

(5, 7, 8). Earlier work revealed that airways of smokers with lung cancer exhibit significant differential expression profiles compared with airways of cancer-free smokers (7). Of note, gene expression changes in the airway field of injury have been translated to viable markers for early lung cancer detection. A 232-gene classifier in the bronchial airway (Percepta) was derived that is able to significantly detect lung cancer in suspect current or former smokers (9). Also, a recent study developed and tested a bronchial gene expression classifier that improved the diagnostic performance of endoscopic bronchoscopy for detection of lung cancer in smokers with indeterminate nodules, particularly in intermediate-risk patients when bronchoscopy was nondiagnostic (10).

Studies reporting diagnostic expression profiles in the airway field of injury of smokers were for the most part cross-sectional. We still do not know how and what airway field expression patterns evolve with time following the onset of smoking to signify the development of a nearby lung cancer. To understand the evolution of the airway field of injury, we sought to survey genome-wide temporal airway gene expression changes by RNA sequencing (RNA-Seq) *in vivo*. We studied airway expression changes in mice with knockout of the G-protein-coupled receptor *Gprc5a* (*Gprc5a*^{-/-}) and nicotine-specific nitrosamine ketone (NNK) exposure, which we recently found to develop the most common type of lung cancer, lung adenocarcinoma, within 6 months following exposure (11). Here, we underscore evolutionarily conserved gene expression profiles in the cytologically normal airway field of injury that progress *in vivo* following tobacco carcinogen exposure and up to the development of lung tumors. Notably, we find that a subset of these progressive airway expression changes is significantly enriched in cytologically normal airways of human smokers with lung cancer and is indicative of lung malignancy in suspect smokers with indeterminate nodules. Our current study sheds light on evolutionarily conserved early changes in the normal bronchial airway preceding onset of lung cancer.

Materials and Methods

Chemical and reagents

NNK (purity > 99%) and hematoxylin and eosin (H&E) staining reagents were purchased from Sigma-Aldrich.

Animal housing and carcinogenesis experiments

Gprc5a^{-/-} mice (C57BL/6 × 129sv) were generated as described previously (12) and were maintained according to a protocol approved by the MD Anderson Cancer Center Institutional Animal Care and Use Committee. The mice were housed in a pathogen-free animal facility approved by the American Association for Accreditation of Laboratory Animal Care. Twenty-eight 8-week-old *Gprc5a*^{-/-} mice (16 females and 12 males) were divided into groups of 5 to 6 mice per time point (1:1, 2:3, or 1:2 male to female ratios)

and were injected intraperitoneally with 50 mg/kg of body weight NNK three times per week for 8 weeks (150 mg/kg total per week) dissolved in PBS. Mice were sacrificed at baseline (prior to NNK treatment), immediately after completion of NNK treatment and at 2, 4, and 6 months following treatment. After sacrifice, lungs were excised and inflated by injection with formalin. Lung surface lesions were examined by macroscopic observation. Formalin-fixed lung specimens were then embedded in paraffin (FFPE) and histologic sections were prepared, stained with H&E, and analyzed for lung tumor development by an experienced pathologist (Junya Fujimoto) using previously reported criteria for characterization of murine lung lesions (13).

Collection of mouse tracheal epithelial brushings

Epithelial brushings of the cytologically normal airway from mouse trachea were obtained after mouse sacrifice and based on the method previously described and reported by Sugimoto and colleagues (14). Brushes were made from 60-grit sandpaper-polished polyethylene tube (PE-10 tube; BD Biosciences) with an inserted stainless wire. Brushes (three/mouse) were inserted through an incision into the midsection of the trachea. Initially, brushes were placed in 1 × PBS and cell pellets were stained with pan-cytokeratin to confirm epithelial content and ensure lack of preneoplastic and neoplastic cells. Harvested epithelia from three brushes were collected into 500 μL of Qiazol lysis buffer (Qiagen) and immediately placed on dry ice. Brushes were stored at -80°C until further processing for RNA isolation.

Total RNA isolation from mouse airway brushings

Tubes containing brushes in lysis buffer were vortexed gently to dissociate harvested epithelia and brushes were then removed. Total RNA were purified from the brushings using the miRNeasy Kit from Qiagen according to the manufacturer's instructions. Purified RNA samples were then cleaned and resuspended in nuclease-free water using the RNeasy MinElute Cleanup Kit from Qiagen according to the manufacturer's protocol. RNA quantity was measured using the NanoDrop 1000. Quality of RNA samples was quantitatively assessed by computation of RNA integrity numbers (RIN) using the Agilent RNA 6000 Pico Kit (Agilent) following the manufacturer's protocol.

RNA-Seq

RNA samples with RNA integrity numbers ≥ 7 were selected for RNA-Seq. Libraries were then prepared from poly-A RNAs using the Ion Total RNA-Seq Kit v2 (Thermo Fisher Scientific) according to the manufacturer's protocol for barcoded whole-transcriptome libraries. RNA-Seq was performed using the Ion Torrent Proton platform according to the manufacturer's instructions. On average, we obtained approximately 39.5 million reads per sample. Alignment of

reads was performed using the STAR algorithm v2.4.1d (15) and Bowtie2 v. 2.1.0 (16). Transcripts were quantified (mm10) using a modified version of the expectation-maximization (E/M) algorithm (17). Raw sequencing and normalized expression data of the mouse normal-appearing airways ($n = 28$ samples) are deposited in the gene expression omnibus (GEO) under series GSE102707 (samples GSM2743377 – GSM2743404).

Gene expression analysis

A linear model was used to identify gene expression profiles that were differentially expressed in the mouse airway following NNK exposure ($t = 0$) relative to baseline. A Benjamini–Hochberg controlled FDR was used to identify significant features based on a q -value cutoff. A linear model was also applied to identify profiles that were differentially expressed at each time point following NNK exposure ($t = 2$, $t = 4$, and $t = 6$) relative to $t = 0$. All statistical analyses and hierarchical clustering were conducted using the R statistical language environment v3.1.1 and the *limma* package v3.22.7.

Comparative genomics analysis

Comparative genomics analysis was used to analyze mouse airway expression changes in human airway datasets. To analyze changes influenced by NNK (at end of exposure vs. baseline), the previously reported human expression dataset by Spira and colleagues (18) consisting of airway samples from phenotypically healthy current smokers ($n = 34$) and nonsmokers ($n = 23$) was interrogated. Orthologues from the murine airway expression profiles were determined, ranked by the t -statistic of the coefficient for smoking status from a linear model, and analyzed by gene set enrichment analysis (GSEA; ref. 19) in the human smoking dataset. Temporal mouse airway expression changes were similarly analyzed by GSEA in the recently reported array dataset by Silvestri and colleagues (10) of bronchial brushings from smokers with suspicion of lung malignancy ($n = 253$ and $n = 90$ with and without lung cancer, respectively) and who had undergone endoscopic bronchoscopy for diagnosis. The expression of a recently reported bronchial 232-gene classifier (9) was also analyzed in the temporal post-NNK mouse airway samples.

Detailed methods are available in the Supplementary Methods accompanying the article.

Results

Analysis of temporal genome-wide gene expression changes in the mouse airway following tobacco carcinogen exposure and preceding lung adenocarcinoma

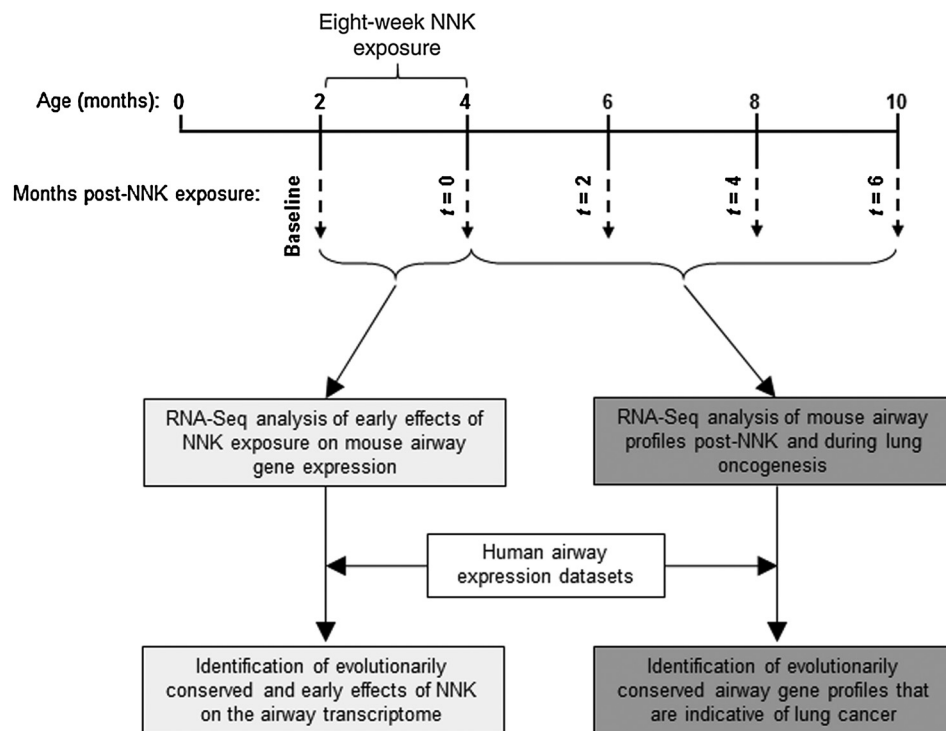
The progression of airway expression changes following smoking and prior to lung cancer onset is poorly understood. This is largely due to the long latency of human lung

malignancy and, thus, the difficulty in temporally tracking airway field effects following smoking. To fill this void, we surveyed, by RNA-Seq, genome-wide temporal expression changes in the normal-appearing airway that occur prior to and during oncogenesis of a nearby lung tumor (Fig. 1). We studied *Gprc5a*^{-/-} mice exposed to the tobacco-specific carcinogen NNK, which we recently found to develop the most common histologic subtype of lung malignancy, lung adenocarcinoma, within 6 months following exposure (Fig. 1; ref. 11). Groups of five to six 8-week-old mice ($n = 28$ total) were intraperitoneally administered 50 mg/kg bodyweight NNK three times per week for 8 weeks. Airway (tracheal) epithelia were collected by brushing and studied at baseline (prior to NNK exposure) and immediately following the 8-week exposure ($t = 0$) to understand early effects of tobacco carcinogen exposure on the airway transcriptome (Fig. 1). Mice were followed up at two ($t = 2$), four ($t = 4$), and six ($t = 6$) months following completion of NNK exposure for lung tumorigenesis. Lung tumor burden was significantly increased with time post-NNK particularly at $t = 4$ and $t = 6$ ($P < 0.001$ of the ANOVA test, Supplementary Fig. S1). Airway epithelia were also collected every 2 months following completion of NNK treatment ($t = 2$, $t = 4$, and $t = 6$) and studied to understand progressive or lasting field effects following exposure and until tumor development ($t = 6$; Fig. 1). Comparative genomics analyses were also performed, using expression datasets of normal airways from human smokers, to determine evolutionarily conserved profiles including those that can diagnose lung cancer in smokers with suspicion of the disease (Fig. 1).

Early evolutionarily conserved expression changes in the mouse airway following tobacco carcinogen exposure

We first compared and contrasted the transcriptome of mouse airways prior to (baseline) and immediately following completion of NNK exposure ($t = 0$; $n = 6$ samples each). Using a linear model and a controlled (by the Benjamini–Hochberg method) FDR threshold of 0.1%, we found 926 gene features (529 upregulated and 397 downregulated) that were significantly differentially expressed in the airway following NNK exposure (Fig. 2A; Supplementary Table S1). Hierarchical clustering revealed that the identified profiles clustered airway samples according to NNK exposure status (Fig. 2A). We then sought to determine from the murine airway profiles those that are enriched in cytologically normal airways of human smokers. We interrogated a reported expression dataset of cytologically normal airway (mainstem bronchi) brushings obtained from phenotypically healthy (cancer-free) current smokers ($n = 34$) and nonsmokers ($n = 23$; ref. 18). After matching gene features between our study and the human airway dataset, 853 remained (491 upregulated and 362 downregulated by NNK). These orthologous

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**Figure 1.**

Identification of global gene expression changes in the normal-appearing airway during the development of lung adenocarcinoma. Study design depicting analysis of genome-wide changes in gene expression in the tobacco-carcinogen normal-appearing airway *in vivo*. Two-month-old *Gprc5a*^{-/-} were treated intraperitoneally with 50 mg/kg bodyweight nicotine-specific NNK three times per week for 8 weeks. Upper airway epithelia were collected by brushings before NNK treatment (baseline) and at the indicated four time points after completion of NNK exposure as described in the Materials and Methods section. Total RNA was isolated and interrogated by RNA-Seq using the Ion Torrent Proton platform. Gene profiles were statistically analyzed to determine patterns that are impacted early on in the airway by NNK exposure ($t = 0$ vs. baseline) and those that temporally evolve for 6 months following completion of the treatment and during the development of lung adenocarcinoma. Using publicly available human airway expression datasets, cross-species comparative genomics analysis (see Materials and Methods and Supplementary Methods) was performed to identify genes from the mouse airway expression profiles that are evolutionarily conserved in airways of human suspect smokers, including patients with lung malignancy.

profiles were then ranked according to the t statistic between human smokers and nonsmokers and analyzed by GSEA in the human dataset. Of the 362 downregulated genes in the airway by the tobacco carcinogen, 54 genes were found to be statistically significantly enriched ($P < 0.001$) within gene features downregulated in airways of human current smokers compared with airways of never smokers (Supplementary Fig. S2; Supplementary Table S2). In addition, the enriched gene features, following hierarchical clustering analysis, not only separately clustered mouse airway samples by NNK exposure status (Fig. 2B, left) but also significantly ($P < 0.001$ of the Fisher exact test) distinguished airways of human smokers from those of nonsmokers (Fig. 2B, right). Of note, similar effects were found with gene features upregulated by NNK, albeit they did not reach statistical significance (data not shown). These data underscore early expression changes by tobacco carcinogen (NNK) in the mouse airway that are also present in airways of humans smokers and are thus likely to be evolutionarily conserved in their response to smoking.

Early expression changes in the mouse normal-appearing airway field of injury that signify lung cancer in human smokers

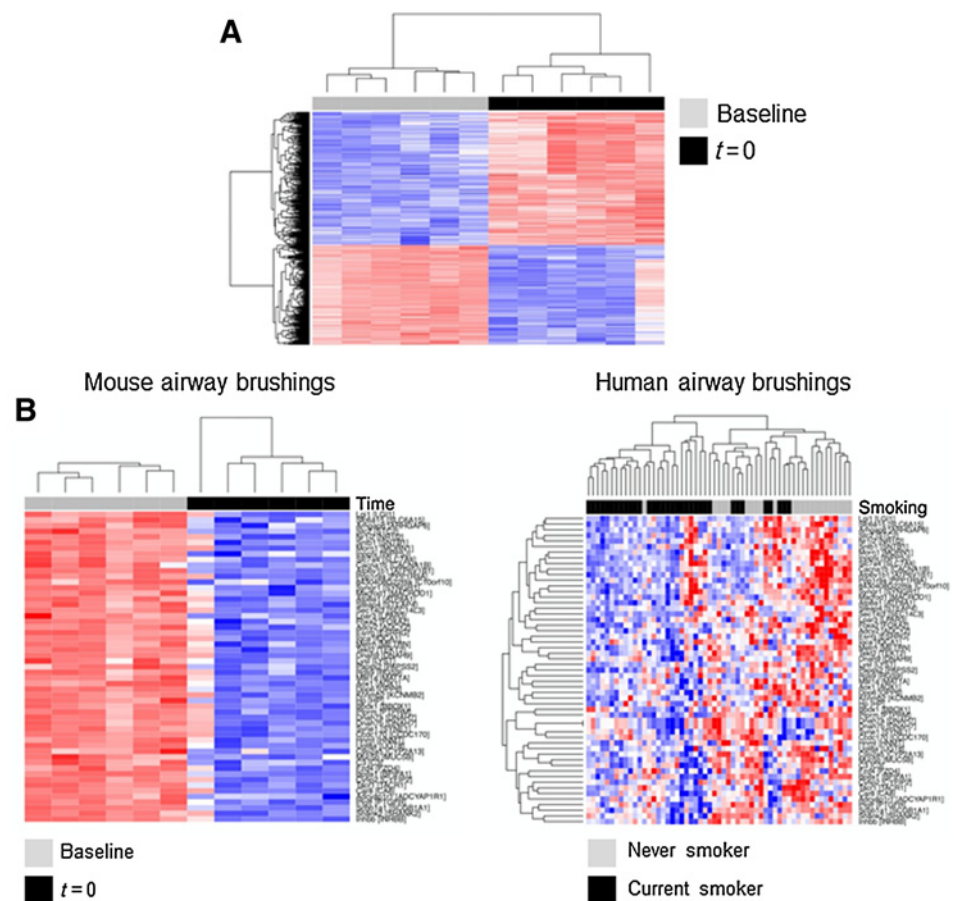
We then sought to pinpoint global gene expression profiles in the airway that progress following tobacco carcinogen exposure and during lung oncogenesis. Using linear models and a controlled FDR threshold of 0.5%, we identified gene profiles in the mouse airway that were differentially expressed at 2 ($n = 362$), 4 ($n = 795$), and 6 ($n = 980$) months following NNK relative to the time ($t = 0$) at completion of carcinogen treatment (Supplementary Table S3). Notably, changes occurring early on at 2 months following NNK ($t = 2$) were still evident in the airways of mice at both 4 and 6 months following carcinogen treatment (Supplementary Fig. S3). Next, we probed the relevance of the mouse airway field of injury expression changes to the airway in human smokers and in the context of lung cancer. We studied the recently reported expression dataset by Silvestri and colleagues (10) comprised of cytologically normal airway (mainstem bronchi) brushings obtained from smokers who had undergone bronchoscopy for

Figure 2.

Evolutionary conserved effects of NNK tobacco carcinogen exposure on airway gene expression.

A, Differentially expressed profiles ($n = 926$; 529 upregulated and 397 downregulated) between mouse airways at the time of completion of NNK exposure ($t = 0$) relative to baseline were identified using a linear model and a controlled FDR threshold of 0.1% (see Materials and Methods).

B, Using orthologues from the identified genes, GSEA was performed to identify expression profiles that are concordantly (in same direction) enriched in airways of human cancer-free current smokers relative to nonsmokers. Statistically significant leading edge set of 54 downregulated orthologous genes was analyzed by semisupervised clustering in the mouse airway brushings (left). The same genes were assessed by unsupervised clustering in the human airway brushings (right). Columns represent samples and rows denote gene features; red and blue: higher and lower expression, respectively.



suspected lung cancer (with lung cancer: $n = 253$; without: $n = 90$). After matching orthologous mouse and human features, genes were ranked according to the t statistic between airways of human smokers with and without lung cancer and based on a linear model incorporating final diagnosis of lung cancer as well as other clinicopathologic variables including chronic obstructive pulmonary disease status, age, and gender (Supplementary Methods). Following GSEA, we identified subsets within gene features differentially expressed at each time point following NNK exposure in the mouse airway that were overall enriched and concordantly (i.e., in same direction) modulated in airways of smokers with lung cancer relative to cancer-free smokers (Fig. 3A). We paid particular attention to orthologous profiles that were induced at $t = 2$, the earliest time point we had sampled following NNK. Of these early changes, we identified a profile of 22 orthologous genes (Supplementary Table S4) that not only persisted (downregulated) at later time points (4 and 6 months) following NNK (Fig. 3B, left), but were also significantly ($P < 0.001$) and concordantly enriched (downregulated) in the airways of human smokers with lung cancer relative to cancer-free smokers (Fig. 3B, right).

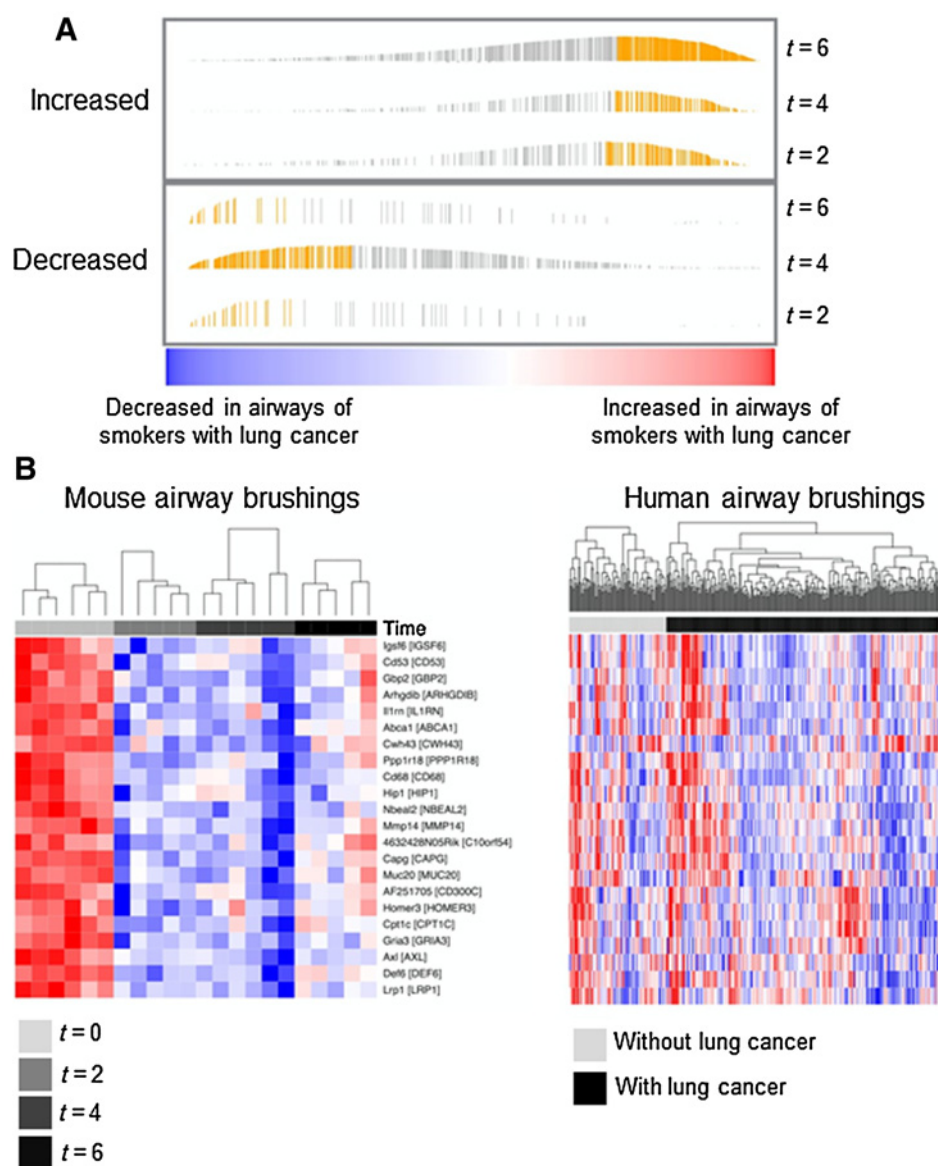
We then sought to interrogate the relevance of the identified early downregulated airway field of injury gene

expression changes in the mouse (at $t = 2$, $n = 68$ genes) to profiles in lung tumors themselves. We interrogated by GSEA orthologous features of the 68-gene profile we found to be downregulated in the mouse normal airway at $t = 2$ in publicly available expression datasets of lung adenocarcinomas/NSCLCs and paired normal lung tissues (20–22). The profile was also assessed in six NNK-exposed *Gprc5a*^{-/-} lung tumors (that developed in 5 *Gprc5a*^{-/-} mice at 4–7 months following NNK exposure) and adjacent normal lung tissues that we also surveyed by RNA-Seq (See Supplementary Methods). We found that the 68-gene profile was significantly (all $P < 0.05$) suppressed in human lung adenocarcinomas/NSCLCs (Supplementary Fig. S4A–S4C) as well as in mouse lung tumors relative to their respective normal lung tissues (Supplementary Fig. S4D). These findings point to early changes in the evolution of the molecular airway field of injury following tobacco carcinogen exposure that persist even after tobacco cessation and may signify the ensuing development of a cancer in the lung epithelial field.

Cross-species temporal analysis of a human bronchial genomic classifier

A recent report studying cytologically normal airways in suspect lifetime smokers who have undergone diagnostic

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**Figure 3.**

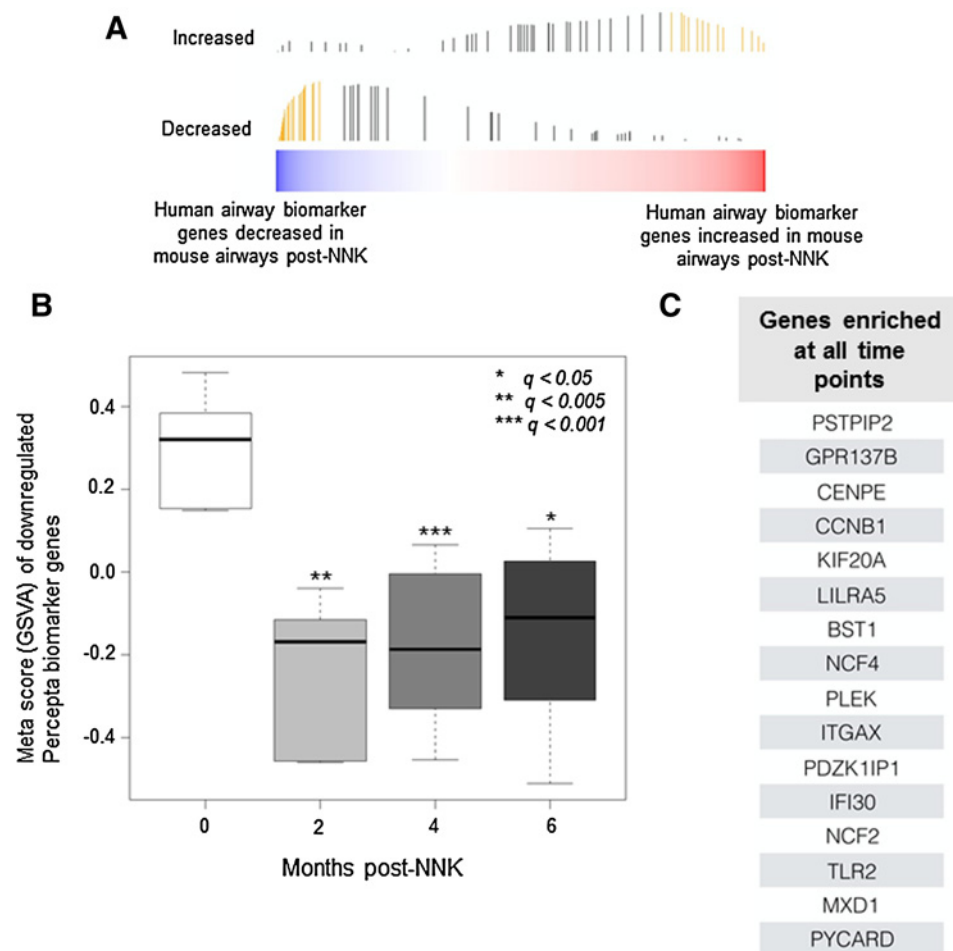
Temporal mapping of the molecular airway field of injury upon following tobacco carcinogen exposure and during *in vivo* development of lung adenocarcinoma. **A**, Murine genes changing in the normal-appearing airway at two ($t = 2$), four ($t = 4$), and six ($t = 6$) months following completion of NNK exposure and relative to the time at the end of the treatment ($t = 0$) were statistically identified as described in the Materials and Methods section and Supplementary Methods. The identified temporal mouse profiles were analyzed by comparative genomics and GSEA in the reported array dataset by Silvestri and colleagues (10), comprised of bronchial brushings from smokers with suspicion of lung malignancy (253 with and 90 without lung cancer). Enrichment plots generated by GSEA of ranked gene expression are depicted demonstrating enriched up- and down-regulated murine genes post NNK and at the indicated time points. Bar heights correspond to enrichment scores and yellow bars correspond to genes in the leading edge subset that were concordantly enriched in airways of human smokers with lung cancer relative to cancer-free smokers. **B**, Concordantly enriched genes that were differentially expressed at the earliest time point ($t = 2$) following NNK exposure were interrogated to derive a leading edge set consisting of 22 downregulated genes. The leading edge set was then analyzed by hierarchical semisupervised clustering in mouse (left) and human smoker (right) airway brushings. Columns, samples; rows, gene features (red, higher; blue, lower expression).

bronchoscopy derived bronchial gene expression profiles that are significantly indicative of lung cancer status (114 upregulated and 118 downregulated genes in airways of smokers with lung cancer; ref. 9). Here, we sought to begin to understand the nature of development and progression of this human bronchial gene signature prior to lung cancer onset. We cross-species examined the human signature (9) in our temporal murine airway field of injury RNA-Seq dataset. Following gene set variation analysis (GSVA), we found that 56 of the 118 downregulated genes were significantly ($q < 0.005$) and concordantly reduced in the mouse airway field of injury (Fig. 4A). Collectively, when summarized as a metacore, these conserved markers were significantly downregulated in the mouse airway field of injury following NNK exposure, a subset of which was

suppressed as early as 2 months following NNK treatment (Fig. 4B). Among these leading edge genes, 16 were found alternatively by GSEA to be statistically significantly enriched in the mouse airway at all tested time points following NNK exposure (Fig. 4C). Pathway analysis of the leading edge set revealed significant modulation of molecular and biological processes associated with the inflammatory response as well as leukocyte migration, suggesting early aberration of immune signaling in the tobacco carcinogen-exposed airway field (Supplementary Table S5). These findings point to airway field genes that are not only indicative of lung cancer in smokers with indeterminate nodules, but are also induced or evolve prior to tumor onset and, thus, may serve as high-potential surrogate markers to predict lung cancer in high-risk smokers.

Figure 4.

Human to mouse cross-species analysis of a human bronchial genomic classifier for lung cancer detection. **A**, A recently reported human bronchial 232-gene classifier (9) was analyzed by GSEA (see Supplementary Methods) in the identified mouse airway profiles that were found to be modulated *in vivo* following NNK exposure (bar heights correspond to running enrichment score calculated in GSEA and yellow bars correspond to the leading edge subset). **B**, Meta scores were plotted based on GSEA of a 56-gene leading edge set from the Percepta classifier that is downregulated in airways of human smokers with lung cancer relative to cancer-free smokers and concordantly decreased in the temporal mouse airway brushings. Meta scores were derived and plotted for conserved markers, which were concordantly downregulated at 2, 4, or 6 months after NNK treatment. Statistically significant differences among the different time points were based on a q -value cutoff (*, < 0.05 ; **, < 0.005 ; ***, < 0.001). **C**, GSEA (see Materials and Methods) of the 232-gene human classifier was performed to identify, from the human biomarker, a gene set that was concordantly enriched in the mouse airway samples across all time points.



Discussion

The nature of the development and progression of the molecular airway field of injury following tobacco is poorly understood. In this study, we sought to determine temporally progressive airway field effects that can elucidate quintessential airway expression signatures during tobacco carcinogen-induced lung tumorigenesis. Following RNA-Seq analysis, we identified evolutionarily conserved airway field changes that are not only modulated early on by the tobacco carcinogen NNK, but are also prevalent and enriched in smoking-exposed cytologically normal airway epithelia of human smokers relative to airway epithelia of human nonsmokers. This airway whole-transcriptome survey also revealed expression profiles that are temporally modulated *in vivo* following tobacco carcinogen exposure prior to and during lung tumor development. By cross-species comparative analysis, we found that a subset of these early post-tobacco murine airway expression changes was significantly enriched in airways of human suspect smokers with lung cancer relative to those without the malignancy. In addition, a recently derived airway expression classifier for lung cancer among suspect smokers (9)

was found to be enriched in the identified mouse orthologous airway profiles. Furthermore, genes found to be suppressed in the mouse normal airway as early as 2 months post-NNK were overall concordantly decreased in mouse and human lung tumors relative to normal lung tissue, pointing to the probable roles of these early deregulated airway profiles in lung oncogenesis. Our study highlights airway expression changes that occur and progress early on during tobacco carcinogen-associated lung cancer development. These changes, otherwise difficult to survey (e.g., longitudinally) in humans, may epitomize the early development of nearby lung lesions within this mutagenized field of injury and, thus, may serve as a low hanging fruit of biomarkers for early detection.

Earlier work from our group and others characterized gene expression changes in the airway field of injury of human smokers and derived markers that are indicative/diagnostic of lung cancer among suspect smokers (5, 7, 9, 10, 20, 23, 24). These studies were largely cross-sectional, that is, they compared, following diagnostic bronchoscopy, airway profiles of smokers with lung cancer relative to those without the malignancy at a specific point in time.

Because these studies were not aligned with time, the question remains on how airway expression patterns change following tobacco carcinogen exposure and prior to lung cancer onset. Specifically, it is not clear whether airway expression profiles that are differentially expressed in smokers with lung cancer relative to cancer-free smokers are related, phenotypically, to the already developed tumor or whether these airway profiles are a consequence that emerged sometime after nearby tumor development. This is particularly significant in light of the long latency of lung cancer development in humans (25). In this context, temporal tracking of tobacco-induced and progressive airway field effects, presumed to recapitulate the genotype of nearby lung tumors that emerged from the field, are arduous to characterize and thus have remained elusive. We sought to fill this void by surveying airway expression changes that occur with time following tobacco carcinogen exposure *in vivo*. We employed RNA-Seq and a lung cancer model comprising knockout of the lung-specific tumor suppressor *Gprc5a* and that we previously showed to develop lung adenocarcinomas following NNK exposure (11, 26). Our first goal here was to characterize effects of NNK exposure on the airway transcriptome by RNA-Seq analysis of airway brushings pre- and post-NNK ($t = 0$). We first pinpointed murine airway expression profiles that were not only altered by tobacco carcinogen but were also evolutionarily conserved in cancer-free human smokers relative to airways in nonsmokers. Biological processes (gene ontology annotated) of a number of gene products identified in those profiles include response to oxidative stress (*Hspb6*, *Txnrd3*, *Hopx*, *Bcl2l11*, *Cat*), inflammation (*Mgst2*, *Ccl28*, *Tgfb1*, *Il7ra*, *Il1rn*, *Tlr2*, *Itgb2*) as well as immune regulation (*Cybb*, *Tnfrsf19*, *Csf2ra*, *Csf2rb*, *Csf2rb2*, *Stk39*, *Tlr7*, *Elf4*, *Cd93*, *Fcer1g*), and those genes were similarly modulated in both NNK-exposed mouse airway and human smokers. Of note, previous studies have demonstrated modulation of oxidative stress, inflammation, and immune reaction related pathways in the molecular airway field of injury (2). These findings suggest that the effects of NNK on the mouse airway transcriptome recapitulate genome-wide effects of cigarette smoke in human airways. It is noteworthy that the results presented herein showed that the murine NNK-modulated genes that were evolutionarily conserved and enriched in human smokers (relative to nonsmokers) were particularly downregulated (compared with nonsmokers). This is in agreement with previous reports showing prevalence of dampened gene expression in the airway field of injury in smokers with early-stage NSCLC (20). Our findings warrant further studies that can potentially survey these profiles as possible early molecular markers of smoking exposure.

By serial analysis of the airway transcriptome up to the time of lung adenocarcinoma development, our study next underscored airway expression changes that develop and/or progress following completion of tobacco carcinogen

exposure and during lung oncogenesis. Our study design enabled temporal tracking of changes in airway gene expression before, immediately after exposure to NNK, and at various time points following exposure to the tobacco carcinogen. Notably, from these temporal airway expression profiles, we identified by cross-species comparative analysis those that were significantly enriched and concordantly modulated in airways of human smokers with lung cancer relative to brushings from cancer-free smokers. In retrospect, our analysis thus identified, from gene profiles previously reported to be dissimilar between airways of smokers with cancer relative to cancer-free smokers (10), profiles that likely emerged following tobacco exposure and prior to lung cancer development. It is noteworthy that mouse airway field of injury profiles that were downregulated, but not elevated, post-NNK were suppressed in cytologically normal airways of human smokers with lung cancer relative to cancer-free smokers. We also showed that these early downregulated profiles in the mouse normal airway field of injury (at $t = 2$) were also modulated in the same direction (downregulated) in both human and mouse lung tumors themselves relative to normal lung tissues. It is plausible that these progressive airway profiles are not only biomarkers for lung malignancy but also causally related to ensuing lung tumor development within the mutagenized airway field of injury. It is reasonable to surmise that downregulated expression of the identified genes in the airway field of injury may be, in part, mediated by methylation, a known molecular genome-wide mechanism in smoking-associated lung oncogenesis (27). It is important to note that upon completion of 8 weeks of NNK exposure, we compared gene signatures after 2, 4, and 6 months to the onset signature ($t = 0$), thereby excluding direct/immediate effects post-NNK and those possibly occurring during carcinogen exposure. By virtue of this analytic design, it is thus plausible that our study centered on airway field of injury profiles that are induced or progressed following cessation of smoking activity, thereby signifying the human former smoker phenotype and the capacity to elucidate the significance of residual effects of smoking in lung cancer development. Indeed, despite reduction in relative lung cancer risk in former smokers, the risk never returns to baseline (6). Also, former smokers constitute up to half of patients diagnosed with lung cancer (6). Future studies are warranted to determine the precise relevance of profiles in our study to former smokers (as compared with current smokers).

Further analysis revealed that a significant fraction of the identified temporal airway field of injury profiles play crucial roles in the host immune response. For example, immunoregulatory receptors *C10orf54* and *Cd300c* can affect T-cell/natural killer cell-mediated ligand-receptor interactions or inflammatory responses (28, 29). Other previously reported functions of the profiles identified include Th1 responses (*Def/SLAT*; ref. 30), evasion of

immunosurveillance (*Mmp14*; ref. 31), immune infiltration, tumor proliferation, progression, and metastases (*Lrp1*; ref. 32), cancer cell survival (loss of *Abca1*; ref. 33), while some have emerged as risk markers for lung cancer (*Il1rn*; ref. 34). Of note, our temporal RNA-Seq analysis demonstrated that the immune marker *Cd68* and the tyrosine receptor kinase *Axl* were downregulated across all surveyed time points post-NNK exposure. *Cd68* plays a crucial role in tumor-associated macrophage infiltration, and its expression was found to be significantly associated with favorable prognosis in NSCLC (35). The *Axl* kinase was found to influence immunosurveillance of tumor cells (36). To further support the significance of immune-related functions of the identified temporal airway field of injury profiles, several of the concordantly enriched Percepta (9) biomarker orthologues were shown to possess immunomodulatory roles (Supplementary Table S5). These findings suggest that the identified early airway field of injury changes (i.e., 68 downregulated genes at $t = 2$) comprise early immunomodulatory effects. Although expression of immune markers (e.g., the immune checkpoint PD-L1; ref. 37) or immunotherapeutic response (38) has been shown to be associated with specific genomic features (e.g., mutation burden), the interplay between the molecular airway field of injury and markers of the immune response is still extremely poorly understood. Emerging and ongoing efforts have been put into understanding the clinical ramifications of the dynamic interplay between tumors and the host immune response (39). In lung cancer, compromised antitumor immunity is associated with poor prognosis (35), and immune therapies that recuperate the host immunity have proven successful (40, 41). In this context, it is intriguing to suggest that modulation of the host immune response occurs very early on following smoking and in the pathogenesis of lung cancer (42), thus providing a framework for immune-based prevention of this fatal disease.

Our study is not without limitations. We acquired epithelial brushings from mice following euthanasia (14) given the difficulty of obtaining such samples from mice that continue to survive over a long course. Thus, true longitudinal assessment of temporal expression patterns in the same variables (mice) was limited by mice euthanasia. Yet, our study underscored global expression patterns that denoted temporal progression of the airway field of injury *in vivo*, findings that warrant future confirmation. In addition, although our study employed the nicotine-specific and derived carcinogen NNK, cigarette smoke comprises other potentially carcinogenic constituents, including polycyclic aromatic hydrocarbons among others (43). Considering this, cigarette smoke exposure may exhibit additional or disparate effects on the airway transcriptome. Nonetheless, earlier work showed that NNK is one of the most prevalent carcinogenic constituents in cigarette smoke (43). In addition, in our current study, mice were not

exposed to a single dose of NNK; rather, they were exposed to NNK continuously over a course of 2 months to attempt to mimic prototypical chronic cigarette/nicotine exposure in human smokers. Also, our cross-species genomic analysis demonstrated that NNK exposure in the mouse does recapitulate known transcriptomic effects of cigarette smoke in human airways. An additional factor to consider in our study is the specificity of the mouse model (*Gprc5a*^{-/-}) employed to lung adenocarcinoma (11, 44). It is reasonable to surmise that the temporally progressive airway field of injury profiles presented in this study are most likely relevant to the pathogenesis of lung adenocarcinomas. Considering that lung adenocarcinomas constitute the major histologic subtype (~65%) among NSCLCs (5), our study's findings are applicable to understanding the pathogenesis of a significant fraction of lung malignancies. Adding to this, our group previously reported that protein expression of human GPRC5A itself was significantly decreased in the normal-appearing airway field of injury of smokers with NSCLC relative to cancer-free smokers (45). Also, we recently found, by whole-exome sequencing, that lung adenocarcinomas from tobacco carcinogen-exposed *Gprc5a*^{-/-} mice exhibit activating mutations in *Kras*, the most commonly mutated oncogene in smoker lung adenocarcinomas (11). Thus, it is conceivable to construe that the tobacco carcinogen-exposed *Gprc5a*^{-/-} mouse is useful to interrogate the temporal evolution of at least a significant molecular fraction (e.g., *Kras* mutant) of smoking-associated lung adenocarcinomas. Future studies are warranted to probe the evolution of airway field effects following tobacco carcinogen and during oncogenesis of other lung cancer subtypes. Along these lines, Xiong and colleagues (46) identified evolutionary conserved early airway expression alterations in a mouse model of lung squamous cell carcinoma and that are modulated in response to chemopreventive agents.

In conclusion, by surveying transcriptome-wide alterations in cytologically normal airways *in vivo*, we delineated evolutionarily conserved expression changes that occur early on during tobacco-associated lung oncogenesis and that are otherwise difficult to ascertain during the long latency of lung cancer development in human smokers. We underscored evolutionarily conserved airway expression profiles that are modulated by NNK in the mouse and that are significantly and concordantly enriched in airways of human smokers relative to nonsmokers. We also identified murine airway profiles that temporally progress in the epithelial field following NNK exposure and that are significantly enriched in airways of human smokers with lung cancer relative to cancer-free smokers, as well as in both murine and human lung tumors relative to normal lung tissues. Our findings point to early evolutionarily conserved changes in the normal-appearing exposed airway that embody the development of nearby lung lesions within the mutagenized field and thus serve as minimally invasive biomarkers and targets for early detection and prevention.

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Disclosure of Potential Conflicts of Interest

A.E. Spira has ownership interest (including patents) in and is a consultant/advisory board member for Veracyte Inc. No potential conflicts of interest were disclosed by the other authors.

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