The Differential Effects of Mutant p53 Alleles on Advanced Murine Lung Cancer

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Abstract

We report a direct comparison of the differential effects of individual p53 mutations on tumor growth and progression, and the creation of a murine model of spontaneous advanced lung adenocarcinoma that closely recapitulates several aspects of advanced human pulmonary adenocarcinoma. We generated compound conditional knock-in mice with mutations in K-ras combined with one of three p53 alleles: a contact mutant, a structural mutant, or a null allele. p53 loss strongly promoted the progression of K-ras–induced lung adenocarcinomas, yielding a mouse model that is strikingly reminiscent of advanced human lung adenocarcinoma. The influence of p53 loss on malignant progression was observed as early as 6 weeks after tumor initiation. Furthermore, we found that the contact mutant p53R270H, but not the structural mutant p53R172H, acted in a partially dominant-negative fashion to promote K-ras–initiated lung adenocarcinomas. However, for both mutants, loss-of-heterozygosity occurred uniformly in advanced tumors, highlighting a residual tumor-suppressive function conferred by the remaining wild-type allele of p53. Finally, a subset of mice also developed sinonasal adenocarcinomas. In contrast to the lung tumors, expression of the point-mutant p53 alleles strongly promoted the development of sinonasal adenocarcinomas compared with simple loss-of-function, suggesting a tissue-specific gain-of-function. (Cancer Res 2005; 65(22): 10280-8)

Introduction

Lung cancer is the leading cause of cancer deaths worldwide, with 157,200 deaths predicted for the year 2003 in the United States alone (1). Adenocarcinoma, a subtype of non–small-cell lung cancer, is the single most common form of lung cancer, comprising ~40% of cases (1). Activating mutations in the K-ras proto-oncogene are found in ~30% of human non–small-cell lung cancers (2). Recently, several mouse lung cancer models have been created using conditionally or spontaneously activatable alleles of oncogenic K-ras (3–6). These models have contributed significantly to our understanding of the role of K-ras in tumor initiation and maintenance of the tumor phenotype but do not recapitulate all aspects of the human disease. In particular, the lung tumors in these models resemble early-stage human lung adenocarcinomas.

Features of late-stage disease including invasion, stromal desmoplasia, and metastasis are not prominent.

Alterations in the p53 tumor suppressor gene are also common in non–small-cell lung cancer, occurring in ~50% of cases (7). As in other cancers, the majority of these alterations are missense mutations that result in the accumulation of high levels of mutant p53 protein. Missense mutations in p53 occur primarily in the DNA-binding domain but the frequency of mutation at individual codons varies dramatically between tumor types. The most common p53 missense mutations in human pulmonary adenocarcinomas occur in descending order at codons 273, 248, 249, 245, and 158 (8). Of note, codon 175 mutations are significantly less common in lung adenocarcinomas than in other cancers. This discrepancy is typically explained by the fact that different codons vary in their exposure and susceptibility to mutagens. About 90% of lung tumors are associated with exposure to carcinogens from tobacco smoke and the polycyclic aromatic hydrocarbons found in tobacco smoke preferentially bind to and form adducts with several of the p53 codons frequently mutated in lung cancer. However, these compounds also efficiently form adducts at codon 175, weakening this theory (9).

An alternative explanation for the variation in mutation frequency between tumor types is a difference in the tumorigenic potential of individual mutant p53 proteins in different tissues. p53 mutations are commonly grouped into two classes that display different behaviors in several in vitro assays (10). Contact mutations alter residues that directly contact DNA whereas structural mutations alter residues that are critical for maintaining global domain structure (11). Structural mutants are frequently described as more potent than contact mutants in promoting cancer. In general, point-mutant p53 confers an adverse prognosis in patients with pulmonary adenocarcinoma but the tumorigenicity of individual mutations has not been determined (12, 13).

Tumor-derived p53 mutations may have two effects beyond loss of function. There is evidence that mutant p53 alleles can act in a dominant-negative manner to inhibit the function of wild-type p53 through hetero-oligomerization between mutant and wild-type p53 polypeptides. In vitro studies have shown that both classes of mutants can exert dominant-negative effects on wild-type p53 activity with the effects being stronger for structural mutants (14). Furthermore, certain point-mutant p53 alleles have been shown to promote tumorigenesis through gain-of-function effects that are not dependent on wild-type p53 (15).

We recently generated two strains of conditional point-mutant p53 mice that allow for endogenous expression of mutant p53 on Cre-mediated recombination. The first strain carries the contact mutation p53R270H (homologous to human p53R273H) whereas the second strain carries the structural mutation p53R172H (homologous to human p53R175H; ref. 16).
We report here the development of an improved murine lung cancer model through the generation of compound mutant animals with conditional mutations in K-ras and p53. This model closely recapitulates several aspects of advanced human pulmonary adenocarcinoma not commonly seen in existing models. We have used this model to directly compare the oncogenic properties of different p53 mutations (p53R270H and p53R172H and a large deletion mutation of p53) in the context of a single tumor type initiated by a defined oncogenic stimulus. Our comparison of K-ras-induced lung tumors with mutant or null alleles of p53 clearly shows a dominant-negative effect by the p53R270H allele that is not conferred by the p53R172H allele. Furthermore, we describe a tissue-specific gain of oncogenic potential conferred by both the p53R172H and the p53R270H alleles toward the development of carcinomas of the sinonasal mucosa.

Materials and Methods

Breeding schemes. LSL-KrasG12D mice were crossed to p53Fl/+ mice to generate LSL-KrasG12D; p53Fl/+ (KFl/+). The p53R172H and p53R270H alleles were each crossed to the p533' deletion strain to generate p53R172H; p533'Fl/+ and p53R270H; p533'Fl/+ mice. By crossing KFl/+ mice with p53R172H; p533'Fl/+ and p53R270H; p533'Fl/+ mice, offspring of the following genotypes were generated for use in tumor studies: KFl/+; K172/+; KFl/Fl; K270/+; and K270/Fl.

Molecular analysis of recombination efficiency. DNA was prepared from tissue samples. For the K-ras allele, PCR was done using the Advantage-GC-eDNA kit from Clontech (Mountain View, CA) with primers flanking the lox-stop-lox cassette: K5' forward, 5'GGGATGTTTGGGAGTAGCTG-3' and K3' reverse, 5'TCCGAATCTAGACGATAGTACAGA-3'. The wild-type and recombined conditional K-ras alleles yield 265- and 305-bp products, respectively. Analysis of the conditional p53 point-mutant alleles was done using standard PCR buffer with primers flanking the lox-stop-lox cassette: dt02000.1 forward, 5'AGCTGTGCTACTCTCTCAGG-3' and dt011200.3 reverse, 5'TCTTGGGGACATACCCACAGT-3'. The wild-type and Flox p53 alleles produce a 291-bp product and the recombined conditional allele produces a 325-bp product. PCR analysis of the floxed allele was done using standard PCR buffer with primers flanking theloxP sites in exons 2 and 10: forward, 5'CATAAAAACAGGTTAACCACG-3' and reverse, 5'GGAGACGAGAAAAGGGGGGAGG-3'. The recombined allele yields a 612-bp product; no product is produced from the wild-type allele.

AdenoCre infection. Mice were infected with 5 x 106 plaque-forming units (pfu) of AdenoCre virus at 6 to 8 weeks of age. AdenoCre:CaPi coprecipitates were prepared as described (17). Mice were anesthetized with avertin. AdenoCre:CaPi coprecipitates were administered intranasally in 20 µL of 0.9% saline. Immunohistochemistry and trichrome staining. All lungs were sectioned as follows: five-step sections were taken at 100 µm apart, with five unstained sections taken after section 3. p53 immunohistochemistry was done following Trifecta dextran-citrate buffer unmasking according to instructions of the manufacturer. p53 CMS rabbit polyclonal antibody (Novo Castra, Newcastle upon Tyne, United Kingdom) was used at a 1:500 dilution. Staining was completed with a horseradish peroxidase anti-rabbit kit (Vector Labs). For phospho-mitogen-activated protein kinase (MAPK) immunohistochemistry, high-temperature citrate buffer unmasking was done and anti-phospho-p42, MAPK (Cell Signaling, Beverly, MA) was diluted 1:100. Sections were incubated in primary antibody overnight at 4°C. Trichrome staining was done using a Masson's trichrome kit from Pol Scientific (R&B) (Bayside, NY).

Tumor grading. Tumor grading was done without knowledge of genotype on the level 3 section of each lung. Each tumor was given a score of 1 to 5 based on predetermined criteria (see Results for specific grading criteria).

Tumor size measurements. Lung and tumor areas were determined using Bioquant Image Analysis software in manual measurement mode.

Results

Generation of K-ras/p53 compound conditional mutant mice. To assess the effect of p53 mutation or loss on K-ras-induced lung tumorigenesis, compound mutant mice were generated that harbored the conditional activatable LSL-KrasG12D allele (6, 18) and combinations of three different conditional p53 alleles: p53Fl/LSL-R270H, p53Fl/LSL-R172H, and p53FlBox (refs. 16, 19; Fig. 1A). For clarity, the genotypes of the compound mutant mice will be described in this article using the following abbreviations: the conditional LSL-KrasG12D allele will be referred to as K and the wild-type p53, p53Fl/LSL-R270H, p53Fl/LSL-R172H, and p53FlBox alleles will be referred to as +, 270, 172, and Fl, respectively. The compound mutant mice will be referred to collectively as K,P mice. The following K,P mice were generated: KFl/+; K270/+; K172/+; KFl/Fl;
K270/Fl, and K172/Fl, allowing us to examine the effects of point-mutant p53 proteins in the presence and absence of wild-type p53 compared with the complete loss of p53. Crosses were designed to minimize effects of mixed genetic backgrounds (see Materials and Methods). Cohorts of Kp mice were infected with 5 × 10^5 pfu of a recombinant adenovirus expressing Cre recombinase (AdenoCre) by intranasal instillation and then sacrificed at various time points (described below). As expected, all of the mice developed multiple primary lung tumors following infection with AdenoCre.

Infection with AdenoCre induces recombination of the conditional alleles. To assess the recombination efficiency of each allele when multiple alleles were present, DNA was isolated from tumors dissected from the lungs of K270/Fl mice and PCR was done to detect recombination. The p53^R270H and the p53^R172H alleles only differ at the single base mutations downstream of the LSL cassette, so the recombination of these alleles was assumed to be identical. In every tumor, PCR amplification of the K-ras allele produced a product 40 bp larger than the wild-type allele due to the single loxP site remaining after Cre-mediated recombination (Fig. 1B, top). Therefore, the K-ras^[LSL] allele recombined with 100% efficiency. Similar analyses showed that the p53^[LSL] allele recombined in 90% (9 of 10) of the tumors analyzed (Fig. 1B, middle). The recombination efficiency of the p53^[box] allele was comparable, occurring in 80% (8 of 10) of tumors analyzed (Fig. 1B, bottom).

Loss of p53 function promotes malignant progression of lung tumors. To investigate the effects of p53 loss, we evaluated the lung tumor phenotypes of K/FI/+ and K/FI/Fl mice at various time points after AdenoCre infection. A cohort of K/FI/+ mice were euthanized at a late time point (26 weeks) after infection. The tumors present in K/FI/+ mice resembled those seen previously in L-K-ras^[G12D] single mutant mice, ranging from adenomas to early-stage adenocarcinomas (6). This indicates that the loss of a single copy of p53 does not significantly affect the progression of K-ras-initiated lung adenocarcinomas.

A cohort of K/FI/Fl mice was also generated for analysis at this time point. However, it became apparent that these mice would not survive for 26 weeks after infection, so they were sacrificed after 19 weeks. In contrast to the K/FI/+ mice, all of the K/FI/Fl animals had advanced pulmonary adenocarcinomas. Multiple tumors were present in each mouse ranging from well-differentiated tumors to advanced highly dysplastic lesions containing large sheets of dysplastic cells, abnormal mitoses, and multinucleate giant cells.

To quantify the extent of progression of K/FI/Fl tumors, we devised a grading system by which to evaluate the stage of every tumor in each mouse. Tumors were scored in a blinded manner on a scale of 1 to 5 with grade 5 indicating the most advanced tumor phenotype. The criteria for each grade are as follows: Grade 1 tumors have uniform nuclei showing no nuclear atypia. Grade 2 tumors contain cells with uniform but slightly enlarged nuclei that exhibit prominent nucleoli. Grade 3 tumors have cells with enlarged, pleomorphic nuclei showing prominent nucleoli and nuclear molding. Grade 4 tumors have very large, pleomorphic nuclei exhibiting a high degree of nuclear atypia, including abnormal mitoses and hyperchromatism, and contain multinucleate giant cells. Grade 5 tumors have all the features of grade 4 tumors and also show stromal desmoplasia surrounding nests of tumor cells (see Fig. 2 for examples).

Using this grading scheme to analyze 944 tumors from 12 K/FI/+ mice and 1161 tumors from 15 K/FI/Fl mice, we confirmed that loss of p53 resulted in a markedly more severe tumor phenotype in K-ras-initiated lung adenocarcinomas (Fig. 3A). In addition, K/FI/Fl tumors showed several characteristics of human lung tumors that were lacking in previous murine lung cancer models (20). Perhaps, most striking was the presence of tumors containing a large stromal component in which nests of tumor cells could be found growing within a field of desmoplastic stroma (Fig. 3B). Masson’s trichrome staining of these tumors showed the abundant production of collagen by the stromal fibroblasts (Fig. 3C) and immunohistochemistry confirmed the expression of smooth muscle actin by some of the stromal fibroblasts (data not shown).

A subset of K/FI/Fl tumors was highly invasive, growing into the hilus, heart, and overlying pleura (Fig. 3D). Furthermore, tumor cells were observed along the luminal surfaces of blood and lymphatic vessels within the tumor mass (data not shown) showing extravasation of the tumor cells, which is required for metastatic spread. In addition, lymph node metastases were present in over 50% (8 of 15) of K/FI/Fl mice (Fig. 3E) and a small percentage had metastases to distant organs (Fig. 3F).

Having established that p53 loss accelerates the progression of K-ras-initiated lung adenocarcinomas, we were curious how rapidly this contributed to tumor progression. Therefore, we examined the lungs of K/FI/+ and K/FI/Fl mice 6 weeks after infection with AdenoCre. The lungs of K/FI/+ mice contained lesions ranging from atypical adenomatous hyperplasia to small adenomas (Fig. 4A). The adenomas present in these mice were very small and, with rare exceptions, had regular nuclei (Fig. 4B). Atypical adenomatous hyperplasia and adenomas were also present in K/FI/Fl mice but were often larger and frequently displayed nuclear atypia (Fig. 4C). Even some of the smallest lesions had aberrant nuclei (Fig. 4D) and a few tumors even contained multinucleate giant cells (Fig. 4C). Grading of 111 tumors from 12 K/FI/+ mice and 215 tumors from K/FI/Fl mice showed a clear shift (P < 0.001) towards more malignant lesions in K/FI/Fl mice (Fig. 4A), indicating that p53 loss promotes rapid lung tumor progression.

A partial dominant-negative effect by the p53^[R270H] allele towards malignant progression. Endogenous expression of mutant p53 can result in a gain-of-function effect on tumor development in some tissues (16, 21). Therefore, we examined whether endogenous expression of point-mutant p53 in K-ras-initiated lung adenocarcinomas would result in a more severe tumor phenotype than homozygous deletion of p53. Cohorts of K270/Fl and K172/Fl mice were sacrificed 19 weeks after AdenoCre infection and their lung tumors were compared with K/FI/Fl mice described above. Surprisingly, grading of 841 tumors from 11 K270/Fl mice and 839 tumors from 10 K172/Fl mice did not reveal any differences in the distribution of tumor grades compared with K/FI/Fl mice (Fig. 5A). All three genotypes had substantially fewer low-grade and more high-grade tumors than K/FI/+ animals sacrificed at 26 weeks postinfection, and all had tumors that were highly invasive and often metastatic to the regional lymph nodes. Together, these data support the loss of wild-type p53 function as a factor in the progression of K-ras-initiated lung adenocarcinomas but do not support a gain-of-function effect by point-mutant p53 in these tumors.

The absence of a gain-of-function effect allowed us to cleanly evaluate the role of dominant-negative effects of mutant p53 in the progression of K-ras-initiated lung adenocarcinomas. Cohorts of K270/+ and K172/+ mice were sacrificed 26 weeks after AdenoCre infection and compared with K/FI/+ mice described above. Grading was done on 932 tumors from 11 K270/+ mice and 773 tumors from 10 K172/+ mice. Strikingly, whereas the distribution of tumor grades was similar between K/FI/+ and K172/+ mice, the K270/+
mice displayed a significant shift in the tumor distribution ($\chi^2, P < 0.001$) with fewer low-grade and more high-grade tumors (Fig. 5 and B). Therefore, $p53^{R270H}$, but not $p53^{R172H}$, acted as a partial dominant-negative allele in the development of high-grade lung adenocarcinoma.

To determine whether this effect also extended to the size of the tumors, we examined the tumor burden after 26 weeks of growth. The average number of tumors in K;Fl/+, K;172/+, and K;270/+ mice were similar (79 ± 21, 77 ± 24, and 85 ± 17, respectively). We measured tumor burden as the percentage of total lung occupied by tumor. Like the grading results, we found that K;Fl/+ and K;172/+ mice had similar tumor burdens with tumor occupying 24% and 27% of the total lung area, respectively. However, the tumor burden in K;270/+ mice was consistently higher, with tumor comprising 36% of the total lung area ($P = 0.024$; Fig. 5C). These findings further support a dominant-negative effect specifically conferred by the $p53^{R270H}$ allele on lung tumor growth and progression.

**Up-regulation of the Raf/mitogen-activated protein kinase pathway occurs in advanced tumors.** Studies in primary fibroblasts have linked high levels of Raf/MAPK activity to Ras-induced premature senescence, in part through activation of p19ARF, a positive regulator of p53 (22). Therefore, we reasoned that mutation of p53 might allow for up-regulation of this signaling pathway, which could account for its tumor-promoting effects. To address this question, immunohistochemistry was done on both early- and late-stage tumors from the K,P mice of all six genotypes using anti–phospho-p42/44MAPK antibodies.

At 6 weeks postinfection, no phospho-MAPK positive tumors were present in any of the animals evaluated, indicating that loss of p53 alone is not sufficient to allow up-regulation of the Raf/MAPK pathway. However, a subset of late-stage tumors contained large foci of phospho-MAPK positive cells and sometimes the entire tumors were immunoreactive. The percentage of phospho-MAPK positive tumors varied depending on the p53 genotype (Fig. 5D). Roughly half of the tumors in K;Fl/Fl, K;172/Fl, and K;270/Fl animals stained positively for phospho-MAPK (52.2%, 51.2%, and 50.6% respectively). Strikingly, the frequency of MAPK activation was identical in K;270/+ mice, occurring in 53.1% of the tumors, whereas only 13.7% of K;Fl/+ tumors were phospho-MAPK positive. Therefore, although p53 loss alone is not sufficient to enable up-regulation of the Raf/MAPK pathway, it may promote the accumulation of additional events that allow MAPK activation. Furthermore, the $p53^{R270H}$ allele dominantly interferes with the ability of wild-type p53 to restrain MAPK up-regulation.

**Figure 2.** Tumor grades in K,P compound conditional mutant mice. A, a region of advanced adenomatous hyperplasia, adjacent to normal lung (left). Hyperplastic cells are enlarged and may accumulate slightly but generally follow the underlying lung architecture. B, grade 1 lesions form a solid tumor but have regular nuclei. C, grade 2 lesions may have slightly irregular nuclei and prominent nucleoli. D, cells in grade 3 lesions exhibit pleomorphic nuclei, prominent nucleoli, and nuclear molding. E, grade 4 lesions have enlarged, pleomorphic nuclei (single arrows), aberrant mitoses (double arrows), and tumor giant cells (triple arrows). F, grade 5 lesions exhibit all the criteria of grade 4 lesions as well as nests of tumors cells surrounded by a desmoplastic stroma.
Wild type–specific loss of heterozygosity in K;270/+ tumors despite dominant-negative effects by the p53<sup>R270H</sup> allele. To determine whether the dominant-negative effect of the p53<sup>R270H</sup> allele alleviates the selective pressure to lose the wild-type allele of p53, we examined the frequency of loss of heterozygosity (LOH). Quantitative PCR analysis was done on genomic DNA isolated from tumors microdissected from the lungs of K;Fl/+ and K;270/+ mice using TaqMan MGB probes. For both genotypes, 100% of the tumors analyzed (n = 11 for K;270/+ and n = 8 for K;Fl/+ ) showed loss of the wild-type p53 allele. These data show that there is still selective pressure for full loss of p53 function despite the dominant-negative effects of the p53<sup>R270H</sup> allele. This may explain why the tumors found in K;270/+ mice, although larger and more advanced than those in K;Fl/+ mice, were not as large and advanced as those in K;Fl/Fi mice, and may help to explain the wild type–specific p53 LOH that is frequently observed in human tumors (12, 13, 23).

Mutant p53 promotes the development of sinonasal adenocarcinomas. The method of AdenoCre administration used here exposes the entire respiratory tract to the virus. Thus, infection and subsequent recombination of the conditional alleles could occur in cells of the nasal passages or the trachea. However, none of the conditional K-ras<sup>G12D</sup> single mutant mice developed tumors in the upper respiratory tract, suggesting that the expression of K-ras<sup>G12D</sup> in these cells is not sufficient to induce tumorigenesis. In the course of performing the experiments described above, we observed that K;172/Fl and K;270/Fl compound mutant mice had a higher mortality than K;Fi/Fi compound mutants, which could not be explained by differences in their lung tumor phenotype, as there was none. Histologic analysis of head sections revealed the presence of anaplastic adenocarcinomas in the nasal passages of a subset of the 19-week K;Fl/Fi, K;270/Fi, and K;172/Fl mice that seemed to arise from the mucous glands of the nasal sinuses. The tumors were extremely poorly differentiated but pan-cytokeratin immunostaining confirmed their epithelial origin (data not shown). The tumor cells displayed dramatically pleomorphic, vesiculated nuclei with prominent nucleoli (Fig. 6B). Furthermore, the tumors were destructively invasive, often penetrating through the cribriform plate into the olfactory bulb (Fig. 6C). Consistent with their increased mortality, the incidence of sinonasal adenocarcinomas was >4-fold higher in K;172/Fl (64%) and K;270/Fl (69%) mice than in p53Fi/Fl (14%) mice (Fig. 6D). The increased incidence of tumors in K;270/Fi and K;172/Fi mice was...
highly statistically significant ($\chi^2, P = 0.001$). Given the significantly higher incidence of sinonasal carcinoma on expression of point-mutant p53 compared with loss of p53 expression, these data provide strong evidence for a gain of oncogenic potential by point-mutant p53 in vivo.

Discussion

In this work, we present compound mutant mice incorporating somatically manipulatable conditional mutations in the K-ras proto-oncogene and the p53 tumor suppressor gene to allow for analysis of mutant cell behavior in the context of an otherwise normal animal. Cooperation between oncogenic ras and point-mutant p53 was one of the earliest examples of genetic interaction between two cancer genes (24). Based on the observation that both K-ras and p53 are frequently mutated in human tumors, we expect that the K,P compound mutant mice described in this work will be useful in accurately modeling a number of different types of cancer beyond the advanced pulmonary adenocarcinoma described here.

K,P mice develop advanced lung adenocarcinoma. Through the generation of K-ras$^{G12D}$;p53$^{R270H}$ compound conditional mice, we have created an improved murine lung adenocarcinoma model that faithfully recapitulates several characteristics of the human disease not commonly observed in previous models. The tumors in these mice exhibit a high degree of nuclear atypia, elicit stromal desmoplasia, and are invasive and metastatic. Although it has been shown in other mouse lung cancer models that K-ras and p53 cooperate in the promotion of tumor growth and progression, the induction of a desmoplastic response and the development of metastatic disease were not reported (4, 5).

Human pulmonary adenocarcinomas often contain a large stromal component. The microenvironment in which a tumor resides may play a critical role in tumor growth and progression, and interactions between cells in the tumor and the microenvironment have been shown to elicit changes in cell behavior (25, 26). The induction of desmoplastic stroma is commonly seen in non–small-cell lung cancer and may contribute to disease progression. The production by stromal fibroblasts of hyaluronan and various matrix metalloproteinases involved in reorganization of the extracellular matrix are thought to influence the invasive and metastatic properties of the tumor (25). Correlative studies suggest that stromal productions of the cell adhesion molecule hyaluronan and of thymidine phosphorylase, which has angiogenic activity, correlate with poor prognosis (27, 28).

Human non–small-cell lung cancer is highly metastatic. Patients diagnosed with early-stage disease that has not spread to lymph nodes or distant organs have an average 5-year survival of about 50%. However, only 15% of lung cancer patients are diagnosed at this stage; most are diagnosed with metastatic disease and have an overall 5-year survival rate only 14%. The current therapeutic regimens are not effective in the treatment of advanced non–small-cell lung cancer. Many murine lung cancer models have been created that provide valuable insights into the molecular alterations driving lung tumorigenesis. However, because these models do not generally develop advanced disease, their growth properties and response to therapeutics may not accurately reflect those of human tumors. Thus, the model described here will provide a valuable system for the study of lung tumor biology and the preclinical testing of novel therapeutics.

The contact mutation p53$^{R270H}$ contributes more strongly to lung tumorigenesis than the structural mutation p53$^{R172H}$. Although it is generally accepted that p53 mutation is an important event in the development of lung cancer, the relative contribution of different p53 mutations has not been previously analyzed in an in vivo setting. By comparing the effects of different p53 alleles under identical conditions, we have established that loss of p53 function is sufficient to promote the progression of K-ras initiated lung adenocarcinomas in mice and that the p53$^{R270H}$ allele more
strongly contributes to lung tumorigenesis by acting as a partial dominant-negative. The primary functions of p53 are to respond to cellular stresses by transactivating genes involved in cell cycle arrest, apoptosis, and genome maintenance. Although we could not determine the contribution of specific p53 effector pathways, it is unlikely that the apoptotic functions of p53 are significant in this setting because we could not detect any differences in the rates of apoptosis within lung tumors of K,P mice by immunostaining for cleaved caspase-3 (data not shown).

Instead, it seems likely that loss of the genome maintenance and antiproliferative functions of p53 are critical to the development of advanced lung adenocarcinoma. One of the most prominent features of the tumors in K,Fl/Fl mice is the development of severe nuclear dysplasia. The observation of lesions with prominent nuclear atypia just 6 weeks after infection with AdenoCre indicates a rapid accumulation of genetic abnormalities following loss of p53. Our finding that early-stage K,Fl/Fl, K,270/Fl, and K,172/Fl tumors did not up-regulate the MAPK pathway whereas late-stage tumors did is consistent with this hypothesis. The data show that p53 mutation is not sufficient for up-regulation of this pathway but facilitates the acquisition of additional events leading to increased signaling through the Raf/MAPK pathway.

Loss of p53 seems to affect the progression of lung adenocarcinomas rather than their initiation. Lung tumors are extremely rare in p53<sup>−/−</sup> mice, presumably due to the early onset of lymphomas and sarcomas. A small subset of p53<sup>−/−</sup> mice does develop lung adenocarcinomas but the tumors are generally solitary and have a long latency (>18 months). Likewise, p53<sup>F1/F1</sup> mice treated with AdenoCre also develop lung adenocarcinomas but only 1 to 1.5 years after treatment (29). Therefore, p53 mutation alone is not sufficient for the initiation of lung cancer. Instead, activation of K-ras serves to initiate the tumor and p53 loss contributes to the progression.

Although p53 loss of function is sufficient to drive tumor progression in K-ras-induced lung adenocarcinomas, p53 is rarely deleted in human lung tumors. Rather, p53 is commonly point-mutated at specific hotspot residues in the DNA-binding domain in lung and other tumors. In particular, contact mutations at codons 248 and 273 are very common in human lung adenocarcinomas. In contrast, an analysis of spontaneous tumors from the IARC p53 mutation database (version R9) confirms that codon 175 mutations are substantially underrepresented in adenocarcinomas and non–small-cell lung cancers compared with all tumors (χ², P = 0.000026). Previously, explanations for this variation focused on the mutagenic effects of cigarette smoke on different p53 codons, especially in light of in vitro evidence showing that codon 175 mutations are more potent in oncogenic assays. However, we found that the p53<sup>R270H</sup> mutation is more strongly dominant-negative over wild-type p53 than is the p53<sup>R172H</sup> mutation and therefore more effective in tumor promotion as shown by the increased tumor burden and higher proportion of high-grade tumors in K,270/+ mice.

There are many possible explanations for this effect. For example, p53<sup>R270H</sup> may have a stronger intrinsic capacity to interfere with wild-type p53 than p53<sup>R172H</sup>. Alternatively, p53<sup>R270H</sup> may accumulate to higher levels in tumor cells than p53<sup>R172H</sup>. Consistently, we found that a higher proportion of K,270/+ tumors showed strong immunoreactivity for p53 (data not shown). One final possibility is that K,270/+ tumors undergo LOH at an earlier point than K,172/+ tumors, perhaps due to a specific dominant-negative effect by p53<sup>R270H</sup> on the genome maintenance functions of p53. Once again, our data on tumor grade and MAPK up-regulation are consistent with this hypothesis as discussed above.

**Gain-of-function effects of point-mutant p53 in sinonasal adenocarcinoma.** In addition to non–small-cell lung cancer, some of the K,P mice developed anaplastic adenocarcinoma of the nasal mucosa arising from the mucous gland epithelium. A significantly
A higher proportion of both K;270/Fl and K;172/Fl mice developed sinonasal adenocarcinomas (69% and 64%, respectively) compared with K;Fl/Fl mice (14%). Because the lung tumor phenotype did not vary between K;Fl/Fl and K;270/Fl or K;172/Fl, the discrepancy in sinonasal adenocarcinoma incidence strongly suggests a tissue-specific gain-of-function of point-mutant p53. Also intriguing is the fact that the relative effects of the p53R270H and the p53R172H alleles are similar in the mucous gland epithelium but appreciably different in the lung epithelium, also pointing to the tissue specificity of the effects of individual p53 mutations. Given the prevalence of p53 mutations in human cancer, elucidating the different allele- and tissue-specific effects of mutant p53 is critical to developing a more complete understanding of molecular carcinogenesis.

Sinonasal adenocarcinomas are rare in humans and the tumors seen in our mice are not readily diagnosed based on human classification criteria. However, we believe they arise from the epithelium of the mucous glands and, thus, are most similar to minor salivary gland tumors in humans. The majority of over 500 minor salivary glands in the head and neck are mucous glands (http://www.bcm.edu/oto/grand/7992.html). Minor salivary gland tumors in humans account for only 10% to 15% of all salivary gland tumors but, unlike tumors of the major salivary glands, which are mostly benign, ≥80% of minor salivary gland tumors are malignant (30, 31). Interestingly, although p53 alterations are rare when evaluated in salivary gland carcinomas as a whole, their frequency may be much higher in carcinomas of the minor salivary glands. An analysis of adenoid cystic and mucoepidermoid carcinomas of the salivary glands revealed p53 alterations in 17.6% and 14.8% of the tumors, respectively, but all of these were tumors arising from the minor salivary glands (32). Therefore, p53 mutation may play a critical role in the generation of minor salivary gland tumors.

Surgery is the primary mode of treatment for minor salivary gland tumors, followed by radiation therapy in some cases (31). Unfortunately, due to the rarity, multiplicity of sites of origin, and varied histology of the disease, no large, multi-institutional, prospective, randomized trials have been completed. As such, the current treatment strategies might not be optimal. We believe this model may provide a valuable tool for the study of minor salivary gland carcinoma biology and could aid in the development of better treatment strategies.

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