HDAC3-Dependent Epigenetic Pathway Controls Lung Alveolar Epithelial Cell Remodeling and Spreading via miR-17-92 and TGF-β Signaling Regulation

Graphical Abstract

Highlights
- Hdac3 is important for AT1 cell spreading and remodeling during lung sacculation
- Hdac3 represses miR-17-92, which in turn targets components of TGF-β signaling
- Proper levels of TGF-β signaling are crucial for lung AT1 cell remodeling

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In Brief
How the pulmonary system develops the immense surface area required for mammalian respiration is unclear. Wang et al. show that an Hdac3-mediated pathway regulates TGF-β signaling, which is required for the spreading of distal lung epithelial cells, generating the large surface area required for efficient gas exchange.

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HDAC3-Dependent Epigenetic Pathway Controls Lung Alveolar Epithelial Cell Remodeling and Spreading via miR-17-92 and TGF-β Signaling Regulation

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SUMMARY

The terminal stages of pulmonary development, called sacculation and alveologenesis, involve both differentiation of distal lung endoderm progenitors and extensive cellular remodeling of the resultant epithelial lineages. These processes are coupled with dramatic expansion of distal airspace and surface area. Despite the importance of these late developmental processes and their relation to neonatal respiratory diseases, little is understood about the molecular and cellular pathways critical for their successful completion. We show that a histone deacetylase 3 (Hdac3)-mediated epigenetic pathway is critical for the proper remodeling and expansion of the distal lung saccules into primitive alveoli. Loss of Hdac3 in the developing lung epithelium leads to a reduction of alveolar type 1 cell spreading and a disruption of lung sacculation. Hdac3 represses miR-17-92 expression, a microRNA cluster that regulates transforming growth factor β (TGF-β) signaling. De-repression of miR-17-92 in Hdac3-deficient lung epithelium results in decreased TGF-β signaling activity. Importantly, inhibition of TGF-β signaling and overexpression of miR-17-92 can phenocopy the defects observed in Hdac3 null lungs. Conversely, loss of miR-17-92 expression rescues many of the defects caused by loss of Hdac3 in the lung. These studies reveal an intricate epigenetic pathway where Hdac3 is required to repress miR-17-92 expression to allow for proper TGF-β signaling during lung sacculation.

INTRODUCTION

The saccular stage of lung development, which extends from approximately embryonic day 16.5 (E16.5) to E18.5 of mouse gestation, is a pivotal step when the distal airspace saccules are generated as a first step toward alveologenesis. Disruption of this process can lead to serious diseases such as bronchopulmonary dysplasia in neonates. Lung sacculation and alveologenesis involve dramatic changes in the architecture and cellular composition of the distal airways. Prior to sacculation, the narrowed distal airway tubules are lined with epithelial progenitor cells that are cuboidal in shape and express markers such as Sox9 and Id2 (Rawlins et al., 2009). By E17.5 a wave of airspace expansion and alveolar epithelial differentiation occurs at the bronchoalveolar junction, which then progresses toward the distal airway tip at E18.5 (Desai et al., 2014; Treutlein et al., 2014). This results in the differentiation of two major alveolar epithelial cell lineages; the flat squamous alveolar type I (AT1) cells and the small cuboidal alveolar type II (AT2) cells. After specification, AT1 cells spread extensively and cover approximately 95% of the luminal surface of alveoli. While earlier stages of lung development including branching morphogenesis have become relatively well understood in recent studies, far less is known about sacculation and alveologenesis in the lung. In particular, how AT1 cells remodel and form the extensive surface area to mediate efficient oxygen diffusion is unclear.

Recent evidence has begun to shed light on the role of histone deacetylases (Hdacs) during lung endoderm progenitor specification (Wang et al., 2013). The class I Hdacs, Hdac1 and Hdac2, are required for development of early Sox2+ proximal lung endoderm progenitors, through regulation of Bmp4 and cell-cycle regulators including Rb1 (Wang et al., 2013). However, what roles other class I Hdacs including Hdac3 play in lung development and homeostasis has remained unclear. Importantly, Hdac3 associates with the NCoR/SMRT
complex whereas Hdacs 1 and 2 associate with complexes such as NuRD/Sin3a (Guenther et al., 2000, 2001; Li et al., 2000; Zhang et al., 1997), suggesting potentially different roles for these Hdacs and chromatin remodeling complexes during lung development.

In this report, we show that Hdac3-mediated transcriptional regulation is required for the formation of distal alveolar saccules and early lung alveologenesis. Hdac3 acts in a cell-autonomous manner to regulate AT1 cell spreading, a process required for formation of the distal alveoli, without affecting specification or early differentiation of this lineage. Loss of Hdac3 results in derepression of two major microRNA (miRNA) clusters including miR-17-92, a cluster of miRNAs that has been previously reported to be important for lung sacculation (Lu et al., 2007). miR-17-92 targets and inhibits the transforming growth factor (TGF-β) pathway (Dews et al., 2010; Mestdagh et al., 2010), which is known to regulate cell spreading, adhesion, and tissue morphogenesis (Edlund et al., 2002; Heino et al., 1989; Igniotz et al., 1989; Massague, 2012). Overexpression of this miRNA cluster in the developing lung epithelium leads to decreased TGF-β signaling and inhibition of sacculation, whereas epithelial loss of miR-17-92 rescues much of the phenotype caused by epithelial loss of Hdac3 expression, including AT1 cell spreading and TGF-β signaling. These data reveal a molecular program regulated by Hdac3 that is essential for the spreading of AT1 cells during late lung development, a process critical for sacculation and formation of the large alveolar surface area in the lung required for postnatal gas exchange.

RESULTS

Loss of Hdac3 in the Lung Epithelium Leads to Defects in Sacculation

Lung sacculation and alveologenesis results in the extensive dilation and expansion of the distal lung region after initial specification and differentiation of AT1 and AT2 cells (Figure 1A). Examination of Hdac3 expression during lung development showed that it is broadly expressed in both epithelial and mesenchymal lineages (Figures S1A and S1C). Given the importance of Hdac1 and 2 in early lung development (Wang et al., 2013), we sought to determine whether Hdac3 is also essential for lung development through genetic inactivation of Hdac3 using the Shhcre: Hdac3f/f line that drives cre-dependent recombination in the lung endoderm starting at E8.75 (Goss et al., 2009; Sun et al., 2011). Hdac3 was efficiently deleted in the developing lung epithelium of Shhcre:Hdac3f/f mutants as shown by immunostaining (Figures S1B and S1D). Approximately two-thirds of the newborn Shhcre:Hdac3f/f mutant pups died within postnatal day 2 (P2) and the rest died by P10.

Shhcre:Hdac3f/f mutants developed normally and showed no obvious lung defects up to E17.5, a time point that marks early sacculation (Figures 1B–1E). However, histological analysis of
E18.5 Shhcre:Hdac3f/f mutant lungs revealed significantly compacted distal airspaces and increased thickness of the alveolar wall (Figures 1F–1I). These defects were further illustrated by whole-mount staining of the AT1 cell marker Aqp5 and subsequent 3D reconstruction of the saccular surfaces, which showed a compressed distal lung region with little airspace dilation in the Shhcre:Hdac3f/f mutants (Figures 1J and 1K, and Movies S1 and S2).

Previous studies have shown that one common function of Hdac proteins is to regulate cell proliferation and survival through direct de-repression of cell-cycle inhibitors (Montgomery et al., 2007; Wilson et al., 2006; Yamaguchi et al., 2010). Bromodeoxyuridine labeling or phospho-histone H3 immunostaining did not reveal any detectable changes in cell proliferation at E16.5 and E18.5 (Figures S1E–S1J). Moreover, there were no obvious changes in cell apoptosis as noted by cleaved caspase-3 immunostaining at these time points (Figures S1K–S1N). These data suggest that Hdac3 does not control lung sacculation through modulation of cell proliferation or survival.

_Hdac3 Is Required for the Proper Spreading of AT1 Cells during Lung Sacculation_

AT1 and AT2 cells are the major epithelial cell lineages lining the alveolar sacs and are thought to derive from a common precursor (Rawlins et al., 2009; Treutlein et al., 2014). Their differentiation and maturation from the Sox9+ distal progenitor pool is an important process occurring at sacculation, and decreased AT1 cell differentiation in particular is associated with sacculation and alveolarization defects during lung development (Becker et al., 2011; Kalin et al., 2008; Wongtrakool et al., 2003). To assess whether the alveolar epithelial cell-fate commitment occurs properly in the Shhcre:Hdac3f/f mutants, we examined the expression of several markers for AT2 and AT1 cells. AT2 cells appeared to differentiate normally in Shhcre:Hdac3f/f lungs, as the expression levels of Sftpc, Stfpc, and Abca3 were not significantly altered (Figures 2A–2F and S2A–S2C). The AT1 cell marker Aqp5 was unchanged as shown by immunostaining and qPCR (Figures 2A–2F and S2A). Expression of Pdpn, which is present in both AT1 cells and lymphatic endothelial cells, was modestly reduced by qPCR but was unaltered using immunostaining (Figures S2A, S2D, and S2E). We quantified the percentage of AT2 and AT1 cells in the distal region and showed no significant changes in the numbers of these epithelial lineages in Shhcre:Hdac3f/f mutant lungs (Figures S2H and S2I). In addition, expression of markers for proximal airway epithelial lineages and lung mesenchymal lineages were not significantly altered in the Shhcre:Hdac3f/f mutant lungs (Figures S2A and S2J–S2Q). These data suggest that the lineage commitment of multiple cell types, including that of AT1 and AT2 cells, is largely unaffected by loss of Hdac3. This is in contrast to a previous mouse model in which a mutation of SMRT that abrogated its interaction with nuclear receptors resulted in inhibition of AT1 cell differentiation via regulation of transcription factor Kif2. However, _Kif2_ expression in our Shhcre:Hdac3f/f lungs was not altered (Figure S2R), suggesting that Hdac3 and SMRT may have non-overlapping roles in lung development.

After their appearance at E17.5 as noted by Aqp5 expression, AT1 cells gradually spread and covered 95% of the alveolar surface to form the thin gas-diffusible interface between the airspace and pulmonary vasculature. This morphological change started to occur rapidly after AT1 cell commitment at E17.5, and the flattened morphology of AT1 cells was evident in the E18.5 wild-type lungs (Figures 2A–2C). In contrast, the majority of Shhcre:Hdac3f/f mutant AT1 cells appeared densely packed and failed to acquire a flat squamous cell shape at this stage (Figure 2F). Using ImageJ to quantitate the luminal distance between neighboring AT1 cells, we found that this distance was greatly reduced in the Shhcre:Hdac3f/f mutant lungs (Figures 2G–2H). To further examine whether the disruption of AT1 cell morphology was due to a cell-intrinsic defect in spreading or merely an indirect consequence of unexpanded distal airways, we isolated and cultured E18.5 control and Shhcre:Hdac3f/f mutant primary epithelial cells and examined their ability to spread in vitro without the influence of mesenchymal cells. Immunostaining for Pdpn showed that AT1 cells from Shhcre:Hdac3f/f mutant lungs were unable to spread as extensively as those from their control littermate lungs (Figures 2I–2L). These data indicate that while epithelial Hdac3 is dispensable for alveolar AT1 and AT2 cell lineage commitment, it is essential for promoting AT1 cell spreading during lung sacculation (Figure 2M).

_miR-17-92 and the miRNAs in the Dlk1-Dio3 Locus Are De-repressed upon Loss of Hdac3 Expression in the Lung Epithelium_

To further define the molecular changes that occur upon loss of Hdac3 expression in the lung epithelium, we examined the changes in the transcriptome of Shhcre:Hdac3f/f mutant lungs at E18.5. 1,070 genes were differentially expressed upon loss of Hdac3 expression, with 569 genes upregulated and 501 downregulated (Table S1). Strikingly, expression of 62 miRNA primary transcripts was increased in the Shhcre:Hdac3f/f mutant lungs (Figure 3A). Analysis of these miRNAs revealed that the majority of them belonged to two genetic loci: the large imprinted _Dlk1-Dio3_ locus located on chromosome 12 and the _miR-17-92_ cluster located on chromosome 14. To determine whether these miRNAs were de-repressed in a cell-autonomous manner by loss of Hdac3, we performed mature miRNA qPCR on the isolated EpCAM+ epithelial cells and EpCAM mesenchymal cells from E18.5 lungs (Figures 3B and S3A). These experiments confirmed that the _miR-17-92_ cluster was significantly de-repressed in lung epithelial cells but not in mesenchymal cells after an epithelial cell-specific loss of Hdac3 (Figures 3C and S3B), whereas fewer of the mature miRNAs in the _Dlk1-Dio3_ locus were validated to be de-repressed, and the ones that were exhibited a smaller fold change compared with the _miR-17-92_ cluster (Figure 3C).

To investigate whether Hdac3 can directly repress the expression of these miRNAs, we examined whether Hdac3 could associate with the promoters of the _miR-17-92_ cluster and _Mig_, which hosts most of the _Dlk1-Dio3_ miRNAs, using chromatin immunoprecipitation (ChIP) assays on E18.5 lungs. These data showed that Hdac3 associates with the _miR-17-92_ and _Mig_ promoters but not with an unrelated intergenic region (Figures 3D–3F), suggesting that Hdac3 can directly regulate the expression of these miRNAs.
Previous studies have reported that miR-17-92 is highly expressed at early stages of lung development but declines as development proceeds (Bhaskaran et al., 2009; Lu et al., 2007). Transgenic overexpression of miR-17-92 in the developing lung epithelium using the mouse SftpC promoter (SftpC:MIIR17-92) inhibited epithelial differentiation and resulted in immature sacculation, similar to but more severe than the lung epithelium using the mouse Sftpc promoter (Sftpc:MIIR17-92) with the transcriptome analysis we performed on the Shhcre:Hdac3f/f mutant lungs (Lu et al., 2007). This suggests that Hdac3 may play a role in repressing the expression of miR-17-92 to allow proper lung sacculation to proceed. To determine whether the transcriptome changes caused by loss of Hdad3 were similar to those caused by overexpression of miR-17-92, we compared the differentially upregulated and downregulated genes from the previously reported SftpC:MIIR17-92 microarray (Lu et al., 2007) with the transcriptome analysis we performed on the Shhcre:Hdac3f/f mutant lungs. Using gene set enrichment analysis (GSEA), we found that the upregulated gene set from the SftpC:MIIR17-92 microarray were enriched in the genes downregulated in the Shhcre:Hdac3f/f mutant lungs at E18.5 and how the neighboring AT1 cell-cell distances are measured. (I–L) The E18.5 primary lung epithelial cell culture shows that the Shhcre:Hdac3f/f mutant AT1 cells have a cell-intrinsic defect in spreading as marked by the significantly decreased area of Pdpn+ cells in the culture (I–K, arrows and dotted lines). The spreading ability of AT1 cells is quantified by the mean areas of Pdpn+ cells (L).
Elevated miRNA Expression Represses TGF-β Signaling and AT1 Cell Spreading

Since both the miR-17-92 and Dlk1-Dio3 locus miRNAs were derepressed upon loss of Hdac3 in the developing lung epithelium, we compared the predicted target gene lists of these two miRNA clusters to identify common downstream molecular pathways. A total of 305 common target genes were differentially expressed in the Shh<sup>cre:lox<sup>f/f</sup> mutant lungs, and signaling pathway impact analysis (SPIA) on those genes revealed TGF-β signaling and extracellular matrix (ECM)-receptor interaction as top candidate pathways targeted by these two miRNA clusters (Figure 4A and Table S2). We then performed SPIA on the predicted targets of all the upregulated miRNAs (Table S3), as well as on all the differentially downregulated genes in our microarray data (Table S4). We found that the TGF-β pathway was also among the top candidates in these analyses. Consistent with our results, previous studies have shown that miR-17-92 cluster is a potent inhibitor of TGF-β signaling through targeting key components of the TGF-β pathway as well as TGF-β target genes (Dews et al., 2010; Mestdagh et al., 2010). To verify that these TGF-β signaling components and cell-ECM genes were downregulated in the Shh<sup>cre:lox<sup>f/f</sup> mutant lungs, we performed qPCR analysis, which revealed that most of these genes were significantly downregulated in the lung epithelium but not mesenchyme of Shh<sup>cre:lox<sup>f/f</sup> mutants (Figures 4B, 4C, and S4A). Immunostaining showed that one of the ECM genes, Col4a3, a basement membrane component preferentially expressed in AT1 cells (Treutlein et al., 2014), was significantly downregulated in the Shh<sup>cre:lox<sup>f/f</sup> mutants (Figures S4B and S4C). To determine whether the changes in TGF-β signaling components caused a decrease in TGF-β signaling activity, we performed immunostaining for p-Smad2 at E18.5. p-Smad2 level was significantly decreased in the distal epithelium and mesenchyme including the bronchoalveolar duct junction region, as noted by co-staining for the lung epithelial marker Nkx2.1, consistent with a decrease in TGF-β signaling components including TGF-β ligands (Figures 4D–4I and S4D–S4I). However, the p-Smad2 level in the proximal airway epithelium of Shh<sup>cre:lox<sup>f/f</sup> mutants was relatively unaffected (Figures 4D–4I, dotted outline).
Figure 4. Elevation of miRNAs Leads to Decreased TGF-β Signaling, which in Turn Regulates Lung Sacculation and AT1 Cell Spreading

(A) Comparison of predicted targets of miR-17-92 and miRNAs in Dlk1-Dio3 locus shows a total of 305 common genes targeted by both loci that are also differentially expressed in the Shhcre: Hdac3f/f mutant lungs. SPIA on those genes identified several top candidate pathways including TGF-β signaling and ECM-receptor interaction.

(B and C) qPCR analysis shows that multiple components of TGF-β pathway and genes involved in cell-ECM interaction are downregulated in the epithelial cells of Shhcre: Hdac3f/f mutant lungs.

(D–I) p-Smad2 is downregulated in the distal portion of Shhcre: Hdac3f/f mutant lungs (arrows) while remaining relatively unaffected in the proximal epithelium (white dotted lines). Co-staining with epithelial marker Nkx2.1 shows that p-Smad2 is downregulated in both distal epithelium and mesenchyme. Br, bronchiole.

(J–R) Treatment of TgfβRI inhibitor SB431542 on E16.5 lung explants for 48 hr disrupts lung sacculation (J and K), which mimics the Shhcre: Hdac3f/f mutant explants cultured in parallel (L). TGF-β inhibition also leads to a defect in AT1 cell spreading as shown by Aqp5 staining (M–O, arrows). The defects in saccular formation and AT1 cell spreading are quantified, which shows significant alteration under SB431542 treatment (P–R).

(S–U) Treatment of TgfβRI inhibitor SB431542 reduces AT1 cell spreading in the E18.5 lung epithelial cell culture. Dotted lines in (S) and (T) outline the Pdpn+ AT1 cells.

(legend continued on next page)
TGF-β signaling is widely recognized as a crucial player in cell spreading, adhesion, and tissue morphogenesis (Edlund et al., 2002; Heino et al., 1989; Ignotz et al., 1989; Massague, 2012). Global loss of either TGF-β2 or TGF-β3 led to perinatal lung defects (Kaartinen et al., 1995; Sanford et al., 1997). TGF-β3 null lungs, in particular, were characterized by alveolar hypoplasia and mesenchymal thickening that is reminiscent of Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lungs (Kaartinen et al., 1995). Moreover, lung epithelial-specific loss of TGF-β receptor type I (TgfβRI) (Alk5) also led to collapsed distal airways before birth, although detailed defects in the distal epithelial development were not explored in this study (Xing et al., 2010). To further investigate whether decreased TGF-β signaling could lead to improper remodeling of AT1 cells including reduced cell spreading, we cultured E16.5 wild-type lung explants at the air-liquid interface in the absence or presence of the TgfβRI inhibitor SB431542. The development of wild-type lung explants partially mimicked lung sacculation in vivo, as evidenced by an expansion of distal airways and the specification and early differentiation of AT1 cells, which is noted by Aqp5 expression and their flattened appearance (Figures 4J and 4M). Treatment with SB431542 reduced p-Smad2 level in the lung explants (Figures S4K and S4L), resulting in greatly narrowed distal airway lumen, and disrupted the morphology and remodeling of AT1 cells, including reduced spreading that resembled Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lung explants cultured in parallel (Figures 4J–4R). This was accompanied by the downregulation of several cell-ECM genes which were also decreased in Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lung epithelium (Figure S4A–4J), suggesting that these cell-ECM genes can be regulated by TGF-β signaling. Similarly, treatment of wild-type lung epithelial cell culture with SB431542 also reduced p-Smad2 level and the spreading of AT1 cells in vitro (Figures S4M, S4N, and 4S–4L). Importantly, addition of recombinant TGF-β ligands to the Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant epithelial cell culture increased p-Smad2 expression and partially restored the ability of AT1 cells to spread (Figures S4O, S4P, and 4V–4Y). Together, these data suggest that epithelial-specific loss of Hdac3 in the lung suppresses TGF-β signaling and genes related to cell-ECM interaction, through upregulation of the miR-17-92 and the Dik1-Dio3 miRNA loci. Moreover, these data demonstrate that loss of TGF-β signaling can lead to reduced AT1 cell spreading and defective lung sacculation (Figure 4Z).

**Overexpression of miR-17-92 Can Phenocopy the Loss of Hdac3 and Lead to Defective AT1 Spreading and Lung Sacculation**

A previous study showed that transgenic overexpression of miR-17-92 led to a block in lung epithelial differentiation with an increase in Sox9+ progenitors (Lu et al., 2007). However, our Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lungs did not display a change in Sox9 expression or alveolar epithelial differentiation (Figures SSA–SSD and 2A–2F). Such differences in the overall phenotype observed between these two models could be due to the differing levels of increased miR-17-92 expression, given that the level of miR-17-92 upregulation in Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lungs was more modest than when using the powerful mouse Sftpc promoter (Figure 3C and Lu et al., 2007). Since our data suggest significant molecular similarities between miR-17-92 overexpression and loss of Hdac3 expression during lung epithelial development, we overexpressed miR-17-92 by utilizing the Shh<sup>cre</sup> line and a Rosa26-MIR17-92 knockin allele (R26<sup>miR17-92</sup>) that has been previously described to deliver a moderate level of miR-17-92 expression upon cre recombination (Xiao et al., 2008). We confirmed that the epithelial miR-17-92 expression level in these Shh<sup>cre:</sup>R26<sup>miR17-92</sup> mutants was upregulated by approximately 2- to 4-fold, which was similar to that in Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lungs (Figure S5E). Histological analysis showed that E18.5 Shh<sup>cre:</sup>R26<sup>miR17-92</sup> mutant lungs exhibited a decrease in the distal airway luminal size and an increase in the alveolar wall thickness, similar to but less severe than in the Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lungs (Figures 5A–5H). Aqp5 staining revealed that a large proportion of AT1 cells in Shh<sup>cre:</sup>R26<sup>miR17-92</sup> lungs were densely packed and had not spread properly (Figures 5I–5K). Multiple components of the TGF-β signaling pathway, along with genes related to cell-ECM interaction, were decreased in Shh<sup>cre:</sup>R26<sup>miR17-92</sup> epithelial cells (Figures 5L and 5M), as well as in the MLE12 mouse lung epithelial cells overexpressing miR-17-92 (Figure S5F). These data suggest that elevated levels of this miRNA cluster can directly suppress TGF-β signaling in lung epithelial cells. Similarly to the Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> lungs, we did not observe an increase in Sox9+ or Sox2+ epithelial progenitor cells in the Shh<sup>cre:</sup>R26<sup>miR17-92</sup> lungs at E18.5 (Figures SSA–SSD and S5G–S5L), suggesting that a higher expression level of miR-17-92 may be required to trigger such a block in distal epithelial differentiation. The phenotype observed in Shh<sup>cre:</sup>R26<sup>miR17-92</sup> lungs including the extent of TGF-β signaling downregulation was generally milder than in Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> lungs, suggesting that additional mechanisms, such as the upregulation of Dik1-Dio3 miRNAs, might also play a role.

**Lung Sacculation Requires Hdac3-Mediated Repression of miR-17-92**

To determine whether the sacculation defects in Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> lungs are directly mediated by increased expression of miR-17-92, we generated mice lacking both Hdac3 and miR-17-92 expression (Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup>MIR17-92<sup>−/−</sup>). In contrast to a previous report of a lung hypoplasia phenotype in the miR-17-92 global knockout mice (Ventura et al., 2008), Shh<sup>cre:</sup>MIR17-92<sup>−/−</sup> mutants were viable with no obvious lung phenotype (data not shown). Next, we generated Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup>MIR17-92<sup>−/−</sup> mutants and compared these with Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutants at E18.5. The sacculation defects observed in Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> mutant lungs, including thickened alveolar wall and compacted

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(V–Y) Treatment of Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup> epithelial culture with recombinant TGF-β1/2/3 partially restores the ability of AT1 cells to spread in vitro. Dotted lines in V–X outline the boundaries of AT1 cells in culture.

(2) A diagram shows that TGF-β signaling promotes the spreading and expansion of AT1 cells during lung sacculation.

Ctrl. Control; KO, Shh<sup>cre:</sup>Hdac3<sup>3f/f</sup>. Two-tailed Student’s t test: *p < 0.05; **p < 0.01; NS, not significant; n = 3–5. qPCR and quantification data are represented as mean ± SD. Scale bars: (I), L, T, X) 100 μm; (O) 25 μm.

See also Figure S4.
distal saccular lumen, were significantly alleviated in Shhcre::MIR17-92f/f mutant lungs (Figures 6A–6J and Figure S6). AT1 cells in Shhcre::Hdac3f/f:MIR17-92f/f mutant lungs also exhibited greater spreading with a more flattened appearance compared with those in the Shhcre::Hdac3f/f mutant lungs (Figures 6K–6N). Together, these studies revealed that loss of miR-17-92 significantly restores the AT1 cell remodelling and sacculation defects in Shhcre::Hdac3f/f lungs (Figure 6O).

To test whether loss of miR-17-92 expression would normalize TGF-β signaling as well as the gene expression changes observed in Shhcre::Hdac3f/f mutants, we isolated EpCAM+ cells from Shhcre::Hdac3f/f:MIR17-92f/f mutant lungs and performed qPCR analysis. Most of the gene expression changes were completely or partially normalized in Shhcre::Hdac3f/f:MIR17-92f/f mutant lungs compared with Shhcre::Hdac3f/f mutant lungs (Figures 7A and 7B). Moreover, loss of miR-17-92 expression also increased p-Smad2 level in the distal alveolar region of Shhcre::Hdac3f/f:MIR17-92f/f mutant lungs (Figures 7C–7K). These data suggest that miR-17-92, through targeting the TGF-β pathway, is a critical modulator of Hdac3-mediated regulation of AT1 cell spreading and remodeling of the distal saccules and alveoli required for postnatal lung function (Figure 7L).

**DISCUSSION**

Lung sacculation and alveologenesis remain poorly understood developmental processes that comprise elegant and complicated events of cellular differentiation and remodeling. Defects in these developmental processes can lead to serious lung diseases in the newborn, including bronchopulmonary dysplasia. How alveolar epithelial cells differentiate and remodel to form the intricate gas exchange interface required for postnatal respiration is poorly understood. In particular, how AT1 cells acquire their unique morphology during late lung development that allows them to cover 95% of alveolar luminal surface and facilitate the rapid diffusion of oxygen to the pulmonary vascular plexus is unknown. Our study demonstrates that an epigenetic pathway directed by Hdac3 regulates alveolar epithelial cell remodeling and saccular development in the mouse lung. Hdac3 is essential for AT1 cells to spread and acquire a sheet-like morphology during sacculation. Hdac3 accomplishes this developmental role through repression of the miR-17-92 and Dlk1-Dio3 miRNA clusters, which in turn allows for appropriate levels of TGF-β signaling.

Hdac3 is a class I Hdac that interacts with the co-repressive complexes NCoR and SMRT (Guenther et al., 2000, 2001; Li et al., 2000). A previous study reported that an SMRT knockin allele that abrogated its interaction with nuclear receptors
Figure 6. Loss of Epithelial miR-17-92 Alleviates the Defects of Lung Sacculation and AT1 Cell Spreading in the Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f} Mutant Lungs (A–H) H&E staining shows that the sacculation defects at E18.5 are alleviated in Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f}:MIR17-92\textsuperscript{f/f} lungs compared with Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f} lungs. (I and J) Quantification of alveolar wall thickness and distal airway area indicates that these indices are significantly restored in Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f}:MIR17-92\textsuperscript{f/f} lungs at E18.5. (K–N) Aqp5 immunostaining (red) and quantification of luminal distance (arrows and yellow dotted lines) between two neighboring AT1 cells show that the defect in AT1 cell spreading is significantly improved in Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f}:MIR17-92\textsuperscript{f/f} lungs compared with Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f} lungs at E18.5. (O) Diagram showing that repression of miR-17-92 expression by Hdac3 is required for AT1 cell spreading.

Ctrl, control; KO, Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f}; Rescue, Shh\textsuperscript{cre}:Hdac3\textsuperscript{f/f}:MIR17-92\textsuperscript{f/f}. Two-tailed Student’s t test: *p < 0.05; **p < 0.01; NS, not significant; n = 3–4. Quantification data are represented as mean ± SD. Scale bars: (H) 100 μm; (M) 50 μm.

See also Figure S6.
and inhibit multiple components of TGF-β signaling. miR-17-92 can directly target miRNAs in the lung epithelium, and that elevated levels of miR-17-92 disrupt formation of distal saccules and primitive alveoli by inhibiting AT1 cell spreading. MiR-17-92 also plays a regulatory role in repressing this cluster of miRNAs also plays an important role in repressing TGF-β signaling specifically in the distal region of E18.5 ShhCre: Hdac3f/f:MIR17-92f/f lungs compared with ShhCre: Hdac3f/f lungs. (C–K) p-Smad2 staining level is improved in the distal region of E18.5 ShhCre: Hdac3f/f:MIR17-92f/f lungs compared with ShhCre: Hdac3f/f lungs. Nkx2.1 co-staining shows that p-Smad2 level is improved in both distal epithelial and mesenchymal (C–K, arrows). Dotted lines mark the proximal airways. Br, bronchiale.

(L) A model diagram summarizes the molecular and cellular mechanisms by which epithelial Hdac3 regulates mouse lung sacculation. Hdac3 directs lung sacculation and AT1 cell spreading through suppressing miRNAs including miR-17-92, which in turn allow for proper level of TGF-β signaling and expression of cell-ECM genes that are crucial for cell spreading and tissue morphogenesis. Ctrl, control; KO, ShhCre: Hdac3f/f; Rescue, ShhCre: Hdac3f/f:MIR17-92f/f. Two-tailed Student’s t test: *p < 0.05; **p < 0.01; NS, not significant; n = 3–4, qPCR data are represented as mean ± SD. Scale bar in (K), 100 µm.

Figure 7. Loss of Epithelial miR-17-92 Significantly Restores TGF-β Pathway in the ShhCre: Hdac3f/f Mutant Lungs

(A and B) qPCR analysis shows that multiple components of TGF-β pathway and genes involved in cell-ECM interaction are significantly restored in the E18.5 ShhCre: Hdac3f/f:MIR17-92f/f lungs compared with ShhCre: Hdac3f/f lungs.

produced defects in lung sacculation and inhibited AT1 cell differentiation (Pei et al., 2011). This phenotype was partially attributed to the downstream target Klf2 and its role in promoting the AT1 cell differentiation program. In contrast to these results, loss of Hdac3 does not significantly perturb the specification of the AT1 cell lineage, and Klf2 expression levels appeared to be unchanged. One possibility for this discrepancy could be that our model examined the epithelial cell-specific function of Hdac3 while the SMRT knockin mutation is a globally expressed point mutation. Moreover, the SMRT knockin mutation used in these studies only abrogated its interaction with nuclear receptors but retained its ability to bind Hdac3. Recent studies have also implicated that SMRT has Hdac3-independent functions in some contexts (Adikesavan et al., 2014), and that SMRT and NCoR have non-redundant in vivo functions (Mottis et al., 2013; Sun et al., 2013), suggesting that despite the co-existence of Hdac3 and SMRT in the same complex, these two proteins are not completely interchangeable in their molecular functions.

Previous studies have shown that the miR-17-92 cluster is highly expressed in the early stages of lung development and declines as development proceeds (Bhaskaran et al., 2009; Lu et al., 2007). The downregulation of this cluster is important for late lung development, as transgenic epithelial-specific overexpression of miR-17-92 impeded differentiation and saccular development (Lu et al., 2007). Our study demonstrates that Hdac3 plays a regulatory role in repressing this cluster of miRNAs in the lung epithelium, and that elevated levels of miR-17-92 disrupt formation of distal saccules and primitive alveoli by inhibiting AT1 cell spreading. MiR-17-92 can directly target and inhibit multiple components of TGF-β signaling including TGF-β responsive genes (Dews et al., 2010; Mestdagh et al., 2010). Consistent with this observation, we found a downregulation of genes involved in TGF-β signaling specifically in the Hdac3-deficient lung epithelium that can be partially restored by epithelial-specific loss of miR-17-92. TGF-β signaling is known as a critical regulator of cell spreading, adhesion, and migration. Previous studies have suggested that it plays a role in lung sacculation and alveologenesis, but the cellular and molecular processes it regulates during this stage of development have remained unclear (Kaartinen et al., 1995; Sanford et al., 1997; Xing et al., 2010). We found that TGF-β signaling promotes the spreading of AT1 cells during lung sacculation, supporting a model in which the morphogenetic changes of AT1 cells are directed by an Hdac3-miRNA-TGF-β molecular cascade. This process of AT1 cell spreading is likely to be important for the rapid expansion of distal saccular surface area that occurs during late lung development in preparation for alveologenesis. We also observed increased expression of miRNAs in the maternally imprinted Dlk1-Dio3 locus, which is regulated by histone acetylation in induced pluripotent stem cells (Stadtfeld et al., 2010). Several miRNAs in this locus, including miR376c, are reported or predicted to directly target components of the TGF-β pathway (Fu et al., 2013; Liu et al., 2014), suggesting that this cluster of miRNAs also plays an important role in repressing TGF-β signaling.

The generation of the thin gas-diffusible interface between the external environment and the vasculature remains a unique characteristic of the mammalian respiratory system. Our data suggest that Hdac3 plays a critical role in the initial establishment of alveolar structure through the promotion of AT1 cell spreading.
by inhibiting miR-17-92-mediated repression of TGF-β signaling. Such information will be beneficial to our understanding of how neonatal respiratory diseases occur and how an alveolus reestablishes its structure after postnatal injury.

EXPERIMENTAL PROCEDURES

Animals
The generation and genotyping of Hdac3<sup>−/−</sup>, Shh<sup>cre</sup>, R26<sup>R26</sup>MIR17-92, and MIR17-92<sup>lox/lox</sup> alleles have been previously described (Goss et al., 2009; Sun et al., 2011; Ventura et al., 2008; Xiao et al., 2008). All animal work was performed in accordance to the guidance of the University of Pennsylvania Institutional Animal Care and Use Committee.

qPCR
For mRNA qPCR, total RNA was isolated from lungs or cells at indicated time points by an RNeasy Mini Kit (Qiagen) or RNeasy Micro Kit (Qiagen). First-strand cDNA synthesis was performed using SuperScript III Reverse Transcriptase (Life Technologies). qPCR was performed using the SYBR green system (Applied Biosystems), GAPDH expression level was used as an internal control. See Supplemental Experimental Procedures for qPCR primer sequences. For mature mRNA qPCR, a MirVana miRNA isolation kit (Life Technologies) was used to isolate total RNA that contains small RNAs. The TaqMan MicroRNA Assays (Life Technologies) was used to quantitate various mRNA levels. Sn202 was used as an internal control for miRNA quantity.

Histology
Tissues were fixed in 4% paraformaldehyde overnight and subject to paraffin sectioning. Immunohistochemistry was performed by using the following antibodies and concentrations: Hdac3 (Santa Cruz Biotechnology, 1:10), Nkx2.1 (Santa Cruz, 1:50), p-Smad2 (Millipore, 1:1000), Sox2 (Seven Hills, 1:500), Sox9 (Santa Cruz, 1:100), SftpC (Santa Cruz, 1:50), SftpB (Chemicon, 1:100), Aqp5 (Abbcom, 1:100), Pdn1 (Hybridoma Bank, 1:50), Sgcbl1A1 (Santa Cruz, 1:20), TubbIV (BioGenex, 1:20), Pdgfrα (Cell Signaling, 1:25), Pecam (Pharmin-gen, 1:500), Coll4A3 (Santa Cruz, 1:50), RAGE (R&D Systems, 1:50) c-Caspase 3 (Cell Signaling, 1:50), BrdU (Abbcom, 1:100), and phospho-histone H3 (Cell Signaling, 1:200). Whole-mount staining was performed as previously described (Metzger et al., 2008).

Quantification of Alveolar Wall Thickness and Airway Lumen Area
For each sample, three pictures were taken under a 40× objective lens. The method for quantification of alveolar wall thickness was previously described (Herriges et al., 2014). The areas for distal airways was measured by using the area measurement function in ImageJ and calculated for mean value and SD.

Transcriptome and Bioinformatics Analysis
RNA was isolated from E18.5 lungs from Shh<sup>cre</sup> control and Shh<sup>cre</sup>:Hdac3<sup>−/−</sup> mutant embryos. Five lungs were collected for either control or Shh<sup>cre</sup>:Hdac3<sup>−/−</sup> groups. The RNA was then used to generate a biotinylated cRNA probe library for Affymetrix Mouse Gene 2.0ST array. Microarray data were analyzed using the Oligo package available at the Bioconductor website (www.bioconductor.org). The raw data were background-corrected by the Robust Multichip Average method and then normalized by an invariant set method. Differential gene expression between the control and mutant mice was analyzed by the Limma package available at the Bioconductor website. p Values obtained from the multiple comparison tests were corrected by false-discovery rates. Heatmap displays were created using the heatmap.2 function in the R package gplots. See Table S1 for the list of genes that were differentially expressed. The Gene Expression Omnibus accession number for the microarray data is GEO: GSE70864.

Functional enrichment analysis of pathways and gene ontology terms was performed using the GSEA software package (PMID: 16199517). Predicted miRNA-gene targets were obtained from the microRNA.org website, and only target predictions that were conserved and annotated with a “good” mirSVR score were considered for analysis. Pathway analysis of genes targeted by miR-17-92 cluster and Dlk1 Dio3 locus was performed using the SPIA R software package (PMID: 18990722) using the KEGG pathway annotations.

ChIP Assays
E18.5 WT lungs were dissected and cross-linked by 3.7% formaldehyde and sonicated to obtain genomic DNA sizes between 200 and 400 bp. ChIP assays were performed using the ChIP Assay Kit (Millipore) and Hdac3 antibody (Abcam). Rabbit immunoglobulin G was used as a negative control in these assays. See Supplemental Experimental Procedures for ChIP primer sequences.

Lung Explant Culture
Lung explant culture was performed as previously reported (Geng et al., 2011). In brief, E16.5 lungs were dissected and diced into 0.5- to 1-mm thick pieces and cultured on an air-liquid interface with DMEM media for 48 hr. The explants were treated with 10 μM SB431542 or vehicle.

Lung Epithelial Cell Isolation and Culture
E18.5 lungs were dissected and digested by collagenase and Dispase to obtain single-cell solution. A Dynabeads Flow Comp Flexi Kit (Life Technologies) and EpCAM antibody (eBioscience) were used to isolate primary lung epithelial cells. The epithelial cells were cultured on the surface coated by fibronectin for 8 days to achieve a well-spread epithelial sheet (Demaiso et al., 2009). The epithelial cells were treated with 10 μM SB431542 from the first day of culture. For TGF-β ligand treatment, Tgfβ1, Tgfβ2, and Tgfβ3 (R&D) were added to the epithelial culture at 5 ng/ml each. Before immunostaining, cells were fixed in 4% paraformaldehyde for 10 min and permeabilized with 0.1% Tween 20.

Lentivirus
Mouse MIR17-92 overexpression construct (Lu et al., 2007) was cloned into pLenti 7.3/V5-DEST through the Gateway cloning system. Lentivirus containing empty vector or MIR17-92 overexpression construct was packaged and produced as previously reported (Herriges et al., 2014). MLE12 Cells were infected with lentivirus and allowed to grow for 48 hr before RNA was extracted for qPCR.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, four tables, and two movies and can be found with this article online at http://dx.doi.org/10.1016/j.devcel.2015.12.031.

AUTHOR CONTRIBUTIONS

Y.W., D.B.F., M.A.L., and E.E.M. designed experiments and wrote the manuscript. Y.W., D.B.F., S.Z., X.W., and M.M.L. performed experiments and analyzed data. M.P.M. analyzed bioinformatics data.

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