Elimination of p19ARF-expressing cells enhances pulmonary function in mice

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Introduction

Most mammalian somatic cells have infinite replicative life spans and eventually undergo permanent cell cycle arrest, called cellular senescence (1). Senescence is triggered by sustained and irreparable damage that leads to the activation of tumor suppressor pathways. Two major tumor suppressor pathways, namely, the p19ARF (p14ARF in humans)/p53 and p16INK4a/Rb pathways, play critical roles in the induction and maintenance of cell cycle arrest during cellular senescence (2, 3). The inactivation of these pathways bypasses senescence, thereby allowing the propagation of damaged cells, which eventually leads to cancer. Thus, cellular senescence eliminates potential malignant transformation and functions as an essential tumor-suppressive mechanism in mammals.

While there is no doubt that senescence prevents cancer, an increasing amount of evidence suggests that cellular senescence is involved in other biological processes and pathologies. Cellular senescence has been shown to contribute to embryonic development (4, 5), wound healing (6), and tissue regeneration (7). Additionally, it has become more evident that cellular senescence contributes to tissue aging. Senescent cells accumulate in many tissues during aging (8) and are considered to underlie aging-associated pathologies (9). The contribution of senescent cells in aging-associated phenotypes may depend on their non–cell-autonomous functions, such as senescence-associated secretory phenotype (SASP), because the population of senescent cells is very small, even in very old human tissue (10).

Senescence is enhanced in BubR1 mutant mice that show accelerated aging phenotypes (11), and the lifelong elimination of p16INK4a-expressing senescent cells from these mice was found to partially reverse these phenotypes (11). The ablation of cellular senescence by deleting the INK4a gene also restored some progeria-like phenotypes and extended the life span of Klotho mutant mice (12). Collectively, these findings indicate that cellular senescence is responsible for aging-associated phenotypes but may not account for all phenomena in aged animals. It has not yet been established whether cellular senescence contributes to chronological (naturally occurring) aging phenotypes and, more importantly, if aging-associated pheno-
types may be reversed by eliminating senescent cells from old animals. van Deursen and colleagues recently reported that the clearance of senescent cells in old animals extended the life span of the mouse using an INK-ATTAC transgenic model (13). They concluded that age-dependent changes, at least in the kidney, heart, and adipose tissue, are caused by p16\textsuperscript{INK4a}-expressing senescent cells, which strongly influence the life span of this animal.

In the present study, we established a transgenic model from which it was possible to eliminate p19\textsuperscript{ARF}-expressing cells using a toxin-mediated cell knockout system (14, 15). Similar to INK4a, the expression of ARF has been shown to increase during aging in the mouse (8). Using the transgenic model, we successfully eliminated ARF-expressing cells from the lung tissue of 12-month-old animals. The ablation of ARF expression abolished the expression of other senescent markers, including INK4a and p21, suggesting that the expression of ARF reflects the accumulation of senescent cells in tissues. The elimination of p19\textsuperscript{ARF}-expressing cells in lung tissue ameliorates the aging-associated loss of tissue elasticity. Moreover, the expression of a large number of aging-associated genes was reversed after the removal of p19\textsuperscript{ARF}-expressing cells. Taken together, these findings highlight the role of p19\textsuperscript{ARF} in lung tissue aging and indicate that the aging phenotype in lung tissue may be reversed by eliminating p19\textsuperscript{ARF}-expressing cells from tissue.

**Results**

In order to investigate the role of p19\textsuperscript{ARF} in tissue aging, we established a transgenic mouse (ARF–diphtheria toxin receptor [ARF-DTR]) in which the DTR (human HB-EGF I117V/L148V; ref. 14) and luciferase were expressed from the CDKN2A locus (Figure 1A). Embryonic fibroblasts prepared from these mice (MEFs) expressed p19\textsuperscript{ARF}, p16\textsuperscript{INK4a}, p21, and luciferase when senescence was induced by either serial passages or oncogenic Ras (Figure 1, B and C). The expression of senescence markers (p19\textsuperscript{ARF} and p16\textsuperscript{INK4a}) as well as luciferase was diminished when senescent ARF-DTR MEFs were treated with diphtheria toxin (DT) (Figure 1, D–G). An in vivo imaging analysis revealed that luciferase activity was barely detectable in young (2-month-old) female mice but became apparent in multiple tissues by the age of 12 months (Figure 2A). Similar results were obtained in male mice, with the exception of the testis (data not shown); luciferase was detectable in the testis irrespective of the age of animals, which presumably reflected the lifelong expression of p19\textsuperscript{ARF} in mouse spermatogonia (16). The luminescence signals observed in old female mice were attributed to those in the lungs and adipose tissues (Figure 2, C and D), which is consistent with the endogenous expression of ARF in the lungs and adipose tissues at this age (Figure 2B). INK4a mRNA expression was also increased, as previously reported (8), confirming the accumulation of senescent cells in these tissues. However, expression of ARF is increased in many tissues during aging (8), suggesting that the expression of the transgene in ARF-DTR mice does not mimic the expression pattern of endogenous ARF in other tissues.

The intraperitoneal administration of DT eliminated the luminescence signal from the lungs within 48 hours (Figure 2, C and D), and this effect lasted for at least 2 weeks after drug administration (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/jci.insight.87732DS1). In contrast, the DT treatment failed to reproducibly decrease luciferase activity in adipose tissue for an unknown reason. Based on the above results, we analyzed the role of p19\textsuperscript{ARF}-expressing cells in lung function using female ARF-DTR mice.

The aging-associated decline in lung function is characterized by an increase in tissue compliance due to the progressive loss of tissue elasticity, which results in the incomplete contraction of lung tissue during the expiration phase (17–19). Luciferase was not detected in bronchoalveolar lavage fluid cells but was detected in the fraction mainly containing fibroblasts (Figure 3, A and B). Immunohistochemistry revealed that endogenous p19\textsuperscript{ARF} colocalized with luciferase and the senescence marker γH2AX in 12-month-old lung tissue (Figure 3, D and E). Luciferase was observed in cells expressing fibroblast markers (vimentin and ER-TR7) but not in those expressing the alveolar epithelial marker, SP-C (Figure 3, C, F, and G). Furthermore, when lung cells were sorted according to epithelial (CD31\textsuperscript{−}; CD45\textsuperscript{−}; EpCAM\textsuperscript{−}) and mesenchymal (CD31\textsuperscript{+}; CD45\textsuperscript{−}; Sca-1\textsuperscript{+}) markers, luciferase was detected in the mesenchymal population, suggesting that p19\textsuperscript{ARF}-expressing cells in the adult mouse lung are alveolar fibroblasts (Figure 3, H and I). Taken together, these results strongly indicate that cellular senescence is induced in the alveolar fibroblasts of the 12-month-old lung parenchyma.

Twelve-month-old mice were treated with PBS or DT (Figure 4A). Lung luciferase activity in
12-month-old mice decreased to a similar level as that in young mice after the DT treatment (Figure 4B). No significant changes were observed in body weights after the DT treatment (Supplemental Figure 2). The DT treatment reduced ARF, INK4a, and p21 levels in the lungs of ARF-DTR mice, but not in those of wild-type mice, indicating that p19ARF-expressing senescent cells were successfully removed from lung tissue of ARF-DTR mice (Figure 4, C and D).

We then performed pulmonary function tests on these mice. As previously reported (20, 21), static lung compliance (Cst) was significantly higher in older animals than in young animals (Figure 5, A and B). The DT treatment resulted in the marked recovery of lung elasticity (decrease in Cst) in ARF-DTR mice but not in wild-type mice. Similarly, the DT treatment reversed aging-associated changes in dynamic compliance, dynamic resistance, tissue elastance, and tissue damping in ARF-DTR mice but not in wild-type mice (Figure 5, C–F). These results clearly indicated that the p19ARF-expressing cells that accumulated in 12-month-old lung tissue had deleterious effects on pulmonary function and that aging-associated declines in pulmonary function were ameliorated by the elimination of these p19ARF-expressing cells.

We also compared lung morphologies among these mice. An increase in alveolar size is associated with a decline in lung function during aging (18, 19, 22, 23). Alveolar mean linear intercepts (the average distance between alveolar walls that reflects the size of the alveolar space; ref. 24) were increased in 12-month-old ARF-DTR mice (Figure 6, A and B). The DT treatment reversed this phenotype but did not affect alveolar size in wild-type mice. Mean alveolar septal wall thickness was lower in 12-month-old mice than in 2-month-old mice (Figure 6C), as previously reported (25). Furthermore, the ablation of p19ARF-expressing cells fully restored the alveolar wall in ARF-DTR mice but not in wild-type mice. Increases in lung compliance in aged animals have been attributed to a decline in the number of elastic fibers, the major component of which is elastin (19, 22). Therefore, we performed a quantitative analysis on elastin in immunostained sections (Figure 6D). The amount of elastin in older animals was approximately half that in young animals (Figure 6E) and recovered in DT-treated ARF-DTR mice but not in wild-type mice.
We also examined the effects of ARF-expressing cell elimination on even older animals. Tumor-free female ARF-DTR or wild-type mice between 20 and 22 months old were treated with PBS or DT for 1 month (Figure 7A). In vivo imaging analysis confirmed that luciferase signal disappeared from lung tissue (Figure 7B). Pulmonary function tests revealed that tissue compliance in older mice was similar to that in 12-month-old mice (Figure 7, C, D, and F). The DT treatment reduced tissue compliance in older animals; however, this effect was less than that observed in 12-month-old mice. Similarly, the DT treatment reversed aging-associated decreases in dynamic resistance and tissue damping (Figure 7, E and G). Alveolar size, which was larger than that in 12-month-old mice, was slightly restored in DT-treated animals (Figure 7, H and I). Collectively, these results indicated that p19ARF-expressing cells provoked the loss of elastic fibers in lung tissue and were also responsible for the increase in lung compliance in aged animals.

Senescent cells express cell cycle–regulating genes as well as a series of genes that function non–cell autonomously, such as cytokines, thereby affecting the behaviors of their surrounding “nonescent” cells (26, 27). In order to gain an insight into the relationship between p19ARF-expressing cells and aging-associated phenotypes in lungs, we examined the expression of senescence-associated genes in these animals (Figure 8A). Among the MMPs examined, MMP-10 and -12 were significantly increased in older mice and were completely abolished or diminished in DT-treated ARF-DTR mice but not in wild-type animals (Supplemental Figure 3). These MMPs have been shown to exhibit elastase activities (28, 29)
and are, thus, good candidates for the reduced level of elastin observed in the lungs of aged animals. MMP-12 knockout mice were found to resist the degradation of elastin when emphysema was induced by cigarette smoking (30). MMP-10 protein was also downregulated in DT-treated ARF-DTR lungs but not to the same extent as their mRNA levels (Supplemental Figure 4). In order to ascertain whether increases in MMP were responsible for the reduced elastin level, MEFs were infected with retroviruses encoding shRNA against MMP-10. Cells were then subjected to ionizing radiation and cultured under 3% oxygen to induce senescence-associated genes in MEFs (31). MMP-10 level was decreased to approximately half by shRNA (Figure 8B). Elastin levels were reduced to approximately half that of untreated cells cultured under 20% oxygen (data not shown) but recovered when MMP activities were inhibited either by shRNA or a chemical inhibitor (Figure 8C). Taken together, these results suggest that the loss of elastin in senescent cells was, at least partly, attributed to increased MMP.

We also compared gene expression profiles among 2-month-old and 12-month-old (with or without DT) ARF-DTR lungs. Genes differentially expressed (more than 2-fold) between 2-month-old and 12-month-old lungs were identified (Supplemental Table 1, a full list of genes is available from the GEO database, accession GSE64754; http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE64754). Changes in the expression of these genes among young and old DT-treated animals were then analyzed. In order to simplify the results obtained, genes with expression levels that changed more than 2-fold in wild-type animals after the DT treatment were subtracted from the list because they exhibited “nonspecific” responses to DT.

**Figure 3. Luciferase was expressed in alveolar fibroblasts in old ARF-DTR mice.** (A) Lung cell fractionation. Bronchoalveolar lavage (BAL) fluid mainly containing macrophages was initially obtained from the lung. The rest of the tissue was minced, trypsinized, and seeded on a tissue culture dish. After being incubated for 1 hour, floating cells that contained epithelial cells were collected. Attached cells were trypsinized to recover alveolar fibroblasts. (B) Lysates were prepared from the collected cells, and the luciferase assay was performed. Luciferase activity was normalized to the cell number in each sample. Results of 2 independent experiments are shown. (C-E) Immunofluorescence staining of the ARF–diphtheria toxin receptor (ARF-DTR) lung. Sections were stained with antibodies against the indicated proteins. Original magnification, ×40 (C and D), ×100 (E); scale bar: 20 μm (C and D), 10 μm (E). (F) Immunofluorescence staining of the ARF-DTR lung. Sections were costained with surfactant-associated protein C (SP-C) or the ER-TR7 fibroblast marker together with luciferase. Sections were counterstained with DAPI. Original magnification, ×100; scale bar: 10 μm. (G) The luciferase-positive population in SP-C- or ER-TR7–stained cells was counted. At least 100 SP-C– or ER-TR7–positive cells were analyzed in each section. Values represent the mean ± SEM of 3 independent experiments. N.D., not detected. (H) Gating strategy for CD31–; CD45–; EpCAM+ epithelial cells and CD31–; CD45–; Sca-1+ mesenchymal cells. Lung cells from 7 ARF-DTR mice were pooled before sorting. (I) Luciferase activity was determined in gated epithelial and mesenchymal cells shown in (H). Luciferase activity was normalized to cell numbers in each sample. A representative result of 2 independent experiments is shown. Values represent the mean ± SD of triplicate samples. *P < 0.05, unpaired Student’s t test.
A pathway analysis revealed that genes involved in cytokine and chemokine signaling are differentially expressed in 12-month-old lungs, which may reflect the secretory phenotype of senescent cells (26, 27) (Table 1). Of the 305 genes that were upregulated in old lungs, the expression of 170 (55.7%) genes was decreased following the DT treatment (Figure 9 and Supplemental Table 1). Conversely, the DT treatment upregulated the expression of 82 (62.1%) of the 132 downregulated genes in old lungs. Thus, the elimination of p19 ARF-expressing cells led to the “rejuvenation” of gene expression profiles in whole lung tissue; however, approximately 40% of genes with altered expression levels in 12-month-old animals remained intact under these conditions, suggesting that p19ARF contributed to only a certain aspect of the aging-associated phenotypes of the lung or that some aging-associated changes are irreversible.

Discussion

We established a mouse model in which it was possible to eliminate p19ARF-expressing cells using a TRECK system. By using this model, the p19ARF-expressing cells that accumulated during aging were successfully eliminated from lung tissue. Attempts to eliminate senescent cells in vivo have been made by Campisi’s and van Deursen’s groups (6, 13). The former showed that cellular senescence is induced during wound healing and promotes wound closure by enhancing myofibroblast differentiation through SASP. The latter documented the role of senescence in naturally occurring tissue aging, including the kidney and heart, and, more importantly, showed that senescent cells limited the life span of model mice. So far, we obtained only one transgenic mouse line in which ARF-DTR transgene exhibited the senescence-dependent expression. The transgene expression reflected endogenous ARF expression in at least lung and adipose tissues but not in other tissue such as liver in which ARF is induced during aging (8), precluding the analysis of the roles of p19ARF-expressing cells in other tissues or life spans. Nonetheless, our results clearly demonstrated that the elimination of p19ARF-expressing cells from lung tissue ameliorated aging-associated phenotypes, including reduced tissue elasticity, morphological changes, and gene expression. Therefore, aging-associated pulmonary hypofunction was caused, at least partly, by p19ARF, which accumulates during aging.

In our mouse model, DTR and luciferase are expressed under the control of ARF promoter/enhancers. Unlike in human cells, ARF clearly plays essential roles in the induction of cellular senescence in mouse
cells. p16\textsuperscript{INK4a} and p19\textsuperscript{ARF} are both induced during senescence in mouse cells. Nevertheless, INK4a is largely dispensable for senescence in MEFs (32, 33), while ARF-null MEFs do not have a replicative life span and knockout mice are highly prone to cancer (34). Although we were unable to detect senescence-associated β-galactosidase activity in old lung tissue sections, p19 ARF-expressing cells were positive for γH2AX, and the elimination of these cells also blunted the expression of other senescent markers, including INK4a and p21. Hence, our results strongly suggest that p19ARF-expressing cells in lung tissue represent senescent cells. However, it should be noted that ARF is less important or dispensable for the senescence of human cells, since p14ARF is not induced during senescence. Expression of ARF is more limited in humans, and hyper-
physiological activation of the E2F transcription factor is required for its induction (35, 36). Therefore, it is unlikely that p14ARF exhibits aging/senescence-dependent expression in human tissue.

We observed luciferase and p19ARF expression in alveolar fibroblasts. Fibroblasts are thought to be a major source of extracellular matrix and play a crucial role in the maintenance of tissue structure. Thus, changes in fibroblast function caused by p19ARF expression may directly cause a pulmonary dysfunction. However, this is unlikely, because only a small fraction of fibroblasts expressed luciferase and p19ARF. Additionally, it cannot be formally excluded that an extremely small number of epithelial or other cells also express p19ARF and cause pulmonary aging. Further studies are required to determine whether other cells also contribute to the phenotypes observed in 12-month-old mice.

Senescent cells are known to have an effect on their surrounding “nonsenescent” cells through SASP (27). Our results suggest that the population of p19ARF-expressing cells was very small, even in adult lung tissue (approximately 1% of the lung mesenchymal population). Nevertheless, our microarray data indicate that these “rare” p19ARF-expressing cells strongly influence gene expression in lung tissue. Hundreds of genes are upregulated and downregulated during aging in the lung, and more than half of these aging-associated genes show p19ARF dependence. Since senescent cells non–cell autonomously induce senescence-like gene expression in their surrounding cells through SASP (37), it is reasonable to assume that changes in aging-associated genes in lung tissue do not simply reflect the events within p19ARF-expressing cells, but also include global changes in lung tissue cells that are mostly nonsenescent. A pathway analysis revealed that cytokine and chemokine signaling pathways are differentially regulated in 2-month-old and 12-month-old lungs, which is expected since most SASP genes participate in these signaling pathways (38). In addition to cytokine and chemokine signaling pathways, our results suggest that genes involved in circadian rhythms are also differentially regulated in older lung tissues. Previous studies demonstrated that the circadian clock affected pulmonary inflammation (39, 40). Thus, aging may affect lung pathology by modulating circadian clock-dependent gene expression.

Aging is considered to underlie many diseases in humans, and this is also the case in the lungs because the risk of chronic obstructive pulmonary disease (COPD) increases with age and is currently one of the main causes of death worldwide. Although aging-associated changes in lung tissue may not directly cause pulmonary diseases, aging facilitates the development of cigarette smoke–induced inflammation and emphysema (41, 42). Cigarette smoking has been shown to markedly increase the incidence of COPD in humans (43). The number of senescent cells also increases under these circumstances (44, 45). Hence, these findings, taken together with our results, suggest a possibility that emphysema is prevented by suppressing cellular senescence. Furthermore, senescent cells accumulate not only in the lungs, but also in other tissues, and are considered to be responsible for several aging-associated phenotypes and diseases (9, 13, 46, 47). Future studies will clarify the role of senescent cells in the aging of other organs and the availability of a senescent cell–targeted therapeutic approach, such as senolytic agents (48–50), for aging-associated diseases.

<table>
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Two- and twelve-month-old female ARF-DTR mouse lungs were analyzed by microarrays (Figure 9 and Supplemental Table 1), and a pathway analysis of differentially expressed genes in the lungs of 12-month-old mice was performed using KEGG gene sets. Significantly enriched gene sets (FDR-adjusted P < 0.05) are listed.
Methods

**PAC transgenic construct.** A mouse ARF/INK4a/INK4b PAC clone was provided by Gordon Peters (Cancer Research UK, London, United Kingdom). The PAC clone contained the complete 19 kb of the mouse ARF genomic sequence, with an additional 25 kb of the 5′- and 33 kb of the 3′-flanking genomic regions of the mouse ARF gene, respectively (Figure 1A). A chimeric ARF-DTR-2A-luc PAC transgenic construct that contains a tandem cassette of the human heparin-binding epidermal growth factor–like growth factor (HB-EGF) with the DTR-generating I117V/L148V mutation (14), the 2A peptide gene, and a luciferase reporter gene was derived from the pGL3 Basic vector (Promega) in place of the start codon of the ARF gene locus, was generated by recombineering (Figure 1A). The DTR-2A-luc gene was transferred to the ARF PAC clone using the Red/ET Counter Selection BAC Modification Kit (Gene Bridges) (51). In brief, a rpsL-neo counter selection cassette flanked by two adjacent sequences of the start codon of the ARF gene was amplified by PCR. The amplified rpsL-neo counter selection cassette was inserted into the ARF gene of the PAC clone by Red/ET recombination. The DTR-2A-luc gene was subcloned to plasmid pENTR1A (Thermo Fisher Scientific). The subcloned DTR-2A-luc fragment was flanked by 30 nucleotide sequences of the ARF gene untranslated region and the exon sequence directly adjacent to the start codon. The DTR-2A-luc gene fragment was precisely transferred to the ARF gene of the mouse PAC clone by Red/ET recombination in order to construct a chimeric ARF-DTR-2A-luc PAC clone. The PAC modification was verified by sequencing. The ARF-DTR-2A-luc PAC transgenic construct was purified for a microinjection with a slightly modified procedure (52). The PAC transgenic construct was extracted from 250 ml of the E. coli culture using the Nucleobond Plasmid Purification kit (MACHEREY-NAGEL). In order to achieve purification, 10 μg of the PAC transgenic construct was linearized overnight with SfiI (New England Bio-
Linearized PAC DNA was separated with pulsed field gel electrophoresis (PFGE) and extracted from the preparative pulsed field gel by electroelution. After dialysis against TE buffer containing 0.1 mM EDTA, aliquots were subjected to PFGE for size and quality control. The PAC DNA concentration was adjusted to 1 ng/μl for the microinjection. Aliquots of the PAC DNA solution were stored at 4°C for later microinjection.

**ARF-DTR transgenic mice.** ARF-DTR mice were generated by a pronuclear injection of C57BL/6J mouse...
embryos (CLEA Japan Inc.). The transgenic founders of the PAC transgenic construct were assessed by Southern blotting of HincI-digested tail DNA probed by the [32P]-labeled luciferase gene fragment. Fifteen of eighty-three progeny (18%) contained the transgene, as detected by the Southern blotting of tail DNA. No obvious gross phenotypic differences were apparent when transgene-positive and -negative littermates were compared.

Fifteen independent transgenic founder mice were bred with C57BL/6J mice. Genomic DNA isolated from each founder mouse was analyzed by Southern blotting, and 9 founder mice carried more than 1 copy of transgene per genome. Embryonic fibroblasts were prepared from these mice, and luciferase activity was analyzed after the induction of senescence. One transgenic line that carried 10 copies of transgene per genome showed senescence-dependent luciferase expression, which was used for all analyses in this study. Regarding the DT treatment, mice were intraperitoneally injected with DT (50 μg/kg bodyweight) twice with a 2-week interval.

Mice developing tumors were excluded from all analyses. Animals were randomly assigned to groups.

**Cell culture, luciferase assay, and in vitro elastin assay.** MEFs were cultured using a 3T3 protocol and infected with a retrovirus as previously described (53). Oncogenic Ras (V12) retrovirus plasmid was provided by Manuel Serrano (Spanish National Cancer Research Center, Madrid, Spain). Infected cells were selected with puromycin. The luciferase assay was performed using the Luciferase Assay System (Promega). Luciferase activity was normalized to the cell number in each sample.

In order to quantitate the amount of elastin, early-passage MEFs were infected with a retrovirus expressing shRNA targeting MMP-10 or scramble shRNA. The target sequences of sh-MMP-10 were ATGGAAAC-CAGGTGCTAGA (sh-MMP-10-1) and GCGCAGCAGATGTCTTTCA (sh-MMP-10-2). Infected cells were cultured for 10 days under 3% oxygen conditions following exposure to ionizing radiation (10 Gy) in order

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**Figure 8.** Changes in senescence-associated gene expression in DT-treated lungs. (A) Total RNA was extracted from 2- and 12-month-old female ARF–diphtheria toxin receptor (ARF–DTR) mouse lung tissues. Twelve-month-old mice were treated with PBS or diphtheria toxin (DT), as depicted in Figure 4A. The expression of the indicated genes was analyzed by real-time PCR. mRNA levels were normalized to GAPDH mRNA in each sample. Values represent the mean ± SEM. (B) Wild-type MEFs were infected with retroviruses encoding shRNA against MMP-10. Two-independent shRNA were used to knockdown MMP-10. Scrambled short hairpin (sh-SCR) was used as a control in each group. Infected cells were selected with puromycin for 3 days, exposed to ionizing radiation (10 Gy), and cultured under 3% oxygen conditions for 10 days. The expression of MMP-10 was analyzed by immunoblotting. Lamin A/C was used as a loading control. (C) shRNA-infected senescent MEFs were analyzed for the amount of elastin. In the chemical inhibition of MMPs, senescent MEFs were cultured in the presence of an MMP inhibitor [N-isobutyl-N-(4-methoxyphenylsulfonyl)-glycylhydroxamic acid (NNGH)]. Media were replaced every 3 days. The amount of elastin was measured and then normalized to the total protein content in each sample. Values represent the mean ± SD of triplicate samples. Data are representative of 2 independent experiments. Data were analyzed 1-way ANOVA followed by post-hoc Tukey-Kramer multiple comparison test (for comparison of 3 groups) or unpaired Student’s t test. *P < 0.05, **P < 0.01, ***P < 0.001.
to induce SASP (31). Where indicated, 50 μM of N-isobutyl-N-(4-methoxyphenylsulfonyl)-glycylhydroxamic acid (NNGH) was added to the culture medium. Media were replaced with fresh media every 3 days. The amount of elastin was determined using the Fastin Elastin Assay kit (Biocolor) and normalized to total protein.

In vivo imaging analysis. An in vivo luciferase imaging analysis was performed using the IVIS imaging system (Perkin Elmer). Mice were ventrally shaved, anesthetized with isoflurane (Wako Pure Chemicals Industries), and intraperitoneally injected with luciferin according to the manufacturer's instructions (VivoGlo; Promega). Luciferase activity was monitored 10 minutes after the luciferin injection. In order to quantify luciferase activity, signals were analyzed using Living Image software (Perkin Elmer).

Pulmonary function test. Pulmonary function tests were performed using the FlexiVent system (Scireq) as previously described (54, 55). Mice were euthanized by an intraperitoneal injection of pentobarbital sodium (100 mg/kg of body weight) and connected to the FlexiVent system after tracheotomy. Mice were ventilated at a respiratory rate of 150 breaths per minute, with a tidal volume of 10 ml/kg against a positive end expiratory pressure of 3 cmH2O. Deep inflation, Snapshot-150, Quickprime-3, and a pressure-volume loop with constant increasing pressure were consecutively performed 3 times in each subject. Cst values were calculated by fitting the Salazar-Knowles equation to the pressure-volume loop. Dynamic lung compliance and resistance values were calculated using a single frequency forced oscillation technique. Pressure, flow, and volume signals obtained from the response to a sinusoidal waveform were fit to the single compartment model using linear regression. Tissue elastance and damping, which are closely related to tissue resistance and elastance, respectively, were obtained from respiratory system impedance data using a constant phase model. All parameters were calculated using FlexiVent software.

Lung morphometry and immunohistochemistry. All histopathological analyses were performed in a blinded manner. Lungs were fixed with Bouin solution or Mildform 20 N (Wako Pure Chemicals Industries) at 25 cmH2O. Paraffin-embedded tissues were sectioned (5 μm thick), stained with Maeda's Resorcin-Fuchsin solution (Muto Pure Chemicals), and counterstained with hematoxylin-eosin. At least 8 randomly selected fields per mouse were photographed, and the mean linear intercepts and alveolar wall thicknesses were calculated. In the measurement of mean linear intercepts, test lines were randomly drawn on images, and the length between two sequential intersections of the alveolar surface with the test line was measured. Airway and vascular structures were eliminated from the analysis.

Immunostaining was carried out after antigen retrieval using LAB solution (Polysciences Inc.) with an elastin antibody (Abcam). Immunostained sections were mounted without counterstaining, and 4 fields not containing airway or vascular structures per mouse were photographed under equal conditions. Elastin signal intensities were analyzed using ImageJ (NIH).

In the immunofluorescence analysis, sections fixed with Bouin solution or formalin were immunostained using a p19ARF antibody (1:50 dilution, 5-C3-1, Santa Cruz Biotechnology), surfactant-associated protein C (1:50 dilution, SP-C, M-20, Santa Cruz Biotechnology), fibroblast marker antibody (1:50 dilution, ER-TR7, Santa Cruz Biotechnology), luciferase antibody (1:100 dilution, PM016, MBL, Japan), γH2AX antibody (1:400 dilution, ab2893, Abcam), and vimentin antibody (1:100 dilution, RV202, Abcam). Sections were visualized with Alexa Fluor 488–conjugated anti-rat IgG (1:2000 dilution, 712-545-153, Jackson ImmunoResearch), Alexa Fluor 488–conjugated anti-goat IgG (1:2000 dilution, 705-545-147, Jackson ImmunoResearch), Alexa Fluor 488–conjugated anti-mouse IgG (1:2000 dilution, 715-545-151, Jackson ImmunoResearch), and Alexa Fluor 488–conjugated anti-rabbit IgG (1:2000 dilution, 711-545-153, Jackson ImmunoResearch).
ImmunoResearch), and Alexa594-conjugated anti-rabbit IgG (1:2000 dilution, ab150080, Abcam). Sections were counterstained with DAPI.

**Immunoblotting.** Lysates were prepared using RIPA buffer (10 mM Na-phosphate, pH 7.2, 150 mM NaCl, 2 mM EDTA, 0.1% SDS, 1% Na-deoxycholate, and 1% NP-40) containing protease inhibitors. Proteins were separated by SDS-PAGE and transferred to PVDF membranes (Merck-Millipore Billerica). Proteins were detected with antibodies to p19ARF, p16INK4a, p21, Lamin A/C, and α-Tubulin (5-C3-1, M-156, F-5, H-110, and B-7; all from Santa Cruz Biotechnology); MMP-10 (bs-1344R, Bioss Inc., Boston MA); and MMP-12 (EP1261Y, Abcam). Antibodies were diluted to 1:500 (p19ARF, p16INK4a, p21, Lamin A/C, α-Tubulin, MMP-10, and MMP-12) or 1:2000 (Lamin A/C) with a blocking buffer (tris-buffered saline containing 5% skim milk and 0.1% Tween20).

**Real-time PCR analysis.** A real-time PCR analysis was performed as previously described (56). Briefly, total RNA was reverse transcribed using the PrimeScript RT reagent kit with gDNA Eraser (Takara Bio Inc.) and subjected to PCR using the following primers: for ARF, 5′-GCCGCACCGGAATCCT-3′ (sense) and 5′-TTGAGCAAGAAGACTGCTACGT-3′ (antisense); INK4a, 5′-CCCAAGCGCCCGAACT-3′ (sense) and 5′-GCAGAAAGAGCTGCAAGTGA-3′ (antisense); IL-6, 5′-GAATGGGGTAGGAAGAC-3′ (antisense) and 5′-GAAATGGGTAGGAAGAC-3′ (antisense); p16INK4a, 5′-TCAAGCAAGAAGAGAGTCTGGA-3′ (antisense); p19ARF, 5′-GGCGACACTTGAGATGAGCTTCC-3′ (sense) and 5′-ATGGTGAAGGTCGGTGTG-3′ (antisense); β-Actin, 5′-GCCGCACCGGAATCCT-3′ (sense) and 5′-TTGGCTGAGTGGTAGAGTCCC-3′ (antisense); GAPDH, 5′-GAGCCACTAGGCCATCCTGG-3′ (sense) and 5′-CTGAGCAAGATCCATGCTTGG-3′ (antisense); MMP-3, 5′-ACATGGAGACCTTTGGCCTTTTG-3′ (sense) and 5′-CTGAGCAAGATCCATGCTTGG-3′ (antisense); MMP-12, 5′-GGCGACACTTGAGATGAGCTTCC-3′ (sense) and 5′-ATGGTGAAGGTCGGTGTG-3′ (antisense); β-Actin, 5′-GCCGCACCGGAATCCT-3′ (sense) and 5′-TTGGCTGAGTGGTAGAGTCCC-3′ (antisense); VIII and 18S rRNA, 5′-GANCCCTGCAAGCTTTTCTACACA-3′ (sense) and 5′-GATCCGGAGGCGCTCCTAACC-3′ (antisense).

**Cell sorting.** Epithelial and mesenchymal cell separation was performed as previously reported (57). Lungs were dissected from wild-type or ARF-DTR mice, and tracheae and extrapulmonary airways were removed. Lungs were then minced using a razor blade, suspended in PBS, and incubated in 5 mg/ml of Dispase II (Roche Applied Science) at 37°C for 60 minutes with gentle shaking. Cells were disaggregated with an 18-gauge needle, filtered through a 50-μm nylon cell strainer (Sysmex), and washed twice with PBS containing 2% fetal bovine serum. Low density cells were isolated by discontinuous density gradient centrifugation using Nycodren (Nycodren Pharma) in order to remove contaminating erythrocytes, neutrophils, and cell debris. Cells were negatively selected using magnetic bead–conjugated anti-CD31 (1:10 dilution, 130-097-418, Miltenyi Biotec) and –CD45 (1:10 dilution, 132-052-301, Miltenyi Biotec). Selected cells were incubated with Alexa Fluor 488–conjugated anti-mouse CD31 (1:50 dilution, 102513, BioLegend), APC-Cy7–conjugated anti-mouse EpCAM (CD326) (1:200 dilution, 118211, BioLegend), and Pacific Blue–conjugated anti-mouse Sca-1 (1:200 dilution, 108119, BioLegend). Sorting was performed using BD FACSaria II (BD Biosciences).

**Microarray.** Four ARF-DTR mice per group and two wild-type mice per group were independently analyzed. RNA was extracted from lung tissue using the RNeasy mini kit (Qiagen) according to the manufacturer’s instructions. RNA quality and quantity were determined using the Agilent 2100 Bioanalyzer (Agilent Technologies) and Nanodrop ND1000 (Thermo Scientific). An RNA probe was prepared using the Low Input Quick Amp Labeling Kit (Agilent Technologies). Labeled RNA was hybridized to the SurePrint G3 Mouse Gene Expression 8×60K Microarray (Agilent Technologies) and scanned by the Agilent DNA microarray Scanner (G2539a) using Scan Control (version A.8.5.1). The scanned images were analyzed with Agilent Feature Extraction (version 10.10.1.1). Full microarray data and protocol details are available from the GEO database, accession GSE64754; http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE64754).

A pathway analysis was performed using KEGG (Kyoto Encyclopedia of Genes and Genomes) (58, 59) gene sets with Molecular Signatures Database program version 5.0 (http://software.broadinstitute.org/gsea/msigdb/index.jsp).
Statistics. For the comparison of 3 sets of data, 1-way ANOVA was performed. When the statistical model was proven significant, the differences between combinations of the 2 groups were analyzed using a Tukey-Kramer test. For the comparison of 2 sets of experimental data, a 2-tailed unpaired Student’s t test was used. Data displayed normal variance. P values of less that 0.05 were considered significant. No blinding was performed, except for histological quantifications. No statistical method was used to determine sample size.

Study approval. All animal experiments were approved by and conducted in accordance with guidelines established by the National Center for Geriatrics and Gerontology Animal Ethics Committee.

Author contributions
MH, AA, HK, RM, KK, and KS designed and performed experiments. YI performed microarray analysis. MM and TS interpreted data. MS designed experiments and wrote the paper.

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