p16\textsuperscript{INK4a} reporter mice reveal age-promoting effects of environmental toxicants

Jessica A. Sorrentino,1,2 Janakiraman Krishnamurthy,2,3 Stephen Tilley,4 James G. Alb Jr.,2,5 Christin E. Burd,6 and Norman E. Sharpless1,2,3

Introduction
Murine models have been critical to the identification of carcinogens. For example, commercially available models (i.e., BigBlue, MutaMouse, and the p53\textsuperscript{−/−} strain; refs. 1–3) represent core tools in cancer toxicology. These reagents increase the efficiency and sensitivity of carcinogen identification, while reducing animal requirements and costs. In 1987, Martin proposed the concept of gerontogens (4): environmental agents that accelerate molecular aging, analogous to carcinogens promoting neoplasia. While murine models for carcinogen testing are advanced, in vivo tools for gerontogen testing do not exist.

Mammalian aging is complex, with distinct molecular processes contributing to age-related tissue dysfunction. While no single process underpins aging, several lines of evidence suggest senescence activation is an important contributor. Markers of cellular senescence dramatically increase with aging in humans and mice, including expression of the p16\textsuperscript{INK4a} tumor suppressor gene (5–8). Moreover, activation of p16\textsuperscript{INK4a} and senescence has been linked to replicative hypomutation in neural stem cells (9), hematopoietic progenitors (10), lymphocytes (11, 12), and pancreatic β cells (13). Altered regulation of the senescence-promoting CDKN2a/b locus, which encodes p16\textsuperscript{INK4a}, p15\textsuperscript{INK4b}, and ARF, has been linked to aging phenotypes, including expression of the endogenous p16\textsuperscript{INK4a} gene into the Cdkn2a locus (18), with expression under the control of the endogenous p16\textsuperscript{INK4a} promoter and distant cis-regulatory elements (19, 20). Importantly, p16\textsuperscript{INK4a} expression is negligible in healthy cells from young mice, but increases more than 300-fold on a per-cell basis with age-promoting stresses (7, 18). The large, dynamic range of p16\textsuperscript{INK4a} expression makes p16\textsuperscript{LUC} a valuable tool for in vivo imaging of senescence induction. To assess the effect of candidate gerontogens — high-fat diet (HFD), UV light, arsenic, and cigarette smoke (CS) — on senescence, we performed serial analysis of p16\textsuperscript{LUC} cohorts. Prior studies on the effects of obesity and HFD on p16\textsuperscript{INK4a} expression in vivo have yielded inconsistent results. HFD has been reported to increase senescence and/or p16\textsuperscript{INK4a} expression in vascular cells (21, 22), whereas p16\textsuperscript{INK4a} mRNA expression in peripheral blood T lymphocytes (PBTLs) does not correlate with body mass index, a marker of obesity, in humans (8). To examine the effects of HFD on whole-body senescence, littermate p16\textsuperscript{LUC} mice were placed on a 42%-fat HFD or 4%-fat normal diet (ND) for 18 months starting at 8–10 weeks of age. HFD feeding produced a clear pharmacodynamic effect, causing significant weight gain and hepatic steatosis, although no differences in overall survival were noted (Supplemental Figure 1, A–C; supplemental material available online with this article; doi:10.1172/JCI70960DS1). Consistent with prior reports (7, 18), we observed an exponential increase in p16\textsuperscript{INK4a} expression with aging; however, the rate of increase, as assessed by total-body luciferase imaging (TBLI), was not different between cohorts (Figure 1, A and B). Due to the wide range of weight gain among HFD-fed mice, we analyzed the relationship between weight and luciferase intensity at 52 weeks and observed no correlation (Supplemental Figure 1D). Corre-
spondingly, expression of p16<sup>INK4a</sup> mRNA was not altered in liver and spleen harvested from 18-month-old mice fed HFD versus ND (Supplemental Figure 1E). These data suggested that HFD does not accelerate p16<sup>INK4a</sup> expression in vivo, at least within the sensitivity of luciferase detection.

Arsenic is an environmental toxicant linked to human aging-associated phenotypes (i.e., several cancers, type II diabetes, and atherosclerosis; ref. 23). While not a direct clastogen, arsenic may indirectly affect DNA damage and repair (24–26). We examined the effect of chronic arsenic exposure on p16<sup>INK4a</sup> expression in vivo by administering 0 or 50 ppm arsenic to littermate p16<sup>LUC</sup> mice in the drinking water for 12 months starting at 8–10 weeks of age. After 48 weeks of exposure, arsenic-treated mice exhibited significant hepatic accumulation (Supplemental Figure 2A). Consistent with known effects of arsenic (27), treated mice showed small, structurally deranged pancreatic islets with an extreme decrease in total islet mass (Supplemental Figure 2, B and C). Overall survival rates were not different between cohorts (Supplemental Figure 2D). A modest but significant increase in p16<sup>INK4a</sup> expression was observed by 24 weeks of exposure using TBLI (P = 0.023; Figure 1, C and D). Additionally, expression of p16<sup>INK4a</sup> mRNA was higher in spleen, but not liver, of 1-year-old mice administered 50 versus 0 ppm (Supplemental Figure 2E). These data suggest that chronic arsenic exposure modestly accelerates the accumulation of p16<sup>INK4a</sup>-expressing cells.

CS promotes DNA damage (28, 29) and age-related diseases such as emphysema, atherosclerosis, and several cancers (30, 31). Chronic CS exposure in humans is associated with increased p16<sup>INK4a</sup> expression in PBTLs (8, 32). To determine the effects of moderate CS exposure on cellular senescence in vivo, littermate p16<sup>LUC</sup> mice were exposed to CS or ambient air (AA) for 1 hour per day, 5 days per week for 6 months starting at 10–12 weeks of age. After 6 months of exposure, mice were further observed without CS up to 1 year of age. At the end of the study, histological analysis of the sinus cavity of CS-exposed mice showed particulate deposits along the epithelial layer of the nasal papilla, whereas no differences were noted in lung architecture between CS and AA cohorts at this tobacco dose (Figure 2A), which

**Figure 1**

HFD does not increase p16<sup>INK4a</sup> expression, while arsenic moderately increases p16<sup>INK4a</sup> expression. p16<sup>LUC</sup> mice were (A and B) fed ND or HFD or (C and D) exposed to 0 or 50 ppm arsenic. (A and C) Luciferase intensity, normalized to mean luciferase levels of the entire cohort at day 0 and graphed as a function of time. Exposure duration is shown by the orange bar on the x axis. Error bars indicate SEM. P values were determined by linear regression analysis. (B and D) Representative TBLI of experiments in A and C, respectively.
was also not associated with accelerated mortality (Supplemental Figure 3A). A significant increase in \( p16^{INK4a} \) induction between the CS and AA cohorts, as measured by TBLI, was readily noted within weeks of CS exposure and persisted after CS was discontinued (Figure 2B). Consistent with histologic examinations of the nasal cavities and lungs (Figure 2A), luciferase induction was noted in the head and neck region of mice, but not the thoracic and abdominal regions (Figure 2C). Correspondingly, \( p16^{INK4a} \) mRNA levels in the liver, lung, axillary lymph nodes, and spleen were similar in CS- and AA-exposed mice (Supplemental Figure 3B). These data suggest that low-dose CS exposure accelerates accumulation of senescent cells in tissues that receive the highest exposure (i.e., nasal epithelium).

Chronic UVB exposure causes photoaging and skin cancer (33, 34). To directly assess whether chronic UVB induces senescence in vivo, the dorsal surface of \( p16^{LUC} \) mice was exposed to 350 J/m\(^2\) UVB light 3 times a week for 6 months starting at 10–12 weeks of age. Littermate control mice were contemporaneously aged in the absence of UVB. After 6 months of exposure, mice were observed without further UVB exposure until 1 year of age. After 6 months of exposure, UV-exposed mice showed hyperkeratosis and metaplasia and inflammatory infiltration of the dermis and hypodermis compared with the nonexposed cohort (Figure 3A). The DNA damage marker phospho-H2AX was substantially increased in the epithelium of UV-exposed mice (Figure 3A and Supplemental Figure 4A), suggesting a persistent DNA damage response. Additionally, UV-exposed mice developed skin neoplasms within 5 months of exposure, which accelerated mortality (Figure 3, B and C). Luciferase expression was higher in \( p16^{LUC} \) animals treated with UV than in unexposed controls prior to development of visible neoplasms, and TBLI was almost 8 times greater in UV-treated animals after 32 weeks of exposure (Figure 3D). In accord with the TBLI results (Figure 3E), \( p16^{INK4a} \) mRNA levels from abdominal and dorsal skin from both cohorts confirmed the site-specific induction of \( p16^{INK4a} \) expression (Figure 3F). Additionally, the UV-exposed dorsal skin exhibited higher expression than unexposed abdominal skin from the same mouse (Figure 3F).

To strengthen the association of \( p16^{INK4a} \) induction with senescence, additional markers were analyzed in UV-exposed and unexposed skin, including mRNA of the senescence-associated (SA) cytokines \( Cxcl1 \) and \( Il6 \) and SA-\( \beta \)-galactosidase (SA-\( \beta \)-gal) staining. \( Cxcl1 \) and \( Il6 \) expression increased in UV-exposed versus nonexposed mice. Additionally, increased expression only occurred in skin directly exposed to UV light (back) and not in unexposed skin (abdomen) from the same animals (Figure 3, G and H). We observed a similar pattern of SA-\( \beta \)-gal staining (Supplemental Figure 4B). These data suggest that chronic DNA damage from UV light accelerates senescent cell accumulation.

Here, we demonstrated that \( p16^{LUC} \) mice can detect the senescence-promoting effects of environmental exposures. Direct DNA-damaging agents (CS and UV light) appeared potently gerontogenic, and arsenic exposure produced a lesser, but significant, effect. We also identified anatomic gerontogenic effects of these
Figure 3
UVB increases $p16^{INK4a}$ expression. $p16^{JC}$ mice were exposed to 0 J/m² ($n = 13$) or 350 J/m² ($n = 11$) UVB light. (A) Representative histopathological images of skin stained with H&E or phospho-H2AX. Original magnification, ×20 (top), ×40 (middle and bottom), ×200 (top, insets), ×120 (middle and bottom, insets). (B) Representative dorsal images. (C) Kaplan-Meier survival curves. Exposure duration is shown by the orange bar on the x axis. $P$ value was determined by Gehan-Breslow-Wilcoxon test. (D) Luciferase intensity, normalized to mean levels of the entire cohort at day 0 and graphed as a function of time. Exposure duration is shown by the orange bar on the x axis. Inset highlights initial 20 weeks of exposure. Error bars denote SEM. $P$ value was determined by linear regression analysis. (E) Representative TBLI of experiments in D. (F–H) Quantitative real-time PCR of $p16^{INK4a}$ (F), Cxcl1 (G), and Il6 (H) expression in back and abdominal skin at sacrifice. Error bars denote SEM. *$P < 0.05$, **$P < 0.01$, ***$P < 0.005$, Student’s t test.
agents, including a stronger effect of passively inhaled CS on nasal epithelia than lung parenchyma (Figure 2C) and effects of UVB on agents, including a stronger effect of passively inhaled CS on nasal epithelia rather than unexposed skin (Figure 3, E and F).

In summary, we demonstrated using modest-sized cohorts of serially analyzed animals that the p16(INK4a) allele was useful in vivo activation of senescence markers and uncovered gerontogenic effects of environmental toxicants. This resource can supplement other rodent platforms used for toxicological assessments; specifically, it represents an in vivo system with which to assess a compound’s age-promoting activity. We believe this resource will provide an important tool for understanding the relationship between environmental exposures and molecular aging.

Methods

Further information can be found in Supplemental Methods.

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Hairsless SKH1-E1 p16INK4a/c mice were used for all experiments (18). Geno- typing was performed as previously described (18). For histologic analysis, tissues were fixed with 10% formalin overnight, then transferred to 70% etha- nol for paraffin blocking and staining. Quantitative TaqMan RT-PCR strate- gies for detection of p16INK4a were performed as previously described (7, 13).

Statistics. Statistical significance was determined using 2-tailed Student’s t test or linear regression analysis for all comparisons, except survival analy- sis (Gehan-Breslow-Wilcoxon test). A P value less than 0.05 was considered statistically significant.

Study approval. Experiments were performed under protocols approved by the University of North Carolina IACUC.

Acknowledgments

We thank H. Yuan, A. Deal, and S. Gaddameedhi for advice. This work relied on the UNC BRIC Small Animal Imaging Facility, Biostatistics Core, Microscopy Services Laboratory, Histopathology Core, Mouse Phase 1 Unit, and Animal Studies Facility. This work was supported by the NIA (AG024379 and AG036817), the NIHES (T32 ES07126), and an HHMI training grant (Med into Grad Initiative).

Received for publication May 8, 2013, and accepted in revised form October 7, 2013.

Address correspondence to: Norman E. Sharpless, The Lineberger Comprehensive Cancer Center, University of North Carolina School of Medicine, CB #7295, Chapel Hill, North Carolina 27599, USA. Phone: 919.966.1185; Fax: 919.966.8212; E-mail: nes@med.unc.edu.

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