Hedgehog-responsive PDGFRa(+) fibroblasts maintain a unique pool of alveolar epithelial progenitor cells during alveologenesis

Feng Gao¹, Changgong Li¹, Soula Danopoulos⁴, Denise Al Alam⁴, Neil Peinado¹, Sha Webster¹, Zea Borok²,³, GoleNaz Adeli Kohbodi¹, Saverio Belluscio¹,⁵, Parviz Minoo¹,²,⁶,*

¹Division of Neonatology, Department of Pediatrics, LAC+USC Medical Center and Childrens Hospital Los Angeles, Los Angeles, CA 90033, USA
²Hastings Center for Pulmonary Research, Keck School of Medicine of University of Southern California, Los Angeles, CA 90033, USA
³Division of Pulmonary, Critical Care and Sleep Medicine, University of California San Diego School of Medicine, San Diego, CA 92093, USA
⁴Lundquist Institute for Biomedical Innovation at Harbor-UCLA Medical Center, Torrance, CA 90502, USA
⁵Universities of Giessen and Marburg Lung Center (UGMLC), Justus-Liebig-University Giessen, German Center for Lung Research (DZL), 35390 Giessen, Germany
⁶Lead contact

This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).

AUTHOR CONTRIBUTIONS
F.G. performed experiments and curated data and analysis. C.L. analyzed data and directed the study. S.D. performed and analyzed Human RNAscope data. D.A.A. performed and analyzed Human RNAscope data. N.P. analyzed data. S.W. performed experiments and analyzed data. Z.B. critiqued the data and assisted in writing the manuscript. G.A.K. performed experiments and assisted with data analysis. S.B. assisted with data analysis and writing the manuscript. P.M. developed the concept, analyzed data, and wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.celrep.2022.110608.

SUPPORTING CITATIONS
The following references are cited in the supplemental information: Cassandras et al., 2020; Chen et al., 2017; Frazier et al., 2019; Hagan et al., 2020; Huqui et al., 2019; Jin et al., 2018; Kuang et al., 2021; Liu et al., 2018; Manchado et al., 2016; Maynard et al., 2020; Nakamura et al., 2015; Park et al., 2019; Treutlein et al., 2014; Wang et al., 2019; Wei et al., 2021; Wimmer et al., 2019; Wong et al., 2018; Zhang et al., 2021.
SUMMARY

The lung alveolus is lined with alveolar type 1 (AT1) and type 2 (AT2) epithelial cells. During alveologenesis, increasing demand associated with expanding alveolar numbers is met by proliferating progenitor AT2s (pAT2). Little information exists regarding the identity of this population and their niche microenvironment. We show that during alveologenesis, Hedgehog-responsive PDGFRa(+) progenitors (also known as SCMFs) are a source of secreted trophic molecules that maintain a unique pAT2 population. SCMFs are in turn maintained by TGFβ signaling. Compound inactivation of Alk5 TβR2 in SCMFs reduced their numbers and depleted the pAT2 pool without impacting differentiation of daughter cells. In lungs of preterm infants who died with bronchopulmonary dysplasia, PDGFRα is reduced and the number of proliferative AT2s is diminished, indicating that an evolutionarily conserved mechanism governs pAT2 behavior during alveologenesis. SCMFs are a transient cell population, active only during alveologenesis, making them a unique stage-specific niche mesodermal cell type in mammalian organs.

In brief

Much of the information regarding heterogeneity of alveolar epithelial cells in the lung is from adult mice. Gao et al. analyze the complexity of the alveolar epithelial cell population during alveologenesis, a process interrupted in the pathogenesis of the neonatal chronic lung disease bronchopulmonary dysplasia.

Graphical Abstract
INTRODUCTION

From the time its primordium is specified within the anterior foregut, lung development is predicated on mesenchymal-epithelial crosstalk (Basil et al., 2020). This communication is particularly complex during alveologenesis, a process that requires concerted functional interactions among multiple cell types. Significant changes in proliferation and differentiation of embryonic cells accompany alveologenesis, which in humans occurs partly and in the mouse entirely postnataally. By completion, the adult alveoli, which number in the millions in mice and man alike, will consist of functional epithelial, mesenchymal, and immune cells, plus a massive capillary plexus, an evolutionary response to oxygen requirement in terrestrial mammals.

The alveolar inner lining is composed of flat alveolar epithelial type 1 cells (AT1s), which are critical for gas exchange, and cuboidal alveolar type 2 cells (AT2s) that produce and secrete pulmonary surfactant. Some AT2s serve as regenerative stem cells (Zacharias et al., 2018; Nabhan et al., 2018). They are key to maintaining homeostasis and repair of lung injury. Regenerative AT2s are WNT responsive and cohabit in niches with specialized mesenchymal cells that direct their behavior. In organoid cultures, platelet-derived growth factor receptor alpha positive (PDGFRa(+)) cells, a heterogeneous population, are thought to serve as mesenchymal “niche” cells (Barkauskas et al., 2013; Zepp et al., 2017). Some candidate mediators of their communication have been identified and functionally tested in organoid cultures (Chung et al., 2018; Green et al., 2016; Zepp et al., 2017). The identity of niche cells in vivo and the mechanisms by which they modulate AT2 behavior are less understood. Furthermore, whether temporally specific niches are present during lung development remains unknown.

During alveologenesis, Sftpc(+) epithelial progenitors (simply, pAT2s) undergo major expansion in number and differentiate to populate the adult alveoli with AT1s, AT2s, and regenerative AT2s (Frank et al., 2016). The mechanisms are not fully understood. Specifically, whether development of pAT2s is guided within interactive microenvironments akin to the adult niches and the required mechanisms of instructive mesenchymal-epithelial signaling remain unknown. More importantly, whether pAT2s in adult lungs can respond to injuries that occur during alveologenesis is also unknown. These are particularly relevant questions in the context of bronchopulmonary dysplasia (BPD), a chronic lung disease that occurs during alveologenesis in preterm infants.

Alveologenesis requires the formation of “secondary crests” composed of multiple cell types with distinct lineage histories. Located at the tip of secondary crests are PDGFRα(+) cells whose role in alveologenesis has been investigated in a number of studies (Bostrom et al., 2002). A recent study found that 95% of cells lineage-traced with Pdgfra-rtTA are located at septal tips in postnatal day 7 (PN7) lungs, cells we previously referred to as secondary crest myofibroblasts (SCMFs) (Li et al., 2018). In the present work we use SCMFs and PDGFRα(+) interchangeably. SCMFs are Hedgehog (HH) responsive and can be targeted effectively by Gli1-creERT2, a high-fidelity HH reporter (Ahn and Joyner, 2004; Li et al., 2015). PdgfraRTTA also targets SCMFs, and partial ablation of Pdgfra(+) cells arrests alveologenesis (Li et al., 2018). Despite their designation as “myofibroblasts,” indicating
aSma expression, SCMFs are distinct from “myofibroblasts,” a term commonly used in reference to pulmonary fibrosis, by quantitatively low aSma expression only in the late fetal to early postnatal period and enriched expression of matrix proteins required for alveologenesis (Li et al., 2015). Transforming growth factor β (TGFβ) signaling is increased in SCMFs at the onset of alveologenesis (Li et al., 2015). TGFβ functions by interacting with a heteromeric complex of transmembrane receptors, ALK5 and TβR2. In the lung, TGFβ is commonly associated with lung “injury” and fibrosis, but it also has diverse functions in normal development (Kaartinen et al., 1995; Pieretti et al., 2014).

To determine the significance of TGFβ signaling in SCMFs during alveologenesis and its role in differentiation of AT2 and AT1 cells, we used Gli1-creERT2 and generated compound mutants with inactivated Alk5 and TβR2 in SCMFs, specifically on PN2, at the onset of alveologenesis. The mutations blocked alveologenesis, reducing the number of SCMFs and pAT2s. Single-cell RNA sequencing (RNA-seq) identified a unique population of proliferating pAT2s whose pool was profoundly depleted in the mutant lung. We found that SCMFs constitute a key signaling hub, providing ligands with roles in self-renewal of pAT2s. These characteristics suggest that SCMFs are a principal component of the alveolar niche during alveologenesis. Intriguingly, SCMFs are a transient cell population that are depleted at completion of alveologenesis, making them a unique and temporally specific niche cell type in mammalian organ development (Li et al., 2015; Hagan et al., 2020).

RESULTS
Postnatal inactivation of TGFβ receptors in SCMFs arrests alveolar formation, phenocopying human BPD

Both TGFβ receptors were inactivated in Gli1-creERT2; ROSA26tm1TmG;Alk5(+/−);TβR2(+/−) newborn mice (hereafter DKO) with one dose of tamoxifen on PN2, as described in STAR Methods. We previously showed that this regimen targets SCMFs (Li et al., 2015, 2019). We generated and used Gli1-creERT2;Rosa26tm1TmG as control. Histology of multiple biological replicates of DKO lungs at two time points, PN14 and PN45, revealed a phenotype of profoundly arrested alveologenesis (alveolar hypoplasia) resembling human BPD lungs (Figures 1A–1D). Quantitative morphometry (n = 3 biological and >10 experimental replicates) using mean linear intercept (MLI) showed increased alveolar size by 52%–58% (Figures 1E and 1F). MLI did not increase in mutant lungs over time, and thus the DKO phenotype, unlike other mutations in the TGFβ pathway, is not progressive (Figure 1G). These observations clearly indicate a critical requirement for functional TGFβ receptors in SCMFs during alveologenesis.

Inactivation of TGFβ receptors in SCMFs reduces both SCMFs and pAT2 cell numbers

Formation of alveoli occurs in close association with specification and maturation of alveolar epithelial progenitors. Aqp5 and Pdpn, both AT1 markers, decreased in DKO lungs (Figure 1H). However, AT1 numbers, determined by manual counting of HOPX(+) cells in control and mutant lungs (Figures 1J versus 1K) showed a small difference that failed to reach statistical significance (Figure 1I). In contrast, pAT2 numbers (SFPTC(+) decreased significantly in the mutant lungs and were not proportional to reduction in total
cell numbers, indicating it is not simply a consequence of alveolar hypoplasia (Figures 1L–1N). Changes in Sftpc mRNA, if any, did not reach statistical significance (Figure 1O). Reduced pAT2 numbers was confirmed by fluorescence-activated cell sorting (FACS) using GFP-labeled pAT2s (Figure 1P). Total GFP-labeled SCMF numbers also decreased in mutant lungs (Figure 1Q). Neither change was proportional to overall reduction of cell numbers due to hypoplasia. Thus, inactivation of TGFβ receptors in SCMFs results in reduced number of both pAT2s and SCMFs. As adult PDGFRa(+) cells are known to interact with adult AT2s in organoid cultures, we posited that loss of pAT2s in the neonatal lung follows from loss of mutant SCMFs through interruption of SCMFs-pAT2s crosstalk.

Molecular phenotype of PN14 lungs at single-cell resolution

To elucidate the molecular basis of SCMF-pAT2 crosstalk, we performed single-cell RNA-seq (scRNA-seq) using the 10X Genomic pipeline and determined the molecular phenotype of the entire, intact distal PN14, control (Gli1-creERT2; Rosa26mTmG) and mutant (Gli1-creERT2; Rosa26mTmG, Alk5(flox/flox); Tβr2(flox/flox)) lungs at single-cell resolution (STAR Methods). Data were filtered by total unique molecular identifier (UMI), mitochondrial reads, possible doublets, and batch effects, resulting in 12,582 cells from control and 13,819 cells from the mutant PN14 lungs. Median numbers of transcripts per cell were 3,345 and 2,683 for control and mutant lungs, respectively. The median number of genes detected per cell was 1,700 for control and 1,352 for mutant lungs. T-distributed stochastic neighbor embedding (tSNE)-based representation of the combined data revealed four transcriptionally distinct major compartments and 38 cell type subclusters in both genotypes (Figure S1). Well-established cell lineage markers identified the predictive four major compartments, namely mesenchymal or MECs (n = 1,627 expressing Col1a1), endothelial or ENCs (n = 6,421 expressing Cdh5), immune or IMC (n = 10,723 expressing Ptprc), and epithelial or EPC (n = 5,664 expressing Nkx2.1) (Figure S2). To focus on the potential genetic changes that might account for altered SCMF-pAT2 cross-communication, we analyzed in detail the scRNA-seq data for MECs and EPCs.

Identification of the mesenchymal subpopulation disrupted by TGFβ receptor inactivation

Isolation and reclustering of MECs revealed three distinct mesenchymal subclusters C1, C2, and C3 (Figure 2A). Inactivation of TGFβ receptors caused selective loss of C3, contracting it by 81.6% and indicating C3 as the principal target of Gli1-creERT2 (Figure 2, table). In control lungs, C3 accounts for 67.3% of total MECs and 5.5% of the total number of interstitial cells. These numbers are respectively reduced to 12.9% and 0.6% in the mutant lungs. C3 includes the highest number of mesenchymal cells expressing both Tbr2 and Alk5 (Figure S3A) and thus is exquisitely sensitive to loss of TGFβ signaling, which is widely recognized as necessary for growth and maintenance of mesodermal cells.

To assign identity to C3 cells, we analyzed differentially expressed (DE) genes in a three-way comparison among C1, C2, and C3 and determined the enriched transcripts in each of the three subclusters (Figures 2B and 2C) and compared them against signature genes in LungMap (www.lungmap.net). PDGFRα is a widely acknowledged, principal marker of a heterogeneous population of mesenchymal cells found in the alveolar compartment (Chen et al., 2012; Green et al., 2016; McGowan and McCoy, 2014). Pdgfra is highly enriched only
in C3 (Figure 2D). Lineage traced Pdgfra(+) cells are located at the septal tips in PN7 lungs, which we have referred to as SCMFs (Li, et al., 2018). Importantly, Fn1, Inmt, and Mfap4, which encode extracellular matrix (ECM) proteins required for alveolar formation, are also enriched in C3. C1 and C2 were identified as pericytes and smooth muscle cells, respectively (Figure S3). Computational analysis showed that 80% of the DE genes between control and mutant MECs (i.e., C1 + C2 + C3) are enriched in control C3 and lost in the mutant (Figure S4). Thus, for the larger part, the observed changes in mesenchymal gene expression are reflective of the loss of C3 cells and the pAT2 phenotype in the mutant lungs.

Identification of a unique proliferative epithelial subpopulation during alveologenesis

Reclustering of the control and mutant EPC data identified eight distinct subpopulations, C4 to C11 (Figure 3A). There were changes in cell numbers in the mutant subclusters (Figure 3B, table). In the control lung, C4 represented 2.25% of total lung cells and 9.06% of total EPCs. These numbers in the mutant lung were reduced to 0.04% and 0.24%, respectively. Thus, a loss of greater than 96% occurred in mutant C4. The identity of cells in C4 to C11 was determined by the extent of overlap between their individual enriched transcripts and the most recent AT1, AT2, and intermediate AT1/AT2 signature genes available on LungMap (www.lungmap.net). C4 represents a unique proliferative and least differentiated subpopulation of pAT2s based on highest expression of cell-cycle genes Top2a, Mki67, Cdk1, and Spc24, combined with coexpression of low levels of AT1, AT2, and AT1/AT2 markers (Figures 3C–3E). Pathway analysis confirmed enrichment of cell proliferation activity in C4 (Figure S5A). Similar analysis identified C5 to C9 as differentiating pAT2s with progressively increased expression of AT2 signature genes, including lamellar body and pulmonary surfactant genes Sftpb and Sftpa1 (Figures 3C, 3D, and S5B). Importantly, only minor differences exist between the profiles of mutant versus control C5 to C9, indicating no major alterations in lineage or differentiation trajectory in response to loss of TGFβ signaling in SCMFs (Figures 3C versus 3D). However, we were surprised to find that nearly all epithelial clusters displayed coexpression of AT2 and AT1 signature genes (Figures 4A and 4B). Presence of intermediate cells in PN14 lungs was confirmed by RNAscope (Figure 4C). Therefore, C5 through C9 represent intermediate or transitional differentiating pAT2s in PN14 lungs, as late as halfway through the alveologenesis period. Interestingly, in C10 and C11, AT2 markers SftpC and SftpB, and SftpA1, were strongly repressed (Figure S5B). The differences between C10 and C11 are the increased expression of maturing AT1 signature genes Hopx, Pdpn, Igtbp2, and Spock2 (Wang et al., 2018), and metabolic genes in C11. Indeed C11 transcriptome included Hopx, Pdpn, Igtbp2, and Spock2, consistent with maturing/mature AT1 cells (Figure S7). C10, in contrast, displays a low metabolic profile and reduced AT2 signature genes, indicating a potential inflection point along the differentiation trajectory (Figure 4D).

Pseudo-time trajectory of dynamic gene expression and cell fate transitions suggests that C4 serves as the origin of the transitional epithelial cells described above (Figure S6). Computational analysis of DE genes between control and mutant EPCs, and comparison with enriched genes in each of the EPC subclusters, shows that 85% of the DE genes in the mutant lung originated from C4 (Figure S8). Thus, C4 is the major impacted epithelial population in the mutant lung. As the mutation is exclusively targeted to HH-responsive
mesenchymal cells, depletion of the C4 pool likely occurs through disruption of normal SCMFs-pAT2s cross-communication.

**Ligand-receptor (L-R) pairing identifies six major pathways**

AT2-mesenchymal interactions have been examined largely by *in vitro* or organoid cultures. In the present *in vivo* study, we identified ligands enriched in C3 and differentially expressed between control and mutant lungs by using the computational algorithm shown in Figure S9. Among 251 genes, we identified 13 (fold change [FC] > 2 and p ≤ 0.05) that encoded ligands based on the annotated Fantom Reactome database. *Cp* or ceruloplasmin is expressed in airways and thus may represent contamination in RNA-seq data (Yang et al., 1996). The other 12 genes fell into two functional groups. Npnt, Cyr61, Bgn, and Col6a3 are related to ECM-based signaling. The other eight encode signaling molecules, members of the HGF, FGF, BMP, WNT, and IGF pathways. These are selectively enriched in C3 (Figure 5A). RNAscope revealed overlapping expression of a select number of ligands with Pdgfra in PN14 lungs (Figure 5B). Finally, we paired the eight ligand genes in C3 with receptor genes that were defined as actively expressed in pAT2 cells in C4 (i.e., transcripts ≥30 copies/cell) and annotated in our scRNA-seq data using the STAR method (Figure 5C) (Ramilowska et al., 2015). The L-R pairing implicated six major functional pathways, whose reduced signaling, due to SCMF loss in the mutant lung, can account for depletion of the C4 pool (Figure 5C).

**Functional analysis of putative signaling pathways in 3D organoid culture using neonatal pAT2s**

The potential function of the six signaling pathways was examined in *ex vivo* three-dimensional (3D) organoid assays (Barkauskas et al., 2013). FACS-isolated pAT2s from PN14 Sftpc-GFP lungs were cocultured with lung mesenchymal cells. In each organoid assay (n = 5 biological replicates) we blocked one of the six principal signaling pathways shown in Figure 6, using specific small-molecule inhibitors for NOTCH, FGF, and IGF1 and recombinant GREM2 for BMP (see Figure 6 legend and Table S1 for concentrations used). The function of individual inhibitors was validated as shown in Figure S10. Blocking HGF with capmatinib, an ATP-competitive inhibitor of c-MET (Clement et al., 2020; Kong et al., 2020; Oh et al., 2021), while dramatically reducing organoid size, had a moderate impact on colony-forming efficiency (CFE) (Figure 6A, a8 and a9). Inhibition of BMP by GREM2 (Tanwar et al., 2014) had no impact on CFE but reduced the organoid size, suggesting its activity may be more related to pAT2 proliferation (Figure 6A, a8 and a9). The most significant inhibition of CFE occurred by blocking FGF using BDJ398 (De Simone et al., 2021; Qi et al., 2019; van den Brink et al., 2020), but also reduced organoid size (Figure 6A, a8 and a9). Inhibition of NOTCH, WNT, and IGF1, using DAPT (Rognoni et al., 2014), XAV939 (Burridge et al., 2014), and PPP (Gong et al., 2019), respectively (STAR Methods), had similar impact on clonogenicity (Figure 6A, a8). NOTCH inhibition had the most moderate impact and IGF1 and WNT the most profound impact on organoid size (Figure 6A, a9). WNT2 is reported to be associated with self-renewal of stem cells (Zeng and Nusse, 2010), and we found it to be expressed nearly exclusively in C3 (Figure 6B, b1). Using Fgfr2 as a marker of AT2s in PN14 lungs (LungGens Consortium and Figure 6B, b2) RNAscope showed that Wnt3a(+) cells are juxtaposed to proliferating, Mki67;Fgfr2 double-
positive cells, indicating WNT2 may be a likely candidate in regulating self-renewal of C4 cells during alveologenesis (Figure 6B, b3–b8). Collectively, the results implicate a rather complex network of multiple signaling pathways, which functionally converge to regulate both clonogenicity and cell proliferation of PN14 pAT2s during postnatal alveologenesis.

Modeling the impact of mesenchymal cell population changes on neonatal pAT2 self-renewal

In the mutant lung, both SCMFs and pAT2s decreased. To examine the significance of cell number changes (i.e., population shifts), we re-employed the 3D organoid culture combined with variable pAT2/mesenchymal cell stoichiometry using control cells. When cocultured with a fixed number of mesenchymal cells (50K), CFE was independent of pAT2 numbers, indicating that only a limited, constant fraction of pAT2s in each sample is clonogenically competent (i.e., progenitor/stem cell property) (Figure 6C, c1–c6 and c13). However, organoid size, more a measure of pAT2 self-renewal, was inversely proportional to pAT2 numbers (Figure 6C, c15). Conversely, when pAT2 numbers were fixed at 5K, CFE as well as organoid size were directly proportional to the number of cocultured mesenchymal cells (Figure 6C, c7–c12, c14, and c16). Therefore, a threshold level of mesenchymal cell number is required, below which organoid formation is nearly undetectable. Overall, as the mesenchyme is the source of factors required for stimulating organoid formation and growth, the results show that decreasing the signaling amplitude by increasing pAT2 numbers, or decreasing mesenchymal cell number, limits accessibility of individual pAT2s to growth molecules, thus resulting in smaller average size per organoid (i.e., cell numbers). This condition mimics and explains the findings in the TGFβ receptor mutant lungs.

The mechanism governing pAT2 behavior in alveologenesis is evolutionarily conserved in the human neonatal chronic lung disease BPD

BPD, which is characterized by alveolar hypoplasia, is a highly complex, variable, and poorly understood neonatal lung disease (Sahni and Mowes, 2020). Two major etiologic factors are lung immaturity and injury. In adult mice, lung injury triggers AT2 self-renewal and differentiation as part of the regenerative/repair mechanism. Mouse Axin2(+);Sftpc(+) cells, referred to as alveolar epithelial progenitors (AEPs), show enhanced regenerative potential compared with Sftpc(+) cells (Zacharias et al., 2018). The adult human lung contains an equivalent of the mouse Axin2(+);Sftpc(+) cells (Zacharias et al., 2018). Whether such regenerative cells are present in human neonatal lungs and their population size in BPD are unknown. In the present study, loss of Pdgfra(+) SCMFs inhibited alveologenesis and depleted the pAT2 pool. Reduced PDGFRα is a primary feature of BPD (Popova et al., 2014). However, the size of the proliferative pAT2 pool in BPD lungs is unknown. We used a combination of immunofluorescence (IF) staining and RNAscope to determine the size of the proliferative pAT2 pool and the extent of AT2 proliferation using multiple human BPD and control samples (n = 4 biological replicates, n = 15–20 experimental replicates for each). Quantification of the number of AXIN2(+);SFTPC(+) cells as a fraction of total pAT2 cells, which corrects for alveolar hypoplasia, was similar or greater in BPD lungs compared with controls (Figures 7A–7D and 7I). Importantly however, double-positive SFTPC(+)MKI67(+) proliferative cells were distinctly reduced in BPD lungs (Figures 7E–7H and 7J). These data suggest that while BPD lungs may contain a
larger or similar reservoir of the presumptive Axin2(+)–pAT2 regenerative cells, they appear to lack the normal proliferative response to injury.

**DISCUSSION**

We used combination of loss-of-function genetics, scRNA-seq, and 3D organoid cultures to elucidate the role of HH-responsive, Pdgfrα(+) cells, previously referred to as SCMFs, in alveologenesis. Inactivation of TGFβ receptors in SCMFs caused significant loss of these cells, a principal source of trophic molecules, and arrested alveolar formation, a phenocopy of human BPD. Reduced SCMF numbers profoundly depleted a unique pool of proliferative epithelial cells, which by inference from trajectory mapping serve as a reservoir for differentiating AT1 and AT2 cells. These findings support a model in which TGFβ signaling is critical for proliferation of SCMFs, which in turn, via crosstalk, maintains pAT2 self-renewal. Finally, using human BPD lung samples, we show that as in the mouse model, reduced self-renewal of pAT2s and not their absence may underlie pathogenesis of this neonatal chronic lung disease.

Inactivation of TGFβ receptors in the mutant lungs resulted in 81% asymmetric reduction of SCMFs represented by subcluster C3 (Figure 2). C3 comprises the largest number of Alk5(+) and Tbr2(+) expressing cells (Figure S3), making their proliferation particularly sensitive to loss of TGFβ signaling, widely recognized for its role in regulation of mesenchymal growth and proliferation (Grafe et al., 2018). Indeed, isolated mutant GFP-labeled SCMF cell lines display reduced proliferation in vitro compared with controls (Figure S11). We demonstrate that SCMFs are a mesenchymal signaling hub for a suite of trophic paracrine molecules, including ligands for six major signaling networks. The consequences of C3 cell losses may include reduced availability of these ligands to the proliferating and differentiating epithelial progenitor cells that are required during alveologenesis. The mutant lungs also contained significantly fewer pAT2s due to reduced proliferation, as revealed by analysis of proliferating SFPTC(+) cells across PN7 and PN14 control and mutant lungs (Figure S11). Analysis by scRNA-seq showed that the Sfptc(+) compartment is composed of eight subpopulations. The most impacted population was C4, comprising the least differentiated, most highly proliferative epithelial progenitors. Loss of TGFβ and the consequent reduction in SCMF numbers depleted the C4 pool by 96% (Figure 3B).

Gene enrichment analysis and pseudo-time trajectory mapping inferred that C4 serves as a reservoir for differentiating alveolar epithelial transitional subclusters (Figure S6). In these subclusters we found widespread presence of epithelial intermediate cells coexpressing mature AT2 (Sftpc, Fasn) and multiple AT1 markers (e.g., Aqp5 and Ager). Epithelial cells coexpressing AT1 and AT2 genes, so-called bipotential, have been described in the saccular phase of lung development but are exhausted by PN4 (Desai et al., 2014, Treutlein et al., 2014). Intermediate AT1/AT2 cells were also reported in early PN1 lungs (Guo et al., 2019). The abundant presence of intermediate cells in PN14 may represent plasticity and incomplete cell fate determination as late as the mid-alveologenesis period. Interestingly, intermediate AT2/AT1 cells have also been observed during alveolar regeneration in multiple injury models including lipopolysaccharide (Riemondy et al., 2019) and bleomycin...
Thus, it appears that regeneration of injured alveoli may largely recapitulate the developmental process of pAT2s during alveologenesis. It is important to note that in both mutant and control lungs, once the daughter cells exited the C4 pool, their differentiation followed a similar trajectory (Figures 3C and 3D). As SCMFs are a transient cell population depleted at completion of alveologenesis (Li et al., 2015; Hagan et al., 2020), their principal function may be limited to providing trophic factors that stimulate or maintain self-renewal of pAT2s only during alveologenesis.

In general, knowledge of factors regulating AT2 self-renewal in vivo and during alveologenesis remains incomplete. In adult lungs, expression of Fgf7, Bmps, and Il6 are reported as potential regulators of AT2 behavior and have been examined in organoid cultures (Chung et al., 2018; Green et al., 2016; Zepp et al., 2017). We leveraged a loss-of-function approach, specifically during alveologenesis, to identify distinct signaling pathways whose in vivo inhibition negatively impacted pAT2 self-renewal. We identified six major signaling networks, HGF, FGF, BMP, WNT, NOTCH, and IGF1. Of these, NOTCH, IGF1, and HGF are less studied. Blocking HGF had the strongest impact on organoid size. HGF is a recognized growth factor for AT2 proliferation in vitro (Calvi et al., 2013; Ohmichi et al., 1998). Blocking NOTCH or IGF1 also impacted clonogenicity and organoid size. The role of NOTCH in self-renewal of airway basal stem cells and differentiation is recognized (Kiyokawa and Morimoto, 2020; Tsao et al., 2016). Consistent with our results, blocking NOTCH signaling in AT2s is reported to cause abnormal lung development and loss of AT2s (Tsao et al., 2016). Limited information is available on the potential role of IGF1 in AT2 regulation (Ghosh et al., 2013). However, IGF1 sustains stem cell-mediated repair of tubular epithelial cells in acute kidney injury (Imberti et al., 2007). The potential in vivo role of these ligands in pAT2 self-renewal remains to be investigated by utilizing loss-of-function models.

Our findings regarding FGF, BMP, and WNT in organoid cultures are largely consistent with previous reports (Chung et al., 2018; Hu et al., 2020; Zepp and Morrisey, 2019). We found clonogenicity to be particularly sensitive to changes in FGF and WNT. In the adult lung, paracrine WNT signaling from a single Wnt5a(+) fibroblast in close proximity to adult Axin2(+) stem cells governs their behavior (Nabhan et al., 2018). C3 cells express WNT ligands including Wnt5a (data not shown). 3D reconstruction of confocal images in PN14 lungs shows that Gli1-creERT2-targeted GFP(+) and SFPTC(+) cells are in close spatial proximity to one another (Figure S11). Wnt2-expressing cells, which are exclusively in the C3 subcluster, are indeed juxtaposed to pAT2 cells (Figure 6). Importantly, C3 cells are also Prcn(+), indicating that they secrete WNT ligands (Resh, 2016) (Figure S12). In contrast, the majority of C4 cells are Prcn(−), raising the possibility of a paracrine WNT-mediated C3-C4 crosstalk during alveologenesis akin to that described in the adult stem cell niche (Nabhan et al., 2018).

WNT signaling and expression of its target Axin2 have been described in AT2s during lung development and in response to injury (Basil et al., 2020). In early alveologenesis, the majority of proliferating AT2s are reported to be Axin2(+) (Frank et al., 2016). In our study, nearly all C4 cells are proliferating (Figure 3) and yet only 10% are Axin2(+) (Figure S12). This percentage remains the same in mutant C4 cells despite profound depletion of total
AT2s (Figure 3). Adult lung AT2 facultative stem cells that proliferate in response to injury are also \textit{Axin2}(+). These cells are reported to be rare, constituting between 1% and 20% of total AT2s (Nabhan et al., 2018; Zacharias et al., 2018). While we do not as yet know the significance of the rare \textit{Axin2}(+) population in C4, they are proliferative and may represent progenitors of a unique subset of adult AT2s with proliferative potential. It is noteworthy that in the organoid cultures we found percent CFE to remain constant, regardless of the number of input pAT2s (Figure 6). This indicates that during alveologenesis only a given fraction of total pAT2s is clonogenically competent, perhaps by virtue of possessing proliferative potential. In this regard, we found a decreasing trend in the overall ratio of \textit{Axin2}(+) pAT2s to total AT2s as pAT2s differentiated along the trajectory line. To determine whether the proliferative status of the cells is reflected in their metabolic state, we examined the expression of multiple subunits of the NADH pathway, the largest of the respiratory complexes. As shown in Figure 4D, transcripts for the NADH subunits decreased along the C4 to C10 trajectory, which largely followed the pattern of reduction in the \textit{Axin2}(+) pAT2s to total AT2s shown in Figure S12. The metabolic decrease occurred strongly in C8, C9, and C10 at the inflection point, where \textit{Sftpc}, the marker for AT2 cells, began to decrease (Figure 4A). C8, C9, and C10 also had the lowest \textit{Axin2}(+) pAT2s/total AT2 ratios (Figure S12). Metabolic transcripts were restored in C11 cells, which represent differentiated AT1 cells (Figures S5B and S4D). AT1s are thought to be \textit{Axin2}(−), and terminally differentiated, implying loss of proliferative potential. We found residual \textit{Axin2}(+) cells within the C11 cluster, suggesting the presence of perhaps a distinct subpopulation of AT1s that retain proliferative capacity. Developmental plasticity and reversal of the terminally differentiated state have been suggested for AT1 cells (Yang et al., 2016). Our collective observations indicate that proliferative potential may be progressively lost during differentiation of cells exiting the C4 pool. However, as each of the differentiating subclusters still retains a pool of \textit{Axin2}(+) cells, one can speculate that the cells in early alveologenesis (PN14) may maintain residual proliferative capacity. While the validity of this concept needs to be tested, it raises potentially exciting possibilities for future treatment modalities for BPD.

Pathogenesis of BPD is the result of injury to the developing, nascent alveoli caused by the side effects of clinical interventions necessary to maintain life after preterm birth. In adult mice, H1N1 influenza causes extensive alveolar injury. In response, AT2 cells identified as \textit{Axin2}(+);\textit{Sftpc}(+) and referred to as AEPs undergo self-renewal and differentiate to regenerate the injured alveoli (Zacharias et al., 2018). AEPs are present in the adult human lung and account for around 29% of AT2s (Zacharias et al., 2018). Whether AEPs are present and functional in immature lungs of preterm neonates has not been discovered. We found that \textit{AXIN2}(+);\textit{SFTPC}(+) cells are in fact present in the control human neonatal lungs and account for nearly 28% of the total pAT2 cell population, a remarkably conserved parallel to the mature adult human lungs (Figure 7). Intriguingly, in BPD lungs this ratio (and not the absolute number) is increased to 36.5%, likely caused by an incomplete or blunted response to injury. This increase is also observed in the mouse DKO lungs in which the overall ratio of \textit{Axin2}(+) pAT2s to total pAT2s is nearly twice as large in the mutant lungs compared with controls (20% and 10.2%, respectively) (Figure S12). However, while the relative pool of “regenerative” cells is larger, the relative number of proliferating pAT2s is smaller in BPD lungs compared with controls, again similar to the mouse DKO lungs.
Reduced PDGFRα is a primary feature of BPD (Popova et al., 2014) and may indeed be a potential cause of failed mesenchymal-pAT2 crosstalk, ultimately accounting for lack of pAT2 responsiveness in BPD. In addition, altered expression of several of the mouse ligands identified here have been found in human BPD (Alejandre-Alcazar et al., 2007; Bourbon et al., 2005; Chao et al., 2020; Lassus et al., 2003; Lecarpentier et al., 2019; Seedorf et al., 2016; Sucre et al., 2016). Collectively these data indicate that BPD may more likely be caused by failure in recruitment rather than lack of presumptive regenerative epithelial cells. The latter has obvious implications for the pathogenesis of BPD and may offer novel preventive or therapeutic strategies for this neonatal chronic lung disease.

Limitations of the study

The present study was in large part focused on genetic analysis of a compound mutation that affects TGFβ signaling in a select population of mesenchymal cells. In addition, the analyses centered around scRNA-seq and limited analysis of the protein products of the genes involved. While highly focused, such analyses may fall short of revealing the complex interactions that constitute cellular and physiologic function in the mutant lungs.

STAR METHODS

RESOURCE AVAILABILITY

Lead contact—Further information and requests for the resources and reagents should be directed to the lead contact, Parviz Minoo (minoo@usc.edu).

Materials availability—Mouse line generated in this study are available to distribute upon approval of the institutional MTA. The raw and analyzed scRNAseq data in this study are deposited in GEO database: GSE172059.

Data and code availability—The raw and analyzed scRNAseq data in this study are deposited in GEO database: GSE172059.

This paper does not report original code.

Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse breeding and genotyping—All animal studies were conducted strictly according to protocols approved by the USC Institutional Animal Care and Use Committee (IACUC) (Los Angeles, CA, USA). The mice were housed and maintained in pathogen-free conditions at constant room temperature (20–22°C), with a 12 h light/dark cycle, and free access to water and food. Gli1-creERT2, Rosa26mTmG, CAGTomato, Alk5(fox/fox) and Tbr2(flox/flox) mice were purchased from the Jackson Laboratory. CAGTomato mice express Tomato Red in presence of activated Cre (Stock No: 007909). Rosa26mTmg express GFP in presence of activate Cre (Stock No: 007676). Sftp-c-GFP mice express GFP in cells that express the surfactant protein gene C, mostly progenitors of AT2 and differentiated AT2 cells. Gli1-creERT2;Rosa26mTmg mice (control) were generated by breeding Gli1-
CRE2 and Rosa26mTmG mice. Gli1-creERT2;Rosa26mTmG; Alk5(fox/flox); Tbr2(fox/flox) (abbreviated as DKO) mice were generated by breeding control mice with the Alk5(fox/flox) and Tbr2(fox/flox) mice. Gli1-creERT2; CAGTomato mice were generated by breeding Gli1-creERT2 and CAGTomato mice. Gli1-creERT2;Alk5(fox/flox); Tbr2(fox/flox) mice were generated by breeding Gli1-creERT2; Alk5(fox/flox); Tbr2(fox/flox); with Sftpc-GFP mice.

Genotyping of the transgenic mice was performed by PCR with genomic DNA isolated from mouse tails. The primers for mouse genotyping are listed in Table S1.

**Mouse lung tissue**—Mice were euthanized by CO₂ inhalation at the time of tissue harvest. Sex of the mice was not noted. Chest cavity was exposed and lungs cleared of blood by perfusion with cold PBS via the right ventricle. Lungs were inflated with 4% formaldehyde under constant pressure of 30 cm water and allowed to fix overnight in refrigerator. Tissue was dehydrated through a series of ethanol washes after which they were embedded in paraffin and sectioned.

**3D alveolar organoid culture**—3D alveolar organoid assays were performed as originally described (Barkauskas et al., 2013) with some modifications. Briefly, Gli1CreERT2;CAGTomato pups were given tamoxifen at PN2 and the lungs were isolated at PN14. Lungs were also isolated from Sftpc-GFP pups at PN14. Single cell suspensions were made from both groups of mouse lungs as described above. GFP labelled AT2 cells and Tomato labelled mesenchymal cells were sorted out by FACs. Previous studies indicated a 1:10 mix of AT2 and mesenchymal cells as the condition best optimized for organoid’s growth. With this ratio as baseline, a combination of different number of above sorted AT2 and mesenchymal cells, were mixed in 50% Matrigel (growth factor reduced, phenol-red free) (Corning), gelled on Costar Transwell Polyester Membrane Inserts (Fisher Scientific), and cultured in air-liquid interface in basal medium (DMEM/F12, 10%FBS, 1x insulin/transferrin, 1xPSA). Fresh media was replaced every two days, and the organoid growth was monitored and documented under the inverted fluorescent microscope periodically. Ligand-receptor interactions discovered between AT2 and mesenchymal cells were functionally examined by inhibiting the indicated pathways. Inhibitors were carefully selected from literature based on their specificity, quality and effect. A search of the literature was used to determine the range of the concentration to be used for each inhibitor. A gradient of different concentrations around the range were tested, and the lowest concentration with phenotype distinct from the control was the final concentration used for the treatment in the study. Please see the Table S1 for the list of inhibitors and the final concentration used. Inhibitors were added at the time of the first media change, and new inhibitors were added upon each media change or every other day.

**Human fetal lung**—BPD and non-BPD postnatal human lung tissues were provided by the International Institute for the Advancement of Medicine and the National Disease Research Interchange, and were classified exempt from human subject regulations per the University of Rochester Research Subjects Review Board protocol (RSRB00056775). The control lung samples were from female and male neonates born at term gestation (40 weeks) who died of non-pulmonary causes. The BPD samples were from female and male neonates.
born prematurely at between 23 weeks and 27 weeks of gestation and died with a diagnosis of BPD.

**METHOD DETAILS**

**Tamoxifen administration**—A single dose of Tamoxifen (8mg/mL in peanut oil) was administered by oral gavage to neonates at postnatal day 2 (P2, 400 μg each pup) with a plastic feeding needle (Instech Laboratories, PA). Neonatal lungs were collected between P7 and P30 for morphological, immunohistochemical, cellular and molecular biological analyses.

**Immunohistochemistry**—H&E staining was performed as usual, and morphometric measurements were made using ImageJ. Immunofluorescent staining was performed as previously described using paraffin-embedded lung sections (Li et al., 2019). In brief, five micrometer (μm) tissue sections were deparaffinized, rehydrated and subjected to antigen retrieval. After blocking with normal serum, the sections were probed with primary antibodies at 4°C overnight. Combinations of fluorescein anti-mouse and Cy3 anti-rabbit or anti-goat IgG secondary antibodies (Jackson ImmunoResearch Laboratories, ING) were applied to detect specific primary antibodies. Nuclei were counterstained with 4′,6-diamidino-2-phenylindole (DAPI). Primary antibodies used and their sources are listed in the Key Resources Table. Images were made with Leica DMi8 fluorescence microscope and processed with Leica LAS X and ImageJ. MLI or mean linear intercept was determined by measurement of the mean interalveolar septal wall distance determined by the number of interruptions that four drawn lines make in each field. Five to six images (10× magnification) from each lung were used to manually count the intercepts, and at least five random fields were examined per tissue.

**RNAScope**—Samples were fixed in either 10% neutral buffered formalin, dehydrated with ethanol and embedded in paraffin wax or fixed in 4% paraformaldehyde and embedded in OCT compound. Sections from paraffin (5 μm) and OCT (20 μm) blocks were processed using standard pretreatment conditions for each per the RNAScope multiplex fluorescent reagent kit version 2 (Advanced Cell Diagnostics) assay protocol. TSA-plus fluorescein, Cy3 and Cy5 fluorophores were used at different dilution optimized for each probe. RNAScope was followed, when needed, by IF staining same as described above in immunohistochemistry. RNAScope probes used are listed in the KRT Table. Micrographs were acquired with a laser scanning confocal fluorescence microscope (Zeiss LSM780) and processed with ImageJ and Imaris.

**Mouse lung single-cell dissociation**—Single-cell suspension was prepared as described in Adam et al. (2017) with all the procedures run on ice or in cold room. Mice were euthanized and the lungs were perfused with PBS as described above. The lungs were inflated with cold active protease solution (5 mM CaCl2, 10 mg/mL Bacillus Licheniformis protease), dissected and transferred to a petri dish, and the heart, thymus and trachea were removed. The lobes were minced using a razor blade. The minced tissue was then immersed in extra cold active protease solution for 10 min and triturated using a 1 mL pipet. This Homogenate was transferred to a Miltenyi C-tube with 5mL HBSS/DNase
(Hank’s Balanced Salt Buffer) and the Miltenyi gentleMACS lung program was run twice. Subsequently, this suspension was passed through a 100um strainer, pelleted at 300 g for 6 minutes, suspended in 2mL RBC (Red Blood Cell) Lysis Buffer (BioLegend) and incubated for 2 min. At this point, 8mL HBSS was added and centrifuged again. The pellet was suspended in HBSS, filtered through a 30um strainer and pelleted and suspended in MACS separation buffer (Miltenyi Biotec) with 2% FBS for FACs and in HBSS/0.01% BSA for 10x single cell sequencing. Cell separation and viability were confirmed under the microscope and through Vi-CELL Cell Counter after staining with Trypan blue.

**Flow cytometry and cell sorting**—Single cell suspensions were gated on viability by staining with ghost dye red 780, then gated on singlets and separated on GFP from Gli1-creERT2; Rosa26rtTomG or Sftpc-GFP labeling or Tomato from Gli1-creERT2; CAGTomato labeling. Cell sorting was performed on a BD FACS Aria II at stem cell Flow Cytometry Core at Keck school of medicine USC. For cell counting, cell number and percentage as of its appropriate group were recorded and used to calculate the difference between control and mutant. For Organoid culture, cells were sorted in DMEM containing 10% FBS. For RNA, cells of interest were collected in Trizol-LS reagent (ThermoFisher). The RNA was isolated according to the manufacture’s protocol. Following RNA purification, cDNA was generated using the SuperScript IV First-Strand Synthesis System (ThermoFisher). Quantitative Real-Time PCR was then performed on a Light Cycler (Roche) or 7900HT fast real-time PCR system (Applied Biosystems) using SYBR green reagents (ThermoFisher).

**Single-cell RNA-sequencing**—We used the Chromium Single Cell 3′ kit (v3) of 10x Genomics for performing scRNA-sequencing. The single cell suspension prepared above was loaded onto a Chromium Single-Cell instrument (10x Genomics) to generate single-cell barcoded droplets (GEMs) as detailed in manufacture’s protocol. The resulting libraries were pooled and sequenced across two lanes on an Illumina HiSeq2500 instrument in High-output mode by Novogene. Reads were aligned and subsequent analyses performed using the Cell Ranger (Pipeline). We obtained 68k reads per cell with a median genes per cell of 1,017 and median UMI count per cell of 2,040. The accession number for the scRNA-seq data is GEO database: GSE172059.

**Bioinformatics**—The 10x Genomics’ Cell Ranger pipeline was used to demultiplex raw base call (BCL) files into FASTQ files, perform alignment, filtering, barcode counting, and UMI counting, combine and normalize counts from multiple samples, generate feature-barcode matrices, run the dimensionality reduction, clustering, and gene expression analysis. Quality control was done additionally in Partek Flow to filter out cells with excess mitochondrial reads and possible doublets and remove batch effects. Loupe Browser was used for data visualization and analysis including identification of differentially expressed genes and marker genes, cell type classification, cell re-clustering, cell counting, gene expression comparative analysis between different cell clusters at both cell and population levels. DE genes derived above were imported into IPA (Ingenuity Pathway Analysis from QIAGEN) for pathway and functional analysis. To construct single cell pseudo-time trajectory and to identify genes that change as the cells undergo transition, Monocle 3 algorithm was applied to the cells from clusters 4 to 11. Interactions between cell types
were predicted using a published Fantom5 Cell Connectome dataset linking ligands to their receptors (STAR Methods) (Ramilowski et al., 2015).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical analysis was performed in JMP pro 15. A two-tailed non-parametric Mann-Whitney U-test was used for the comparison between two experimental groups and a one-way ANOVA was used for multiple comparisons. Data were considered significant if \( p < 0.05 \).

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**ACKNOWLEDGMENTS**

This work was supported by the Hastings Foundation, NIH grants HL143059 (P.M.), HL144932 (C.L. and P.M.), and HL62569-10 (Z.B. and P.M.).

**REFERENCES**

Adam M, Potter AS, and Potter SS (2017). Psychrophilic proteases dramatically reduce single-cell RNA-seq artifacts: a molecular atlas of kidney development. Development 144, 3625–3632. [PubMed: 28851704]

Ahn S, and Joyner AL (2004). Dynamic changes in the response of cells to positive hedgehog signaling during mouse limb patterning. Cell 118, 505–516. [PubMed: 15315762]

Alejandre-Alcazar MA, Kwapiszewska G, Reiss I, Amarie OV, Marsh LM, Sevilla-Perez J, Wygrecka M, Eul B, Kobrich S, Hesse M, et al. (2007). Hyperoxia modulates TGF-beta/BMP signaling in a mouse model of bronchopulmonary dysplasia. Am. J. Physiol. Lung Cell Mol. Physiol 292, L537–L549. [PubMed: 17071723]

Barkauskas CE, Cronce MJ, Rackley CR, Bowie EJ, Keene DR, Stripp BR, Randell SH, Noble PW, and Hogan BL (2013). Type 2 alveolar cells are stem cells in adult lung. J. Clin. Invest 123, 3025–3036. [PubMed: 23921127]

Basil MC, Katzen J, Engler AE, Guo M, Herriges MJ, Kathiriya JJ, Windmueller R, Ysasi AB, Zacharias WJ, Chapman HA, Kotton DN, Rock JR, Snoeck HW, Yunjak-Novakovic G, Whitsett JA, and Morrissey EE (2020). The Cellular and Physiological Basis for Lung Repair and Regeneration: Past, Present, and Future. Cell Stem Cell 26, 482–502. [PubMed: 32243808]

Bostrom H, Gritli-Linde A, and Betsholtz C (2002). PDGF-A/PDGF alpha-receptor signaling is required for lung growth and the formation of alveoli but not for early lung branching morphogenesis. Dev. Dyn 223, 155–162. [PubMed: 11803579]

Bourbon J, Boucherat O, Chailley-Heu B, and Delacourt C (2005). Control mechanisms of lung alveolar development and their disorders in bronchopulmonary dysplasia. Pediatr. Res 57, 38R–46R.

Burridge PW, Matsa E, Shukla P, Lin ZC, Churko JM, Ebert AD, Lan F, Diecke S, Huber B, Mordwinkin NM, et al. (2014). Chemically defined generation of human cardiomyocytes. Nat. Methods 11, 855–860. [PubMed: 24930130]

Calvi C, Podowski M, Lopez-Mercado A, Metzger S, Misono K, Malinina A, Dikeman D, Poonyagariyon H, Ynalvez L, Derakhshandeh R, et al. (2013). Hepatocyte growth factor, a determinant of airspace homeostasis in the murine lung. PLoS Genet 9, e1003228. [PubMed: 23459311]

Cassandras M, Wang C, Kathiriya J, Tsukui T, Matatia P, Matthay M, Wolters P, Molofsky A, Sheppard D, Chapman H, and Peng T (2020). Gli1(+) mesenchymal stromal cells form a
pathological niche to promote airway progenitor metaplasia in the fibrotic lung. Nat. Cell Biol 22, 1295–1306. [PubMed: 33046884]

Chao CM, Chong L, Chu X, Shrestha A, Behnke J, Ehrhardt H, Zhang J, Chen C, and Bellusc S (2020). Targeting bronchopulmonary dysplasia-associated pulmonary hypertension (BPD-PH): potential role of the FGF signaling pathway in the development of the pulmonary vascular system. Cells 9, 1875.

Chen L, Acciani T, Le Cras T, Lutzko C, and Perl AK (2012). Dynamic regulation of platelet-derived growth factor receptor alpha expression in alveolar fibroblasts during realveolarization. Am J Respir Cell Mol Biol 47, 517–527. [PubMed: 22652199]

Chen L, Chen Q, Rong P, Wang HY, and Chen S (2017). The energy sensing LKB1-AMPKα pathway regulates IGF1 secretion and consequent activation of the IGF1R-PKB pathway in primary hepatocytes. FEBS J. 284, 2096–2109. [PubMed: 28500773]

Chung MI, Bujnis M, Barkauskas CE, Kobayashi Y, and Hogan BLM (2018). Niche-mediated BMP/SMAD signaling regulates lung alveolar stem cell proliferation and differentiation. Development 145, ev163014. [PubMed: 29752282]

Clement MS, Gammelgaard KR, Nielsen AL, and Sorensen BS (2020). Epithelial-to-mesenchymal transition is a resistance mechanism to sequential MET-TKI treatment of MET-amplified EGFR-TKI resistant non-small cell lung cancers. Transl. Lung Cancer Res. 9, 1904–1914. [PubMed: 33209611]

De Simone A, Evanitsky MN, Hayden L, Cox BD, Wang J, Tornini VA, Ou J, Chao A, Poss KD, and Di Talia S (2021). Control of osteoblast regeneration by a train of Erk activity waves. Nature 590, 129–133. [PubMed: 33408418]

Desai TJ, Brownfield DG, and Krasnow MA (2014). Alveolar progenitor and stem cells in lung development, renewal and cancer. Nature 507, 190–194. [PubMed: 24499815]

Frank DB, Peng T, Zepp JA, Snitow M, Vincent TL, Penkala JI, Cui Z, Herriges MJ, Morley MP, Zhou S, et al. (2016). Emergence of a wave of Wnt signaling that regulates lung alveologenesis by controlling epithelial self-renewal and differentiation. Cell Rep. 17, 2312–2325. [PubMed: 27880906]

Frazier NM, Brand T, Gordan JD, Grandis J, and Jura N (2019). Overexpression-mediated activation of MET in the Golgi promotes HER3/ERBB3 phosphorylation. Oncogene 38, 1936–1950. [PubMed: 30390071]

Ghosh MC, Gorantla V, Makena PS, Luellen C, Sinclair SE, Schwingshackl A, and Waters CM (2013). Insulin-like growth factor-I stimulates differentiation of ATII cells to ATI-like cells through activation of Wnt5a. Am. J. Physiol. Lung Cell Mol. Physiol 305, L222–L228. [PubMed: 23709620]

Gong W, Zhang N, Cheng G, Zhang Q, He Y, Shen Y, Zhang Q, Zhu B, Zhang Q, and Qin L (2019). Rehmamnia glutinosa libosch extracts prevent bone loss and architectural deterioration and enhance osteoblastic bone formation by regulating the IGF-1/PI3K/mTOR pathway in Streptozotocin-induced diabetic rats. Int. J. Mol. Sci 20, 3964.

Graf J, Alexander S, Peterson JR, Snider TN, Levi B, Lee B, and Mishina Y (2018). TGF-beta Family Signaling in Mesenchymal Differentiation. Cold Spring Harb Perspect Biol 10.

Green J, Endale M, Auer H, and Perl AK (2016). Diversity of interstitial lung fibroblasts is regulated by platelet-derived growth factor receptor alpha kinase activity. Am. J. Respir. Cell Mol. Biol 54, 532–545. [PubMed: 26414960]

Guo M, Du Y, Gokey JJ, Ray S, Bell SM, Adam M, Sudha P, Perl AK, Deshmukh H, Potter SS, Whitsett JA, and Xu Y (2019). Single cell RNA analysis identifies cellular heterogeneity and adaptive responses of the lung at birth. Nat Commun 10, 37. [PubMed: 30604742]

Hagan AS, Zhang B, and Ornitz DM (2020). Identification of a FGF18-expressing alveolar myofibroblast that is developmentally cleared during alveologenesis. Development 147, dev181032. [PubMed: 31862844]

Hu Y, Ng-Bleichfeldt JP, Ota C, Ciminieri C, Ren W, Hiemstra PS, Stolk J, Gosens R, and Konigshoff M (2020). Wnt/beta-catenin signaling is critical for regenerative potential of distal lung epithelial progenitor cells in homeostasis and emphysema. Stem Cells 38, 1467–1478. [PubMed: 32526076]
Huaqi Y, Caipeng Q, Qiang W, Yiqing D, Xiang D, Xu T, Xiaowei Z, Qing L, Shijun L, and Tao X (2019). Transcription factor SOX18 promotes clear cell renal cell carcinoma progression and alleviates cabozantinib-mediated inhibitory effects. Mol. Cancer Ther 18, 2433–2445. [PubMed: 31527225]

Imberti B, Morigi M, Tomasoni S, Rotta C, Corna D, Longaretti L, Rottoli D, Valscerci F, Benigni A, Wang J, et al. (2007). Insulin-like growth factor-1 sustains stem cell mediated renal repair. J. Am. Soc. Nephrol 18, 2921–2928. [PubMed: 17942965]

Jin K, Ruan L, Pu J, Zhong A, Wang F, Tan S, Huang H, Mu J, and Yang G (2018). Metformin suppresses growth and adrenocorticotropic hormone secretion in mouse pituitary corticotroph tumor AtT20cells. Mol. Cell Endocrinol 478, 53–61. [PubMed: 30025915]

Kaartinen V, Voncken JW, Shuler C, Warburton D, Bu D, Heisterkamp N, and Groffen J (1995). Abnormal lung development and cleft palate in mice lacking TGF-beta 3 indicates defects of epithelial-mesenchymal interaction. Nat. Genet 11, 415–421. [PubMed: 7493022]

Kiyokawa H, and Morimoto M (2020). Notch signaling in the mammalian respiratory system, specifically the trachea and lungs, in development, homeostasis, regeneration, and disease. Dev. Growth Differ 62, 67–79. [PubMed: 31613406]

Kobayashi Y, Tata A, Konkimala A, Katsura H, Lee RF, Ou J, Banovich NE, Kropski JA, and Tata PR (2020). Persistence of a regeneration-associated, transitional alveolar epithelial cell state in pulmonary fibrosis. Nat Cell Biol 22, 934–946. [PubMed: 32661339]

Kong LR, Mohamed Salleh NAB, Ong RW, Tan TZ, Syn NL, Goh RM, Fhu CW, Tan DSW, Iyer NG, Kannan S, et al. (2020). A common MET polymorphism harnesses HER2 signaling to drive aggressive squamous cell carcinoma. Nat. Commun 11, 1556. [PubMed: 32214092]

Kuang J, Zhai Z, Li P, Shi R, Guo W, Yao Y, Guo J, Zhao G, He J, Xu S, et al. (2021). SS18 regulates pluripotent-somatic transition through phase separation. Nat. Commun 12, 4090. [PubMed: 34215745]

Lassus P, Heikkipaila P, Andersson LC, von Boguslawski K, and Andersson S (2003). Lower concentration of pulmonary hepatocyte growth factor is associated with more severe lung disease in preterm infants. J. Pediatr 143, 199–202. [PubMed: 12970632]

Lecarpentier Y, Gourrier E, Gobert V, and Vallee A (2019). Bronchopulmonary dysplasia: crosstalk between PPARgamma, WNT/beta-Catenin and TGF-beta pathways; the potential therapeutic role of PPARgamma agonists. Front. Pediatr 7, 176. [PubMed: 31131268]

Li C, Lee MK, Gao F, Webster S, Di H, Duan J, Yang CY, Bhopal N, Peinado N, Pryhuber G, et al. (2019). Secondary crest myofibroblast PDGFRalpha controls the elastogenesis pathway via a secondary tier of signaling networks during alveologenesis. Development 146, dev176354. [PubMed: 31331942]

Li C, Li M, Li S, Xing Y, Yang CY, Li A, Borok Z, De Langhe S, and Minoo P (2015). Progenitors of secondary crest myofibroblasts are developmentally committed in early lung mesoderm. Stem Cells 33, 999–1012. [PubMed: 25448080]

Li R, Bernau K, Sandbo N, Gu J, Preisel S, and Sun X (2018). Pdgfra marks a cellular lineage with distinct contributions to myofibroblasts in lung maturation and injury response. Elife 7, e36865. [PubMed: 30178747]

Liu L, Charville GW, Cheung TH, Yoo B, Santos PJ, Schroeder M, and Rando TA (2018). Impaired Notch signaling leads to a decrease in p53 activity and mitotic catastrophe in aged muscle stem cells. Cell Stem Cell 23, 544–556 e544. [PubMed: 30244867]

Manchado E, Weissmueller S, Morris J.P.t., Chen CC, Wullenkord R, Lujambio A, de Stanchina E, Poirier JT, Gainor JF, Corcoran RB, et al. (2016). A combinatorial strategy for treating KRAS-mutant lung cancer. Nature 534, 647–651. [PubMed: 27338794]

Maynard A, McCouch CE, Rotow JK, Harris L, Haderk F, Kerr DL, Yu EA, Schenk EL, Tan W, Zee A, et al. (2020). Therapy-induced evolution of human lung cancer revealed by single-cell RNA sequencing. Cell 182, 1232–1251 e1222. [PubMed: 32822576]

McGowan SE, and McCoy DM (2014). Regulation of fibroblast lipid storage and myofibroblast phenotypes during alveolar septation in mice. Am J Physiol Lung Cell Mol Physiol 307, L618–L631. [PubMed: 25150063]
Nabhan AN, Brownfield DG, Harbury PB, Krasnow MA, and Desai TJ (2018). Single-cell Wnt signaling niches maintain stemness of alveolar type 2 cells. Science 359, 1118–1123. [PubMed: 29420258]

Nakamura H, Arai Y, Totoki Y, Shiratori T, Elzawahry A, Kato M, Hama N, Hosoda F, Urunishi T, Ohashi S, et al. (2015). Genomic spectra of biliary tract cancer. Nat. Genet 47, 1003–1010. [PubMed: 26258846]

Oh JW, Oh YJ, Han S, Her NG, and Nam DH (2021). High-content analysis-based sensitivity prediction and novel therapeutics screening for c-met-addicted Glioblastoma. Cancers (Basel) 13, 372. [PubMed: 33498427]

Ohmichi H, Koshimizu U, Matsumoto K, and Nakamura T (1998). Hepatocyte growth factor (HGF) acts as a mesenchyme-derived morphogenic factor during fetal lung development. Development 125, 1315–1324. [PubMed: 9477330]

Park HR, Ahn YO, Kim TM, Kim S, Kim S, Lee-Y, Kim M, Keam B, Kim DW, and Heo DS (2019). NK92-CD16 cells are cytotoxic to non-small cell lung cancer cell lines that have acquired resistance to tyrosine kinase inhibitors. Cytotherapy 21, 603–611. [PubMed: 31010733]

Pieretti AC, Ahmed AM, Roberts JD Jr., and Kelleher CM (2014). A novel in vitro model to study alveologenesis. Am. J. Respir. Cell Mol. Biol 50, 459–469. [PubMed: 24066869]

Popova AP, Bentley JK, Cui TX, Richardson MN, Linn MJ, Lei J, Chen Q, Goldsmith AM, Pryhuber GS, and Hershenson MB (2014). Reduced platelet-derived growth factor receptor expression is a primary feature of human bronchopulmonary dysplasia. Am. J. Physiol. Lung Cell Mol. Physiol 307, L231–L239. [PubMed: 24907056]

Qiu Z, Li H, Zhang Z, Zhu Z, He S, Wang X, Wang P, Qin J, Zhuang L, Wang W, et al. (2019). A pharmacogenomic landscape in human liver cancers. Cancer Cell 36, 179–193 e111. [PubMed: 31378681]

Ramilowski JA, Goldberg T, Harshbarger J, Kloppmann E, Lizio M, Satagopam VP, Itoh M, Kawai H, Carninci P, Rost B, and Forrest AR (2015). A draft network of ligand-receptor-mediated multicellular signaling in human. Nat. Commun 6, 7866. [PubMed: 26198319]

Resh MD (2016). Fatty acylation of proteins: the long and the short of it. Prog. Lipid Res 63, 120–131. [PubMed: 27233110]

Riemondy KA, Jansing NL, Jiang P, Redente EF, Gillen AE, Fu R, Miller AJ, Spence JR, Gerber AN, Hesselberth JR, and Zemans RL (2019). Single cell RNA sequencing identifies TGF-beta as a key regenerative cue following LPS-induced lung injury. JCI Insight 5.

Rognoni E, Widmaier M, Jakobson M, Ruppert R, Ussar S, Katsougkri D, Bottcher RT, Lai-Cheong JE, Rifkin DB, McGrath JA, and Fassler R (2014). Kindlin-1 controls Wnt and TGF-beta availability to regulate cutaneous stem cell proliferation. Nat. Med 20, 350–359. [PubMed: 24681597]

Sahni M, and Mowes AK (2020). Bronchopulmonary Dysplasia. In StatPearls [Internet] (Treasure Island (FL): StatPearls Publishing), pp. 3–15.

Seedorf G, Metoxen AJ, Rock R, Markham N, Ryan S, Vu T, and Abman SH (2016). Hepatocyte growth factor as a downstream mediator of vascular endothelial growth factor-dependent preservation of growth in the developing lung. Am. J. Physiol. Lung Cell Mol. Physiol 310, L1098–L1110. [PubMed: 27036872]

Strunz M, Simon LM, Ansari M, Kathiriya JJ, Angelidis I, Mayr CH, Tsidiridis G, Lange M, Mattner LF, Yee M, Ogar P, Sengupta A, Kukhtevich I, Schneider R, Zhao Z, Voss C, Stoeger T, Neumann JHL, Hilgendorff A, Behr J, O’Reilly M, Lehmann M, Burgstaller G, Konigshoff M, Chapman HA, Theis FJ, and Schiller HB (2020). Alveolar regeneration through a Krt8+ transitional stem cell state that persists in human lung fibrosis. Nat Commun 11, 3559. [PubMed: 32678092]

Sucre JM, Wilkinson D, Vijayaraj P, Paul M, Dunn B, Alva-Ornelas JA, and Gomperts RN (2016). A three-dimensional human model of the fibroblast activation that accompanies bronchopulmonary dysplasia identifies Notch-mediated pathophysiology. Am. J. Physiol. Lung Cell Mol. Physiol 310, L889–L898. [PubMed: 26968771]

Tanwar V, Bylund JB, Hu J, Yan J, Walthall JM, Mukherjee A, Heaton WH, Wang WD, Potet F, Rai M, et al. (2014). Gremlin 2 promotes differentiation of embryonic stem cells to atrial fate by activation of the JNK signaling pathway. Stem Cells 32, 1774–1788. [PubMed: 24648383]
Treutlein B, Brownfield DG, Wu AR, Neff NF, Mantalas GL, Espinoza FH, Desai TJ, Krasnow MA, and Quake SR (2014). Reconstructing lineage hierarchies of the distal lung epithelium using single-cell RNA-seq. Nature 509, 371–375. [PubMed: 24739965]

Tsao PN, Matsuoka C, Wei SC, Sato A, Sato S, Hasegawa K, Chen HK, Ling TY, Mori M, Cardoso WV, and Morimoto M (2016). Epithelial Notch signaling regulates lung alveolar morphogenesis and airway epithelial integrity. Proc. Natl. Acad. Sci. U S A 113, 8242–8247. [PubMed: 27364009]

van den Brink SC, Alemany A, van Batenburg V, Moris N, Blotenburg M, Vivie J, Baillie-Johnson P, Nichols J, Sonnen KF, Martinez Arias A, and van Oudenaarden A (2020). Single-cell and spatial transcriptomics reveal somitogenesis in gastruloids. Nature 582, 405–409. [PubMed: 32076263]

Wang G, Yin L, Peng Y, Gao Y, Gao H, Zhang J, Lv N, Miao Y, and Lu Z (2019). Insulin promotes invasion and migration of KRAS(G12D) mutant HPNE cells by upregulating MMP-2 gelatinolytic activity via ERK- and PI3K-dependent signalling. Cell Prolif. 52, e12575. [PubMed: 30838710]

Wang Y, Tang Z, Huang H, Li J, Wang Z, Yu Y, Zhang C, Li J, Dai H, Wang F, et al. (2018). Pulmonary alveolar type I cell population consists of two distinct subtypes that differ in cell fate. Proc. Natl. Acad. Sci. U S A 115, 2407–2412. [PubMed: 29463737]

Wei Y, Wang T, Ma L, Zhang Y, Zhao Y, Lye K, Xiao L, Chen C, Wang Z, Ma Y, et al. (2021). Efficient derivation of human trophoblast stem cells from primed pluripotent stem cells. Sci. Adv 7, eabf4416. [PubMed: 34380613]

Wimmer RA, Leopoldi A, Aichinger M, Wick N, Hantusch B, Novatchkova M, Taubenschmid J, Hammerle M, Esk C, Bagley JA, et al. (2019). Human blood vessel organoids as a model of diabetic vasculopathy. Nature 565, 505–510. [PubMed: 30651639]

Wong GS, Zhou J, Liu JB, Wu Z, Xu X, Li T, Xu D, Schumacher SE, Puschhof J, McFarland J, et al. (2018). Targeting wild-type KRAS-amplified gastroesophageal cancer through combined MEK and SHP2 inhibition. Nat. Med 24, 968–977. [PubMed: 29808010]

Xu K, Nieuwenhuis E, Cohen BL, Wang W, Canty AJ, Danska JS, Coultas L, Rossant J, Wu MY, Piscione TD, et al. (2010). Lunatic Fringe-mediated Notch signaling is required for lung alveologenesis. Am. J. Physiol. Lung Cell Mol. Physiol 298, L45–L56. [PubMed: 19897741]

Yang J, Hernandez BJ, Martinez Alanis D, Narvaez del Pilar O, Vila-Ellis L, Akizawa H, Evans SE, Ostrin EJ, and Chen J (2016). The development and plasticity of alveolar type 1 cells. Development 143, 54–65. [PubMed: 26586225]

Zacharias WJ, Frank DB, Zepp JA, Morley MP, Alkhaleel FA, Kong J, Zhou S, Cantu E, and Morrisey EE (2018). Regeneration of the lung alveolus by an evolutionarily conserved epithelial progenitor. Nature 555, 251–255. [PubMed: 29489752]

Zeng YA, and Nusse R (2010). Wnt proteins are self-renewal factors for mammary stem cells and promote their long-term expansion in culture. Cell Stem Cell 6, 568–577. [PubMed: 20569694]

Zepp JA, and Morrisey EE (2019). Cellular crosstalk in the development and regeneration of the respiratory system. Nat. Rev. Mol. Cell Biol 20, 551–566. [PubMed: 31217577]

Zepp JA, Zacharias WJ, Frank DB, Cavanaugh CA, Zhou S, Morley MP, and Morrisey EE (2017). Distinct mesenchymal lineages and niches promote epithelial self-renewal and myofibrogenesis in the lung. Cell 170, 1134–1146 e1110. [PubMed: 28886382]

Zhang K, Wang A, Zhong K, Qi S, Wei C, Shu X, Tu WY, Xu W, Xiao C, Xiao Y, et al. (2021). UBQLN2-HSP70 axis reduces poly-Gly-Ala aggregates and alleviates behavioral defects in the C9ORF72 animal model. Neuron 109, 1949–1962 e1946. [PubMed: 33991504]
Highlights

- Lung alveolar type 2 progenitors (pAT2s) are heterogeneous during alveologenesis
- Blocking TGFβ in SCMF fibroblasts reduces their number and blocks alveologenesis
- SCMF signaling regulates proliferation of unique pAT2 population in alveologenesis
- An evolutionarily conserved mechanism appears to govern pAT2s in neonatal human lungs
Figure 1. Postnatal inactivation of TGFβ receptors in Hedgehog-responsive SCMFs arrests alveologenesis

Newborn Gli1-creERT2; Alk5(flox/flox);TβR2(flox/flox) (DKO) and control mice received tamoxifen on PN2, and lungs were analyzed between PN7 and PN45.

(A–D) Histology of control and mutant lungs at PN14 (A, B) and PN45 (C, D). Scale bar, 50 μm.

(E and F) Average of mean linear intercept (MLI) for control versus mutant lungs on PN14 (E) and PN45 (F).

(G) MLI of control and mutant lungs over time from PN7 to PN45.

(H) Fold change in Hopx mRNA between control and mutant lungs.

(I) Ratio of HOPX(+) cells to total cells in control versus mutant lungs.

(J and K) Immunofluorescent (IF) staining for HOPX (red) using PN14 lung sections from control (J) and mutant (K) mice.

(L and M) IF staining for PDPN (red) and SFTPC (green) using PN14 lung sections from control (L) and mutant (M) mice. DAPI (blue) was used to visualize all cells. White arrows point to pAT2s and yellow arrowheads to AT1s. Scale bar, 50μm (J&K), 20μm (L&M).

(N) Ratio of SFTPC(+) cells per total lung cells in control versus mutant lungs.

(O) Fold change in Sftp3 mRNA between control and mutant lungs.
(P) FACS-based quantification of GFP-labeled pAT2 cells isolated from
Sftpc-GFP;Alk5floxflox;Tbr2floxflox (control) compared with Gli1-creERT2;Sftpc-
GFP;Alk5floxflox;Tbr2floxflox (mutant).

(Q) FACS-based quantification of GFP-labeled SCMFs isolated from Gli1-
creERT2;ROSA26mTmG (control) compared with Gli1-creERT2;ROSA26mTmG;
Alk5floxflox;Tbr2floxflox (mutant).

All mice were treated with tamoxifen at PN2 and analyzed on PN14. Data are presented as
mean+/−SD (E-I,N,O) and boxplots (P,Q) with n ≥ 3 biological replicates. CONT, control; MUT, mutant.
Figure 2. Defining the three distinct mesenchymal subclusters in PN14 lungs
(A) tSNE representation of reclustered MECs data revealing three distinct subclusters, C1, C2, and C3, in control versus mutant lungs. The number of cells in each subcluster and their alterations in the mutant lung (PS, population shift) are shown in the table on the right.
(B and C) Heatmaps of cell-type-enriched transcripts in each of the three MEC subclusters in control (B) and mutant (C) lungs.
(D) tSNE representation of select enriched transcripts for C3, Pdgfra, Inmt, Fn1, and Mfap4, and violin plot representations of their abundance and differential expression.
Figure 3. Defining distinct epithelial subclusters in PN14 EPCs

(A) tSNE representation of reclustered EPCs data based on Sftpc expression, showing eight distinct subclusters, C4 to C11, in both control and mutant lungs.

(B) Alterations in cell number in each of the eight subclusters as a fraction of total lung cells, or total epithelial cells between control and mutant lungs. PS, population shift.

(C and D) Heatmaps of enriched transcripts in each of the control and mutant subclusters and their overlap with alveolar epithelial cell lineage signature genes derived from LungGens Consortium.

(E) tSNE representation of select enriched transcripts for C4 (Mki67, Top2a, Cdk1, and Spc24) and their violin plot abundance and differential expression in C4 to C11.
Figure 4. Transitional cells are scattered throughout the epithelial cell population in PN14 lungs

(A) tSNE representation of the pAT2 signature gene transcripts Sftpc and Lamp3 in control and mutant epithelial subclusters C4 to C11 (a, b, e, f) and their abundance (c, d, g, h).

(B) tSNE representation of the AT1 signature gene transcripts Aqp5 and Ager in control and mutant Sftpc(+) subclusters C4 to C11 (i, j, m, n) and their abundance (k, l, o, p).

(C) RNAscope using DAPI (a) and probes for Sftpc (b), Aqp5 (c), and their merged images (d) (n = 3). Scale bar, 10 μm.

(D) Heatmap of metabolic-related transcripts showing low-level expression of genes in C9 and particularly C10.
Figure 5. Differentially expressed ligands in C3 and localization of select ligands by RNAscope and ligand-receptor coupling in PN14 lungs
(A) tSNE representation of eight identified signaling ligands enriched in C3 and DE (FC > 2 and p ≤0.05) between control and mutant lungs.
(B) RNAscope analysis of select ligand transcripts, Wnt2, Fgf10, Fgf2, and Igf1, and their overlap with Pdgfra (arrows in c, f, i, and l) in PN14 control lung samples (n = 3). Scale bar, 10 μm.
(C) Ligand-receptor coupling diagram and the six principal signaling pathways they represent.
Figure 6. Functional examination of candidate ligands in 3D organoid cultures and localization of Wnt2-expressing cells in PN14 lungs

(A) GFP(+) pAT2s were isolated from Sftpc-Gfp PN14 lungs and cultured with PN14 mesenchymal cells as described in STAR Methods. 3D cultures were seeded with 1,250 GFP(+) pAT2 cells and 50K mesenchymal cells and were examined on day 28 for direct GFP(+) cells. Ligand-receptor interactions between pAT2 and mesenchymal cells identified in Figure 5 were functionally examined by inhibiting the indicated pathways as shown. Inhibitors were used to block HGF (capmatinib, 1.0 μM), BMP (recombinant GREM2, 100 ng/mL), NOTCH (DAPT, 25 μM), FGF (BDJ398, 1 μM), WNT (XA V939, 1 μM) and IGF1.
(PPP, 4 μM). (a1-a7) Photomicrographs of representative organoids in each culture. (a8) quantification of the number of clones (clonogenicity). (a9) Quantification of organoid size. Scale bar, 1mm (a1-a7).

(B) (b1 and b2) tSNE representation of Wnt2 and Fgfr2 transcript distribution in control lung. (b3–b8) RNAseq analysis showing DAPI alone (b3), Mki67 (b4), Fgfr2 (b5), Wnt2 (b6), Fgfr2/Mki67 merge (b7), and Wnt2/Mki67 merge (b8). Red arrows show Wnt2(+) mesenchymal cells juxtaposed to Mki67(+) pAT2 cells, shown by green arrows. Scale bar, 10 μm (b3–b8).

(C) Modeling impact of cell population changes on neonatal pAT2 self-renewal. Different ratios of GFP(+) pAT2 cells and mesenchymal cells from PN14 control lungs were cultured as described in STAR Methods. Cell ratios are shown above each panel. 3D cultures were examined on day 21 for direct GFP(+) cells. (c1–c6) Increasing number of GFP(+) pAT2 cells cultured with 50K mesenchymal cells. (c7–c12) 5K GFP(+) pAT2 cells cultured with increasing number of mesenchymal cells as shown. (c13 and c15) Quantification of the number and size of the colonies respectively formed in c1–c6. (c14 and c16) Quantification of the number and size of the colonies, respectively formed in c7–c12. For each culture, n = 3. Scale bar, 1 mm (c1–c12). Data are presented as boxplots with n=4 (a8) and n=6 (c13,c14).
Figure 7. Regenerative pAT2s in human BPD and non-BPD lungs

(A–D) Combination IF for SFTPC (green) and RNAscope for AXIN2 (red) used to identify regenerative SFTPC(+)::AXIN2(+) double-positive cells. (A) Representative control human lung sample from a full-term neonate who died of non-pulmonary causes. (B) High magnification of lung shown in (A). Arrowheads show SFTPC(+) cells and arrows show SFTPC(+)::AXIN2(+) double-positive cells. (C) Representative BPD lung sample from a male preterm neonate born at 23 weeks gestation who died with BPD. (D) High magnification of lung shown in (C).

(E–H) Combination IF for SFTPC (red) and RNAscope for MKI67 (white) used to identify and quantify proliferative AT2s. (E) Representative control human lung sample from a full-term neonate who died of non-pulmonary causes. (F) Higher magnification of the same lung as shown in (E). Arrowheads show SFTPC(+) and arrows show SFTPC(+):MKI67(+) double-positive cells. (G) Representative BPD lung from a male preterm neonate born at 24 weeks gestation who died with BPD. (H) High magnification of lung shown in (G).

(I) Quantification of the number of SFTPC(+)::AXIN2(+) double-positive cells as a fraction of total SFTPC(+) cells.

(J) Quantification of SFTPC(+)::MKI67(+) double-positive cells as a fraction of total SFTPC(+) cells. n = 4 biological and 15–20 experimental replicates. Scale bars, 40 μm (A, C, F, H, G), 10 μm (B, D), and 100 μm (E). Data are presented as boxplots with n=4 (I) and n=5 (J).
## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| GFP                 | Santa Cruz | Cat#: SC-9996; RRID:AB_627695 |
| SFTPC               | EMD Millipore Corp. | Cat #: AB3786; RRID:AB_91588 |
| HOPX                | Santa Cruz | Cat #: SC-30216; RRID:AB_2120833 |
| MKI67               | R&D System | Cat #: AF7649; RRID:AB_2687500 |
| PDGFRA              | Cell Signaling | Cat #: 3174 |
| ACTA2               | Abcam | Cat#: AB5694; RRID:AB_2223021 |
| AQP5                | Alomone | Cat #: AQP-005; RRID:AB_2039736 |
| SFTPC (Human Tissue) | LSBio | Cat #: LS-B10952 |
| Ki67 (Human Tissue) | Cell Signaling | Cat #: 9449T; RRID:AB_2715512 |
| **Biological samples** |        |            |
| Human fetal lungs  | The International Institute for the Advancement of Medicine and the National Disease Research Interchange | [https://ndiresource.org/](https://ndiresource.org/) |
| **Chemicals, peptides, and recombinant proteins** |        |            |
| Tamoxifen           | Sigma | Catalog No. T5648 |
| Capmatinib          | Selleckchem | Catalog No. S2788 |
| Grem2               | R&D Systems | Catalog No. 2069-PR-050 |
| DAPT                | Selleckchem | Catalog No. S2215 |
| BGJ398              | Selleckchem | Catalog No. S2183 |
| XAV939              | Selleckchem | Catalog No. S1180 |
| PPP                 | Selleckchem | Catalog No. S7668 |
| **Critical commercial assays** |        |            |
| Chromium Single Cell 3’ Library & Gel Bead Kit v3 | 10x Genomics | Catalog No. 1000092 |
| Single Cell In-Drop RNA-sequencing | Novogene Corporation | GEO database: GSE172059 |
| RNAscope® Multiplex Fluorescent Reagent Kit V2 | Advanced Cell Diagnostics | Catalog No. 323100 |
| **Deposited data** |        |            |
| Raw and analyzed scRNAseq data | This paper | GEO database: GSE172059 |
| Inhibitor concentrations tested and used in organoid culture | This paper | Table S1 |
| **Experimental models: Organisms/strains** |        |            |
| Gli1-cre<sup>ERT2</sup> | Jackson Laboratories | Catalog No. 007913 |
| Rosa26<sup>TmG</sup> | Jackson Laboratories | Catalog No. 007676 |
| CAG<sup>Tomato</sup> | Jackson Laboratories | Catalog No. 007914 |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Alk5 flox/flox      | Jackson Laboratories | Catalog No. 028701 |
| Tbr2 flox/flox      | Jackson Laboratories | Catalog No. 012603 |

**Oligonucleotides**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Sftpc-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 314101-C1 |
| Hopx-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 405161-C3 |
| Agplf5-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 430021-C2 |
| Fgfr1-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 316851-C1 |
| Pdfrα-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 480661-C2 |
| Wnt2-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 313601-C1 |
| Kgr10-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 446371-C1 |
| Fgf2-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 443501-C2 |
| Mifap4-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 421391-C1 |
| MKI67-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 416771-C3 |
| AXIN2-RNAscope probe | Advanced Cell Diagnostics | Catalog No. 400241-C2 |
| RNAscope® 3-plex Positive Control | Advanced Cell Diagnostics | Catalog No. 320881 |
| RNAscope® 3-plex Negative Control | Advanced Cell Diagnostics | Catalog No. 320871 |
| Primers for mouse strain genotyping | This paper | Table S1 |
| Primers for quantitative RT-PCR | This paper | Table S1 |

**Software and algorithms**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Cell Ranger R kit   | 10x Genomics | https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/krkit |
| Loupe Cell Browser  | 10x Genomics | https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/latest#loupe |
| Partek Flow 9.0.20.1104 | Partek Inc | https://www.partek.com/partek-flow/ |
| Monocle3            | Trapnell Lab | https://cole-trapnell-lab.github.io/monocle3/ |
| IPA                 | Qiagen | https://digitalinsights.qiagen.com/products-overview/discovery-insights-portfolio/analysis-and-visualization/qiagen-ipa/ |
| Image J             | NIH | https://imagej.nih.gov/ij/ |
| STAR 2.5            | PMCID: PMC3530905 | https://github.com/alexdoabin/STAR |
| JMP pro 15          | Statistical Discovery | https://www.jmp.com/en_us/software/predictive-analytics-software.html |
| Fantom5 Cell Connectome | FANTOM5 project | https://fantom.gsc.riken.jp/5/suppl/Ramilowski_et_al_2015/vis/#have |
| Imaris              | BitPlane | http://www.bitplane.com/imaris/imaris |
| R 3.2               | R Project | https://www.r-project.org/ |
| LAS X               | Leica | https://www.leica-microsystems.com/products/microscope-software/pl/leica-las-x-ls/ |