The cell of origin and subtype of K-Ras-induced lung tumors are modified by Notch and Sox2

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Cell type-specific conditional activation of oncogenic K-Ras is a powerful tool for investigating the cell of origin of adenocarcinomas in the mouse lung. Our previous studies showed that K-Ras activation with a CC10(Scgb1a1)-CreER driver leads to adenocarcinoma in a subset of alveolar type II cells and hyperplasia in the bronchioalveolar duct region. However, no tumors develop in the bronchioles, although recombination occurs throughout this region. To explore underlying mechanisms, we simultaneously modulated either Notch signaling or Sox2 levels in the CC10+ cells along with activation of K-Ras. Inhibition of Notch strongly inhibits adenocarcinoma formation but promotes squamous hyperplasia in the alveoli. In contrast, activation of Notch leads to widespread Sox2+, Sox9+, and CC10+ papillary adenocarcinomas throughout the bronchioles. Chromatin immunoprecipitation demonstrates Sox2 binding to NOTCH1 and NOTCH2 regulatory regions. In transgenic mouse models, overexpression of Sox2 leads to a significant reduction of Notch1 and Notch2 transcripts, while a 50% reduction in Sox2 leads to widespread papillary adenocarcinoma in the bronchioles. Taken together, our data demonstrate that the cell of origin of K-Ras-induced tumors in the lung depends on levels of Sox2 expression affecting Notch signaling. In addition, the subtype of tumors arising from type II cells is determined in part by Notch activation or suppression.

Keywords: Notch; Sox2; K-Ras; lung cancer

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Lung adenocarcinoma is a heterogeneous disease, both histologically and genetically. While much progress has been made in therapeutically targeting specific driver mutations [EGFR, ALK, and ROS] (Lynch et al. 2004; Takeuchi et al. 2012), K-RAS mutant tumors remain difficult to treat effectively. Farnesyl transferase inhibitors have failed (Johnson and Heymach 2004), and MEK inhibitors have proven ineffective (Adjei et al. 2008), partly because they block only one pathway downstream from the oncogenic kinase. Synthetic-lethal approaches identify genes that, when inhibited, are lethal to K-RAS-activated but not wild-type cells (Luo et al. 2009; Scholl et al. 2009). However, these have not resulted in effective therapies (Luo et al. 2009). One reason for these failures may be the inadequate consideration of the cellular context of K-RAS activation. We previously demonstrated that in the mouse lung, K-RasG12D induces adenocarcinoma formation preferentially in a subset of type II cells in the alveoli, even when the oncogenic mutation is induced throughout the bronchiolar airways (Xu et al. 2012). Others have subsequently corroborated this cell of origin specificity but have not identified underlying mechanisms (Mainardi et al. 2014; Sutherland et al. 2014). Here, we address this question by identifying molecular differences between bronchiolar cells that are not transformed and alveolar cells that are tumorigenic when K-RasG12D is induced. Specifically, we focus on the role of Notch signaling in controlling this differential response.

The Notch signaling pathway plays an important role in lung development and cell fate decisions (Tsao et al. 2009; Rock et al. 2011; Guha et al. 2012; Morimoto et al. 2012). Mounting data implicate alteration of the Notch
signaling pathway in lung cancer development in both humans and mice [Westhoff et al. 2009; Eliasz et al. 2010; Allen et al. 2011; Osanyingbemi-Obidi et al. 2011; Wang et al. 2011; Yang et al. 2011; Maraver et al. 2012; Licciulli et al. 2013]. Notch receptors transduce signals in response to ligands on neighboring cells, regulating lineage selection and developmental patterning. To activate Notch transcriptional targets, ligands [Jagged1, Jagged2, and Delta1–3] bind to receptors [Notch1–4], leading to two proteolytic cleavages at the membrane. The second of these, mediated by γ-secretase, frees the Notch intracellular domain (ICD), which translocates to the nucleus and forms a DNA-binding complex with coactivators, including Mastermind-like (MAML). This transcriptional complex activates downstream effectors, including Hes1, Hes5, Hey1, Hey2, and Hey5. Recent studies [Wang et al. 2011; Cancer Genome Atlas Research Network 2012] have demonstrated that inhibitory alterations of Notch signaling are frequent in human squamous non-small cell lung cancer (NSCLC).

In this study, we show that inhibition of Notch signaling strongly abrogates K-Ras-induced adenocarcinoma formation, while activation of Notch in bronchiolar cells allows K-Ras-induced tumorigenesis. Further investigation demonstrates that NOTCH is a direct target of the SOX2 transcription factor in human cells and that Sox2 overexpression leads to transcriptional repression of Notch1 and Notch2 in mice. Genetically reducing the level of Sox2 by 50% in CC10-CreER; Lox-Stop-Lox (LSL)-K-RasG12D mice allows bronchiolar tumor formation. Together, these data identify Notch signaling as a key determinant of both tumor initiation and tumor progression in K-Ras-activated respiratory epithelial cells.

Results

Notch signaling is required for K-Ras-induced lung adenocarcinoma

Involvement of the Notch pathway in K-Ras-induced tumors has been controversial. Using mouse models of K-RasG12D activated by inhaled adenoviral Cre recombinase, groups have published conflicting data regarding its necessity for in vivo lung tumorigenesis [Osanyingbemi-Obidi et al. 2011; Maraver et al. 2012]. In order to determine whether Notch activation is necessary for K-Ras-induced tumors in the subset of alveolar type II cells that expresses CC10+, we exploited a conditional Rosa26\textsuperscript{DNMaml1-GFP} allele encoding a dominant-negative Mastermind-like protein [DNMaml1] tagged with GFP [Tu et al. 2005]. Recombination of this allele results in the expression of a competitive inhibitor of the Rpbj coactivator Mamll that is required for transcriptional activation of Notch targets [Nam et al. 2006]. We generated CC10-CreER; LSL-K-RasG12D; Rosa26-DNMaml1-GFP [hereafter, CC10-CreER; K-Ras; DNMaml1] mice and used CC10-CreER; LSL-K-RasG12D; Rosa26-f-GFP [CC10-CreER; K-Ras] mice as controls. At 15 wk after four doses of tamoxifen, the lungs of control CC10-CreER; K-Ras mice contain widespread adenocarcinomas and adenomas [Fig. 1A, left panel]. In contrast, CC10-CreER; K-Ras; DNMaml1 lungs [Fig. 1A, right panel] contain mostly small clusters of hyperplastic GFP+ columnar cells in the alveoli and bronchioalveolar duct junction (BADJ), very few adenomas >100 μm in diameter, and no adenocarcinomas (Fig. 1A). Quantification reveals 12.6 ± 2.6 lesions >100 μm in diameter per left lobe in CC10-CreER; K-Ras mice and only 2.8 ± 0.8 lesions of this size in the CC10-CreER; K-Ras; DNMaml1 mice (P < 0.05). The adenomas and adenocarcinomas in CC10-CreER; K-Ras mice are GFP+ Sftp+ [Fig. 1B]. In contrast, the cells in alveolar hyperplasia/ small adenomas in CC10-CreER; K-Ras; DNMaml1 mice are GFP+ Sftp– (Fig. 1C). Significantly, the very rare larger adenomas (>100 μm) that do develop in experimental lungs are GFP-negative [Fig. 1C, C'], demonstrating activation of the K-Ras allele only and escape from recombination of the DNMaml1 allele. Immunohistochemistry for CC10 shows that the tumorigenic cells in CC10-CreER; K-Ras mice lose CC10 expression when K-Ras is activated [Fig. 1D]. This is consistent with our previous report [Xu et al. 2012]. However, although the majority of the GFP+ cells in CC10-CreER; K-Ras; DNMaml1 mice are CC10-negative [Fig. 1E, E'], some cells in the hyperplasia/small adenomas continue to express CC10. Thus, in our model, active Notch signaling is required for K-RasG12D-induced adenocarcinoma formation in the subset of alveolar type II cells that express CC10.

Inhibition of Notch in K-RasG12D; CC10-expressing alveolar cells leads to a Sox2+Krt5+Trp63+ squamous phenotype

Recently, it was demonstrated that γ-secretase inhibition abrogates K-Ras-induced tumor formation in the mouse lung [Maraver et al. 2012]. However, while Notch ablation caused inhibition of large tumors in our CC10-CreER; K-Ras; DNMaml1 mice, clusters of hyperplastic cells and small adenomas persisted. To further characterize these lesions, we performed immunostaining for lung epithelial markers. As shown in Figure 2, A and A', consistent with our previous report [Xu et al. 2012], the majority of lung adenomas and lung adenocarcinomas in the alveolar region of CC10-CreER; K-Ras mice do not express the transcription factor Sox2. However, expression of DNMaml1 leads to clear expression of Sox2 in all GFP-positive cells [Fig. 2B, B']. Moreover, Foxi1, a marker for ciliated airway epithelial cells, is not seen in lung adenomas and adenocarcinomas in CC10-CreER; K-Ras mice [Fig. 2C, C'] but is expressed in some Sox2+ cells in the hyperplastic lesions in CC10-CreER; K-Ras; DNMaml1 mice [Fig. 2D, D']. To further test whether hyperplastic lesions express other markers of cell types normally present in the Sox2+ proximal airways of the mouse lung, we stained for Trp63 and Krt5. These proteins are normally expressed only in basal cells of the pseudostratified epithelium of the mouse trachea [Fig. 3A, A'] and proximal, extralobar bronchi. They are also markers for squamous carcinoma in the human lung. As shown in Figure 3, B and B', some cells in the hyperplastic...
lesions in the alveoli of CC10-CreER; K-Ras; DNMaml1 mice express Trp63 alone (17.3%), Krt5 alone (5.4%), or both Trp63 and Krt5 (6.1%) (Fig. 3C). These expression patterns are never seen in the alveoli of control lungs (data not shown). These results indicate that Notch inhibition with resultant Sox2 up-regulation drives a squamous phenotype in cells that were originally K-RasG12D-expressing type II cells. These Sox2-high/Trp63+/Krt5+ cells do not progress to larger tumors at the times of sacrifice in this study.

Figure 1. Genetic inhibition of Notch signaling significantly reduces K-Ras-induced lung adenocarcinoma formation. (A) Sections of left lobes of CC10-CreER; LSL-K-RasG12D; Rosa26-f-GFP (CC10-CreER; K-Ras) ([left]) and CC10-CreER; LSL-K-RasG12D; Rosa26-LSL-DNMaml1-GFP (CC10-CreER; K-Ras; DNMaml1) ([right]) mice 15 wk after tamoxifen injection, stained with H&E. Note that the lung expressing both activated K-Ras and DNMaml1 has significantly fewer large tumors (>100 μm), which is quantified in the bottom panel. (B) Section of CC10-CreER; K-Ras mouse lung stained with DAPI, GFP (green), and Sftpc (purple). The inset shows higher magnification of tumor cells (arrow). Note that tumors are GFP’Sftpc-. (C) Section of CC10-CreER; K-Ras; DNMaml1 mouse lung stained with DAPI, GFP (green), and Sftpc (purple). Note that widespread GFP+ hyperplastic clusters are Sftpc- . Rare larger solid adenomas (arrow) are GFP+ Sftpc- (white dashed circles with asterisk), indicating that these rare large adenomas escape DNMaml1 expression. (C9) Higher magnification of the boxed area in C. (D) Section of CC10-CreER; K-Ras mouse lung stained with DAPI, GFP (green), and CC10 (red) showing that tumors are CC10-. (E) Section of CC10-CreER; K-Ras; DNMaml1 mouse lung stained with GFP and CC10. (E9) Higher magnification of the white boxed area in E demonstrating that hyperplastic clusters/small adenomas contain a majority of cells that are GFP+ alone.

Figure 2. Inhibition of Notch signaling leads to high Sox2 and Foxj1 expression in CC10-expressing alveolar cells. (A,B), Mouse lung sections stained with DAPI, GFP (green), and Sox2 [red]. (A) Sections of CC10-CreER; K-Ras mouse lung. Sox2 expression is seen in airway epithelial cells but not tumor [white dashed line and asterisk]. (A9) Higher magnification of boxed region in A. (B) Sections of CC10-CreER; K-Ras; DNMaml1 mouse lung. Hyperplasia/small adenoma with high Sox2 expression in all GFP+ cells. (B9) Higher magnification of the boxed region in B. (C,D) GFP [green], Foxj1 [red], and Sftpc [purple] staining for the indicated mouse lung sections 15 wk after tamoxifen treatment. (C) Tumor cells in CC10-CreER; K-Ras mouse lung are GFP+Sftpc−Foxj1+. (C9) Higher magnification of the boxed region in C. (D) GFP−Foxj1+ cells in some alveolar hyperplastic lesions in CC10-CreER; K-Ras; DNMaml1 mouse lung. Note that all GFP+ cells are Sftpc−. (D9) Higher magnification of the boxed region in D.
Overexpression of Notch1 ICD promotes papillary adenocarcinoma tumor formation in CC10+ airway cells that express K-RasG12D

Because Notch inhibition suppresses K-Ras-induced tumors in the alveoli, we reasoned that Notch activation might allow or promote K-Ras-induced tumor formation in CC10+ airway cells. In order to determine whether bronchiolar cells form tumors when Notch is activated in conjunction with K-Ras, we made use of the Rosa26-NotchICD-GFP allele (Murtaugh et al. 2003). Following Cre-mediated recombination, this allele results in the constitutive expression of the ICD of Notch1 that is sufficient for ligand-independent, cell-autonomous Notch activation as well as nuclear GFP. We treated adult CC10-CreER; LSL-K-RasG12D; Rosa26-NotchICD-GFP (CC10-CreER; K-Ras; NotchICD) mice with tamoxifen. CC10-CreER; LSL-KrasG12D; Rosa26-f-GFP (CC10-CreER; K-Ras) mice of the same age were controls. After 15 wk, we assessed the lung tumor burden of both groups. Strikingly, we observed the widespread appearance of papillary lung adenocarcinomas in airways of CC10-CreER; K-Ras; NotchICD mice (Fig. 4A, right panel). In contrast, such tumors are very rare in CC10-CreER; K-Ras control mice (Fig. 4A, left panel). We quantified tumors per airway and found that CC10-CreER; K-Ras; NotchICD mice had significantly higher numbers of airway adenocarcinomas than CC10-CreER; K-Ras mice (Fig. 4E; Supplemental Table S1). Thus, taken together, our loss- and gain-of-function Notch allele studies demonstrate that Notch signaling activity determines tumor location/cell of origin when K-RasG12D is induced.

Papillary adenocarcinomas in CC10+; K-RasG12D airway cells overexpressing Notch express CC10 and both Sox2 and Sox9

To further characterize the airways of CC10-CreER, K-Ras mice with and without NotchICD, we immunostained lung sections with antibodies to Sftpc, Foxj1, CC10, Krt5, and Trp63. The columnar epithelial cells lining the airways in control CC10-CreER; K-Ras mice have an increased number of GFP+Foxj1+ cells 15 wk after tamoxifen injection compared with CC10-CreER; Rosa26-f-GFP mice (Supplemental Fig. S1). The GFP+Foxj1+ cells in CC10-CreER; K-Ras mice express acetylated tubulin and do not incorporate BrdU (data not shown). This phenotype mimics the effect of inhibiting Notch during lung development, which leads to terminal differentiation of CC10+ cells into Foxj1+ ciliated cells when a proliferative signal is encountered (Rock et al. 2011; Guha et al. 2012). In contrast, consistent with high Notch promoting secretory airway phenotypes, most of the GFP+ tumor cells in the papillary adenocarcinomas of the CC10-CreER; K-Ras; NotchICD mice are CC10+ and Foxj1−/C0. They are also negative for Sftpc (Fig. 4E,F,G,G′), Krt5, and Trp63 (data not shown).

To further analyze the phenotype of the papillary lung adenocarcinomas in CC10-CreER; K-Ras; NotchICD mice, we stained sections for Sox2 and Sox9. Both transcription factors are expressed in the embryonic lung, but Sox9 is normally down-regulated at birth [Chang et al. 2013]. In contrast to the alveolar adenocarcinomas in CC10-CreER; K-Ras mice, which are Sox2− and Sox9+ (data not shown), the papillary adenocarcinomas in CC10-CreER; K-Ras; NotchICD mice express both Sox2 and Sox9, as judged by staining of adjacent sections [Fig. 4H,H′,I′]. These data demonstrate that Notch activation leads to expression of Sox9, a distal embryonic lung
marker that is often expressed in human lung tumors (Jiang et al. 2010; Zhou et al. 2012).

Reduction of Sox2 expression in CC10; K-RasG12D airway cells leads to CC10+ papillary adenocarcinoma

Notch is activated only in tumor-forming type II cells when K-Ras is induced [Maraver et al. 2012]. We next asked what regulates the Notch pathway in these cells. Sox2, a high-mobility box transcription factor, is normally expressed in epithelial cells of the proximal trachea, bronchi, and bronchioles but not in the distal alveolar region of the lung [Tompkins et al. 2009]. Because of this expression pattern and because it binds a Notch1 intergenic region in the retina [Taranova et al. 2006], we reasoned that Sox2 limits Notch transcription
in airway cells. We previously generated a Rosa26-LSL-CAG-Sox2-IRES-GFP [LSL-Sox2] mouse line (Lu et al. 2010). When CC10-CreER; LSL-Sox2 mice are administered tamoxifen, adenocarcinoma with squamous features develops in the alveoli in about half of the mice. In order to identify potential mechanistic effectors of Sox2, we isolated RNA from GFP+ cells present in CC10-CreER; LSL-Sox2 (n = 7) and CC10-CreER; Rosa26-f-GFP (n = 4) lungs 16 wk after tamoxifen treatment. Microarray analysis was performed using Affymetrix 430.2 chips. We found that Hes1, Notch1, and Notch2 are significantly down-regulated when Sox2 is overexpressed (14.4-fold, 18.9-fold, and 15.3-fold, respectively) (Supplemental Table S2; Supplemental Fig. S2A). In addition, we performed chromatin immunoprecipitation (ChIP) using a SOX2 antibody in SOX2-overexpressing H520 human lung cancer cells. The resulting DNA was sequenced, and significant peaks were noted at the promoter region of HES1 and NOTCH1 and in an intergenic region of NOTCH2 (Supplemental Fig. S2B, Supplemental Table S2). Taken together, these analyses suggest that Sox2 directly binds regulatory sequences of Hes1, Notch1, and Notch2, leading to suppression of transcription.

Based on these findings, we reasoned that reducing the level of Sox2 in bronchial/bronchiolar epithelial cells would release the inhibition of Notch signaling when K-RasG12D is activated. Because Sox2 is required for CC10 cell survival (Tompkins et al. 2009), and CC10-CreER; LSL-K-RasG12D, Sox2flox/fox mice die early due to abnormal airways (Supplemental Fig. S3), we generated CC10-CreER; LSL-K-RasG12D, Sox2flox/+ mice, which have 50% of the wild-type Sox2 level (Supplemental Fig. S4). At 12 wk after tamoxifen, CC10-CreER, K-Ras; Sox2flox/fox mice already developed airway papillary adenocarcinomas (Fig. 5A–D). To characterize these lesions, we immunostained sections for Foxj1, CC10, and Sox2. As described earlier, control CC10-CreER, K-Ras mice have a single layer of GFP+ epithelial cells lining the airways 15 wk after tamoxifen injection. In contrast, airways in CC10-CreER, K-Ras; Sox2flox/fox mice contain papillary adenocarcinomas. Most tumor cells that extend into the airway lumen in these mice are Foxj1+, and a few GFP+Foxj1+ cells remain in the normal airway epithelial layer [Fig. 5E, E′, F, F′]. This result suggests that a reduction in Sox2 levels can allow tumor-resistant CC10+ airway cells to initiate tumors when K-Ras is induced.

To ask whether Sox2 overexpression suppresses tumor formation in K-RasG12D-expressing type II cells, we created a CC10-CreER; LSL-K-RasG12D, Sox2flox/fox mouse line and administered tamoxifen at 6 wk after birth. As Figure 6 reveals, these mice have markedly smaller alveolar tumors than CC10-CreER, K-Ras mice at similar ages. Immunohistochemical analysis of these alveolar hyperplastic regions demonstrates positive staining for the squamous markers Krt5 and Trp63. All hyperplastic cells were noted to be strongly GFP+ [Sox2 label]. These data demonstrate that overexpression of Sox2 is tumor-suppressive for large adenocarcinomas in this model.

Figure 5. Reduction of Sox2 in CC10+ airway cells leads to papillary adenocarcinoma formation. (A, right) Sections of CC10-CreER, LSL-K-RasG12D, Sox2flox/+ mouse lungs 12 wk after tamoxifen injection stained with H&E. (B,C) Higher magnification of papillary airway tumors. (D) Bar graph demonstrating percentage of airways with papillary tumors in CC10-CreER, LSL-K-RasG12D mice. (E,F, right) Sections of CC10-CreER, LSL-K-RasG12D, Sox2flox/+ mouse lungs stained with the indicated markers.

Discussions

Collectively, our data provide further insights into the pivotal roles of Sox2 and Notch signaling in K-Ras-driven lung tumors. As the model in Figure 7A summarizes, in the alveoli, CC10+Sftpc+ cells in which K-Ras has been activated progress to lung adenocarcinoma, and this progression requires the ability to activate Notch signaling. When we block Notch signaling, adenocarcinoma formation is inhibited, and only hyperplastic clusters/small
adenomas are found. These cells express proximal lung epithelial markers and do not progress to adenocarcinomas over the course of this study. In the airways (Fig. 7B), K-Ras-activated CC10+ cells differentiate into Foxj1+ ciliated cells and are resistant to tumor formation. However, when Notch is activated or Sox2 is down-regulated, the K-Ras-activated CC10+ cells can form CC10+Foxj1+ papillary adenocarcinomas.

There are at least two subtypes of type II cells in the alveoli [a majority that are CC10+Sftpc+ and a minority that are CC10'Sftpc']. Our current results from CC10-CreER-driven models indicate that inhibition of Notch signaling or overexpression of Sox2 can inhibit K-Ras-activated lung adenocarcinoma formation originating from CC10'Sftpc- cells. Preliminarily, Sftpc-CreER; LSL-K-RasG12D; LSL-Sox2 mice also have markedly abrogated tumor formation [data not shown]. Further analysis of these mice will be the focus of another study. Our data are consistent with a previous report (Sutherland et al. 2014) that lung adenocarcinoma formation requires reduced Sox2 expression.

A previous study suggested that genetic or γ-secretase inhibition of the Notch pathway blocks tumor growth in codon 12 mutant Kras mice (Maraver et al. 2012). However, in humans, missense and nonsense mutations in Notch ligands and receptors lead to squamous cell tumors in the lung [Wang et al. 2011; Cancer Genome Atlas Research Network 2012]. This is consistent with the development of alveolar hyperplasia with squamous markers that we observed with Notch inhibition in the current study. Because of this, we believe that γ-secretase inhibitors should be used with great caution in patients with lung adenocarcinoma. Because γ-secretase inhibition blocks all Notch receptors, a future direction will be to target individual Notch receptors and Notch targets as potential therapeutic strategies.

CC10+ airway papillary adenocarcinomas in CC10-CreER; K-Ras; Sox2flox/+ and CC10-CreER; K-Ras; NotchICD mice, unlike alveolar tumors, do not express Sftpc, a marker for type II cells that is expressed in the majority of lung adenocarcinomas. In humans, K-RAS mutant tumors are strictly alveolar (Cooper et al. 1997), although our data would predict that K-RAS mutant tumors may arise from bronchiolar cells if NOTCH signaling is also activated (or if SOX2 is inactivated). This will be a focus of further study with human lung adenocarcinoma samples.

Terminal differentiation of CC10+ airway cells into Foxj1+ ciliated cells when K-RasG12D is induced is similar to the effect of inhibition of Notch in lung development. As the airway cells proliferate during airway development, low levels of Notch signaling result in the differentiation of progenitors into Foxj1+ ciliated cells (Guha et al. 2012; Morimoto et al. 2012). This emphasizes the importance of cellular context in determining the phenotype of tumors formed in response to oncogene activation. Future therapeutic opportunities may be identified if the specific factors that drive differentiation in cells proliferating in response to K-Ras induction can be delineated.

K-Ras mutant lung adenocarcinomas in humans may be divided into K-Ras-dependent and K-Ras-independent subsets. K-Ras-dependent tumors are epithelial and may respond better to MEK inhibition, while K-Ras-independent tumors are mesenchymal and may be more resistant to treatment (Singh et al. 2009; Corcoran et al. 2013). These distinctions do not correlate with genetic mutations but are epigenetic. In fact, exogenous administration of TGF-β leads to loss of K-Ras dependence. Interestingly, Sox2 overexpression in the respiratory epithelium leads to induction of TGF-β targets
In addition to lung cancer, Notch signaling is required for progression of pancreatic intraepithelial neoplasia and development of pancreatic ductal adenocarcinoma [Mazur et al. 2010]. It would be interesting to test whether altered Sox2 expression plays a role in K-Ras-driven pancreatic adenocarcinoma development. A future direction will be to determine whether Sox2 is inhibiting K-Ras-induced tumor formation in additional ways that are distinct from Notch inhibition.

In summary, our data support a general model depicted in Figure 7C. According to this model, K-Ras

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### Table 1: Genetic Interaction between Notch, Sox2, and K-Ras in Lung Tumor Initiation and Phenotype

| Alveolar epithelium | Airway epithelium |
|---------------------|------------------|
|                      |                  |
| Wildtype K-Ras      | Wildtype K-Ras   |
| K-RasG12D           | K-RasG12D        |
| Notch wildtype      | Notch wildtype   |
| Sox2 wildtype       | Sox2 wildtype    |
| Notch inhibition    | Notch inhibition |
| Sox2 overexpression | Sox2 overexpression |
| Sox2 reduction      | Sox2 reduction   |

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**Figure 7.** Schematic summary of genetic interaction between Notch, Sox2, and K-Ras in lung tumor initiation and phenotype in the alveoli (A) and airway (B). [C] Model for some of the genetic interactions that determine tumor initiation in the epithelium of the mouse lung. (Top line) Known proximal-distal gradient of Sox2 expression. (Bottom lines) Proposed distal-proximal gradients of potential for Notch pathway activation and K-Ras tumor initiation.
activation of type II cells leads to adenocarcinoma because these cells are able to sustain or maintain Notch signaling when K-Ras is activated. Sox2, which is present in airway cells but only at very low levels in type II cells, suppresses the expression of Notch receptors at the transcriptional level, and this in turn inhibits K-Ras-induced tumor formation.

Materials and methods

Mice

The CC10-CreER, LSL-K-RasG12D, Rosa26-r-f-GFP and Rosa26-LSL-CAG-Sox2-IRES-GFP mouse lines have been described previously (Rawlins et al. 2009; Xu et al. 2012). Rosa26-LSL-DNMaml-GFP mice (Tu et al. 2005) were kindly provided by Warren Pear (University of Pennsylvania, Philadelphia, PA). Rosa26-Notch1ICD-GFP mice [Murthaugh et al. 2003] were from The Jackson Laboratories. Sox2 flox/flox mice (Taranova et al. 2006) were generated by Larysa Pevny (University of North Carolina, Chapel Hill, NC) and kindly provided by Barry Stripp (Cedars Sinai, Los Angeles, CA). All animal experiments were approved by the Duke Medicine Institutional Animal Care and Use Committee.

Tamoxifen administration

A 20 mg/mL tamoxifen (Sigma) solution was made by dissolving tamoxifen in Mazola corn oil. Six-week-old to 8-wk-old mice were injected with 0.25 mg per gram of body weight tamoxifen every other day for four doses.

Histology and immunohistochemistry

For immunohistochemistry, lungs were perfused with PBS and then inflated and fixed with 4% PFA overnight. Fixed lungs were washed three times with PBS, gradually dehydrated in ethanol, and embedded in paraffin. Paraffin sections (7 μm) were washed three times with PBS, gradually dehydrated in ethanol, and embedded in paraffin. Paraffin sections (7 μm) were stained with the following primary antibodies: chicken anti-GFP (1:400; Aves Laboratories, GFP1020), rabbit anti-proSftpc (1:500; Millipore, AB3786), goat-anti CC10 (1:10,000; kindly provided by Barry Stripp), mouse anti-Foxj1 (1:200; eBioscience, 14-9965), rabbit anti-Sox2 (1:1000; Millipore, AB3786), goat-anti CC10 (1:10,000; kindly provided by Barry Stripp), rabbit-anti-Krt5 (1:500; Covance), rabbit anti-Krt8 (1:200; Developmental Studies Hybridoma Bank), and rabbit anti-Sox9 (1:1000; Millipore). Alexa-fluor-coupled secondary antibodies (Invitrogen) were used at 1:400 dilution. All images were captured on a Leica Sp2 laser scanning confocal microscope.

ChIP/sequencing

Approximately 50 million NC1-H520 [American Type Culture Collection, HTB-182] cells were fixed with 1% PFA and then subjected to ChIP with an anti-SOX2 antibody [Cell Signaling, catalog no. 5024] using the Cell Signaling SimpleChIP enzymatic ChIP kit [Cell Signaling, catalog no. 9003] as per the manufacturer’s instructions. Input and SOX2-immunoprecipitated chromatin samples were sequenced at the Duke University Genome Sequencing and Analysis Core. Approximately 31 million input and 26 million SOX2-immunoprecipitated 36-base-pair reads were acquired. Reads were mapped to human genome assembly GRCh37/hg19 using Bowtie via the Galaxy online analysis toolkit [http://galaxyproject.org] [Goecks et al. 2010]. SOX2-binding sites were then predicted using Model-Based Analysis of ChIP-Seq (MACS) version 1.4.1 (Zhang et al. 2008).

Microarray analysis

CC10-CreER, LSL-Sox2 mice and CC10-CreER, Rosa26-r-f-GFP mice were sacrificed 20 wk after tamoxifen injection. Lungs were perfused with 10 mL of PBS and lavaged four times with 1 mL of PBS and 0.2 μM EGTA. One milliliter of 4 U/mL Elastase solution (Worthington Biochemical) was instilled in the airways and incubated for 5 min at 37°C followed by three 0.5-mL instillations for 5 min each. Lung lobes were then minced and incubated with DNase I solution for 10 min at 37°C. Cells were passed through a 100-μm cell strainer, and red blood cells (RBCs) were lysed using RBC lysing buffer [eBioscience, Inc.]. Isolated cells were resuspended at 1 × 10⁶ cells in 100 μL of Hanks’ balanced saline solution, 10 mM HEPES, and 2% fetal bovine serum. Propidium iodide (PI) was used for dead cell discrimination. GFP-positive cells were sorted and collected on FACS Vantage SE. RNA was then extracted using Qiagen protocols. The RNA was reverse-transcribed, and the cDNA was hybridized to Affymetrix 420 mouse microarray chips in the Duke Medicine microarray facility. The resulting CEL files were rma-normalized in Bioconductor in the R environment. Differential expression was carried out using the Limma package with multiple comparisons controlled by the method of Benjamini and Hochberg.

Pathologic analysis

Tumor sections were reviewed by Dr. Thomas Sporn of the Department of Pathology, Duke Medicine.

Tumor analysis

For each mouse, three longitudinal sections of the left lung lobe (600 μm, 800 μm, and 1200 μm from the surface) containing the main axial bronchial pathway were stained with H&E. Tumor size and number were scored using ImageJ software. Airway tumors were assessed by ≥30% of airway space covered by tumor cells, and airways were categorized by lobar bronchi, segmental bronchi, and bronchiole.

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