of TRPV1+ neurons resulted in enhanced immune protection to lethal pneumonia induced by S. aureus via increased survival cytokine induction and bacterial clearance. TRPV1 and Nav1.8+ nociceptor sensory neurons were shown to influence bacterial dissemination via the suppression of neutrophil numbers and surveillance activities and the regulation of resident γδT cell numbers. The specific ablation of vagal TRPV1 neurons resulted in enhanced antibacterial immunity via the release of the neuropeptide CGRP.

Epithelial–neuronal interactions facilitate lung inflammation via the secretion of neuropeptides and neurotransmitters. Many immune cells express receptors that facilitate communication with the epithelium but also directly with nerves. The discovery that ILC2s express a range of receptors for neuropeptides and neurotransmitters reveals a mechanism by which neuronal and immune systems integrate to promote a range of type 2 cytokine responses that facilitate antimicrobial, inflammatory and tissue protective type 2 repair responses at mucosal sites. ILC2 residency in the airways is regulated by signals delivered systemically or by the local tissue environment and these cells are enriched at branch points, similar to PNECs. ILC2s interact directly with neurons via neuropeptides, neurotransmitters and neurotrophic factors, including CGRP, neuromedin U (NMU), acetylcholine and vasoactive intestinal peptide (VIP) and such interactions facilitate both stimulatory and inhibitory signals. ILC2 expression of VIPR2 enables the cells to respond to neuronal-derived VIP by secreting IL-5, thus creating a feedback loop whereby IL-5 stimulates nociceptors to produce more VIP. NMUR1 signalling amplifies ILC2-driven allergic inflammation. NMU secreted by thoracic dorsal root ganglia interacts with NMUR1 expressed by lung ILC2s and regulated by IL-25. The loss of NMUR1 reduces ILC2 responses and type 2 inflammation. NMU also promotes the contraction of smooth muscle cells, although this has not been shown specifically in pulmonary smooth muscle cells. Conversely, CGRP has differential effects on the immune response to allergen-induced inflammation. Mice lacking PNECs have a reduced expression of both CGRP and GABA after allergen exposure. CGRP induces IL-5 secretion from ILC2s and inactivation of the CGRP receptor (Calarl1) in ILC2s in vitro ameliorates allergic inflammation. CGRP is also found to negatively regulate type 2 responses to either allergen exposure or helminth infection. CGRP modulates type 2 cytokine production from ILC2s that have been activated by alarmins or NMU, thereby constraining the magnitude of the type 2 responses in vivo. PNEC-derived GABA seems to affect mucus secretion since the inactivation of GABA production results in defective goblet cell hyperplasia and direct instillation of GABA in PNEC-deficient mice restores mucus production. These differential effects of neuropeptides on ILC2 cytokine production and type 2 immune pathology, concomitant with epithelial cell-derived alarmins and neuropeptides, provides an integrated mechanism whereby tissue responses to allergen can be fine-tuned according to changing circumstances.

**Concluding remarks and future directions**

Our views of the human airway epithelium have shifted dramatically, catalysed by the advancement in technologies that facilitate single-cell transcriptomics. The cellular communities of the airway epithelium are diverse, dynamic and yet functionally cohesive. Epithelial cells work collaboratively with localized populations of immune cells to support a multifaceted frontline defence system with highly effective injury-repair responses.

Epithelial, stromal and immune cells in the lung can be modulated by many aspects of their local tissue microenvironment, including the properties of the extracellular matrix, mechanical stress and exposures such as to cigarette smoke. Mapping cell location in a specific anatomical tissue microenvironment using spatial transcriptomic techniques will undoubtedly enhance our understanding of cell molecular phenotype and function within a particular spatial niche.

A multi-omic approach utilizing single-cell transcriptomics and mass spectrometry-based proteomics was used to chart the molecular and cellular alterations that occur in the lung with ageing. This work showed alterations in airway epithelial cell composition and lung extracellular matrix with increasing age. The application of a similar approach to human tissue samples in combination with comprehensive demographic and clinical information will allow us to link changes in cell biology with disease phenotype.

The functional validation of transcriptomic data that has been generated through complex bioinformatics requires robust in vitro culture systems. The human ‘small-airway-on-a-chip’ was developed to recapitulate the airway tissue microenvironment and cell responses in health and disease, and represents a significant development in this area. This system comprises a microfluidic device with a fully differentiated, mucociliary airway epithelium cultured adjacent to a microvascular endothelium in a channel exposed to media under flow. It has been used to model inflammatory airway diseases and study the interplay between epithelial cells and immune cells in vitro.

The airway epithelium is dynamic and continuously replenished by basal progenitor cells. Single-cell transcriptomics has been used to elucidate airway basal cell differentiation and response to injury in vitro. Fascinating mechanisms exist to preserve a healthy barrier in other stratified epithelial surfaces around the body such as the skin. A natural phenomenon of cell competition was observed to operate in a mouse model of skin development — in a mosaic epidermal tissue, so-called ‘winner’ progenitor cells induce apoptosis in neighbouring apoptotic ‘loser’ progenitor cells and eliminate them by engulfment. Cell losers not cleared by this mechanism were later eliminated from the basal cell layer by differentiation through the stratified cell layer. This is physiologically important because, when cell competition was disturbed, epithelial barrier function was compromised. Whether this competitive...
cell behaviour is conserved throughout other epithelial tissues, including the airway lining, remains to be elucidated and warrants further investigation.

Transcriptomic technologies have enhanced our view of the cellular landscape in the human lung but there remain inherent limitations and many unanswerd questions. Loss of fragile cell types in the process of sequencing can bias the analysis and variation in bioinformatic pipelines and cell annotation can hinder comparisons between different studies. Rare epithelial cell types have attracted much interest but, given their low frequency within the lung, the challenge will be to ascertain their contribution to diverse pathologies in the hope that we can harness and manipulate particular cells to promote lung health across the human life course.

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