Premature senescence involving p53 and p16 is activated in response to constitutive MEK/MAPK mitogenic signaling

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Oncogenic Ras transforms immortal rodent cells to a tumorigenic state, in part, by constitutively transmitting mitogenic signals through the mitogen-activated protein kinase (MAPK) cascade. In primary cells, Ras is initially mitogenic but eventually induces premature senescence involving the p53 and p16INK4a tumor suppressors. Constitutive activation of MEK (a component of the MAPK cascade) induces both p53 and p16, and is required for Ras-induced senescence of normal human fibroblasts. Furthermore, activated MEK permanently arrests primary murine fibroblasts but forces uncontrolled mitogenesis and transformation in cells lacking either p53 or INK4a. The precisely opposite response of normal and immortalized cells to constitutive activation of the MAPK cascade implies that premature senescence acts as a fail-safe mechanism to limit the transforming potential of excessive Ras mitogenic signaling. Consequently, constitutive MAPK signaling activates p53 and p16 as tumor suppressors.

[Key Words: MAPK signaling; premature senescence; MEK; tumor suppression; p53; p16]

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transformed by oncogenic Ras alone. Similarly, immortalizing oncogenes disrupt the senescence program retained in primary cells; hence, such changes interfere with the ability of oncogenic Ras to provoke senescence. In either case, cells escape normal growth controls that actively limit transformation by Ras. Consequently, premature senescence may function as a bona fide mechanism of tumor suppression (Serrano et al. 1997; for review, see Weinberg 1997).

Consistent with this view, p53 and p16INK4a appear critical for premature senescence induced by oncogenic Ras (Serrano et al. 1997). Both p53 and INK4a are mutated at high frequency in many tumor types, implying that their action is central to tumor development. Nevertheless, the precise circumstances in which either is engaged to function as tumor suppressors remain poorly understood. p53 can promote cell-cycle arrest or apoptosis in response to a variety of cellular stresses, including DNA damage and hypoxia (Kastan 1993; Graeber et al. 1996). In both settings, cells acquiring p53 mutations have a selective advantage over their p53-normal counterparts. p16 increases gradually as cells proceed towards senescence (Alcorta et al. 1996; Zindy et al. 1997). Consequently, loss of p16 might uncouple the cell-cycle machinery from a cell-doubling clock, allowing developing tumor cells to proliferate beyond their normal life span. The fact that oncogenic Ras activates p53 and p16 provides an additional explanation for their action as tumor suppressors: p53 and p16 act in a compensatory mechanism that suppresses Ras-induced transformation. Hence, cells acquiring p53 and INK4a mutations would tolerate ras mutations occurring early in tumor development (Serrano et al. 1996).

In immortal rodent lines, transformation by oncogenic Ras involves its ability to bind and activate a series of effector proteins, including Raf-1, phosphoinositide 3-OH kinase (PI(3)K), and Ral.GDS (Van Aelst et al. 1993; Rodriguez-Viciiana et al. 1994; Spaargaren and Bischoff 1994). Each of these proteins, in turn, activates distinct downstream targets, thereby producing different aspects of the transformed phenotype (for review, see Katz and McCormick 1997). For example, the Ras–Raf interaction initiates the mitogen-activated protein kinase (MAPK) cascade, which involves the sequential activation of a series of protein kinases that transmit mitogenic signals to nuclear transcription factors (Chen et al. 1992; Gilles et al. 1995). These kinases include Raf-1, the MEKs (MEK1 and MEK2), and the MAPKs (ERK1 and ERK2). In contrast, the ability of Ras to activate PI(3)K promotes membrane ruffling (Joneson et al. 1996; Rodriguez-Viciiana et al. 1997), perhaps through Rac and Rho (for review, see Van Aelst and D’Souza-Schorey 1997), and may also suppress apoptosis through activation of Akt/PKB (for review, see Downward 1998). Finally, the Ras.GDS proteins act as exchange factors that can activate the Ras family of small GTPases (Spaargaren and Bischoff 1994) which, in turn, can regulate phospholipase D (Jiang et al. 1995). Although each of these effector pathways contributes to the transforming activity of Ras in immortal rodent fibroblasts (Khosravi-Far et al. 1995, 1996; White et al. 1995; Urano et al. 1996; Rodriguez-Viciiana et al. 1997), activation of the MAPK cascade is clearly sufficient (Cowley et al. 1994; Mansour et al. 1994; Bottorff et al. 1995; Stang et al. 1997).

Almost all of the studies examining Ras signaling in mammalian systems have utilized immortal or tumor-derived lines harboring unknown genetic alterations. Because premature senescence induced by Ras is restricted largely to nonimmortal cells (Serrano et al. 1997), Ras might signal premature senescence through an unidentified pathway lost during the immortalization process. Alternatively, Ras could promote premature senescence through a known effector pathway or by activating a combination of effector pathways. In the current study, we identified the effector pathway responsible for Ras-induced senescence. We find that Ras induces senescence in normal cells through the primary mechanism whereby it forces uncontrolled proliferation and transformation in immortal cell lines. These results have important implications for our understanding of oncogene cooperation, tumor suppression, and the regulation of cellular senescence.

Results

Oncogenic Ras is initially mitogenic in normal diploid human fibroblasts

Microinjection of oncogenic ras into human diploid fibroblasts induces S-phase entry (Lumpkin et al. 1986). On the other hand, retroviral transduction of oncogenic ras into human fibroblasts provokes permanent cell-cycle arrest (Serrano et al. 1997). These observations raise the possibility that the initial effect of Ras is forced proliferation and, only later, cell-cycle arrest. Therefore, the immediate and long-term consequences of oncogenic Ras expression were examined in the same population of normal diploid human IMR90 fibroblasts. In this and subsequent experiments, oncogenic ras (H-RasV12) was introduced into whole-cell populations using high-titer retroviral vectors coexpressing a selectable marker. After a brief selection to eliminate uninfected cells, cells were plated for all assays described below (designated day −1). Using this protocol, the percentage of transduced cells ranged between 70% and 90% as estimated in parallel infection with viruses expressing a lacZ reporter (data not shown).

To examine the proliferation properties of Ras-expressing populations, [3H]thymidine and BrdU incorporation were measured shortly after ras transduction (48–72 hr postinfection) in the presence and absence of serum. In normal growth conditions (10% serum), both vector and Ras-expressing cells incorporated similar amounts of [3H]thymidine (Fig. 1A). As expected, transfer of vector-containing cells to low serum for 24 hr resulted in a marked reduction in [3H]thymidine incorporation. By contrast, Ras-expressing cells continued to incorporate high levels of [3H]thymidine even after serum depletion. Similarly, Ras-expressing cells placed in low serum incorporated twice as much BrdU as cells containing an
empty vector (Fig. 1B). Initially, proliferation in the presence of oncogenic Ras was accompanied by morphological changes characteristic of constitutive Ras activity in immortal cells, becoming small and refractile with thin cytoplasmic projections (Fig. 1C). Only later do Ras-expressing cells acquire the characteristic senescent morphology (Fig. 1C; see also Serrano et al. 1997). Therefore, the immediate consequences of oncogenic Ras expression are forced proliferation, implying that cell-cycle arrest is an antiproliferative cellular response.

Ras effectors and cell-cycle arrest

As a first step in identifying the Ras-signaling pathway(s) required for premature senescence, we exploited a series of Ras ‘effector loop’ mutants that interact preferentially with specific Ras effector proteins (White et al. 1995; Joneson et al. 1996). For example, H–RasV12/S35 binds preferentially to Raf-1 and activates MAPK but has no effect on membrane ruffling (White et al. 1995). In contrast, H-RasV12/C40 associates with PI(3)K and promotes membrane ruffling and cell survival but does not bind Raf-1 or induce mitogenesis (Joneson et al. 1996; Rodríguez-Viciiana et al. 1997). H-RasV12/G37 binds Ral.GDS but fails to bind Raf-1 or PI(3)K (White et al. 1995). IMR90 human fibroblasts were infected with retroviruses expressing individual Ras mutants, and cell proliferation was determined by assessing relative cell accumulation at various times postplating (Serrano et al. 1997).

Consistent with previous results, constitutive expression of oncogenic Ras induced cell-cycle arrest at subconfluent densities and in the presence of serum, whereas cells harboring the empty vector grew exponentially to confluence (Fig. 2A). Cell populations expressing the H–RasV12/G37 or H-RasV12/C40 mutants failed to arrest, suggesting that neither the Ras-PI(3)K interaction nor the Ras-Ral.GDS interaction is sufficient to in-
duce cell-cycle arrest. In contrast, cells expressing H-RasV12/S35 grew slowly, at rates similar to cells expressing H-RasV12 (Fig. 2A). Of note, each Ras mutant was expressed at levels equal to (or greater than) H-RasV12, ranging from a 5- to 10-fold increase over endogenous Ras expression (data not shown). Therefore, the differential effects of each mutant were not caused by variations in Ras expression but, more likely, by their ability to interact with distinct effector proteins. These results correlate the Ras-Raf interaction with Ras-induced cell-cycle arrest in normal diploid human fibroblasts.

Because the Ras effector loop mutants were characterized in immortal cell lines, it is formally possible that they possess different activities in primary cells. As an alternative approach, selected Ras effectors (Raf-1, PI3K, downstream components (Rac, Akt, RhoA, MEK1) (see Introduction), or Ras-related proteins (R-Ras, TC21) were tested for their ability to cause growth arrest by retroviral transduction into IMR90 human fibroblasts. Of note, R-Ras is highly homologous to H-Ras, and interacts with PI3K and activates the PI3K/Akt pathway but not the MAPK pathway in vivo (Marte et al. 1997). TC21 is more closely related to H-Ras than any other known member of the Ras superfamily but does not activate Raf (Graham et al. 1996). Consequently, H-Ras, R-Ras, and TC21 stimulate cell growth and transformation via distinct signaling pathways. After selection for virus-infected populations, cells were plated at subconfluent densities and pulsed with [3H]thymidine in the presence of 10% serum. Among the Ras effectors and downstream components examined, some enhanced proliferation, whereas others had little effect (Fig. 2B). Only constitutively active Raf-1 and MEK—components of the MAPK cascade—inhibited proliferation (Fig. 2B; see also Zhu et al. 1998). Neither R-Ras nor TC21 produced cell-cycle arrest. Therefore, both the Ras mutant and effector analysis are consistent: Ras-induced arrest in normal diploid human fibroblasts involves activation of the MAPK cascade.

Activation of the MEK/MAPK cascade induces p53, p16, and premature senescence

The MEKs are dual specificity protein kinases that, upon activation, phosphorylate the threonine and tyrosine regulatory sites of MAPKs (for review, see Marshall 1994). MEK1^Q56P is a MEK mutant identified by virtue of its ability to enhance the transforming activity of otherwise transforming-defective Ras mutants. Substitution of glutamate to proline at codon 56 results in a gain-of-function mutant that has increased kinase activity (Bottorff et al. 1995). To better characterize MEK-induced arrest, the proliferation properties of IMR90 cell populations containing a control vector, oncogenic Ras, or MEK1^Q56P were measured by growth curves, or by BrdU-labeling and flow cytometry (Serrano et al. 1997). Activated MEK causes arrest even more rapidly than oncogenic Ras, with cells displaying a dramatic reduction in BrdU incorporation [compare 0.6% (MEK) and 0.3% (Ras) to 24.5% (vector) (Fig. 3A,C)]. Likewise, MEK-expressing cells arrest primarily with a G1 DNA content, although there is a clear G2 component (Fig. 3C). Importantly, both oncogenic Ras and MEK1^Q56P activate MAPK in human fibroblasts (Fig. 3B).

The adenovirus E1A oncoprotein cooperates with ras to transform primary rodent fibroblasts (Ruley 1990) and abrogates Ras-induced senescence (Serrano et al. 1997). To determine the effect of E1A on MEK-induced arrest, IMR90 cells were infected sequentially with retroviruses expressing E1A, and then either MEK1^Q56P or a control
Cell populations coexpressing E1A and MEK1^{Q56P} accumulated at rates comparable to vector-containing cells and incorporated high levels of BrdU, showing no signs of cell-cycle arrest (Fig. 3A,C). Consequently, as in the case of Ras-expressing cells, E1A effectively counters MEK-induced arrest and allows uncontrolled proliferation.

Cellular senescence is accompanied by a series of changes that, together, distinguish senescence from quiescence or differentiation. These changes include altered expression of cell-cycle proteins, upregulation of p53, p16, and p21, and the accumulation of senescence-associated β-galactosidase (SA-β-gal) (Noda et al. 1994; Adadja et al. 1995; Dimri et al. 1995; Alcorta et al. 1996; Reznikoff et al. 1996). Like cells expressing oncogenic Ras, MEK1^{Q56P}-expressing cells harbored underphosphorylated Rb and lost cyclin A expression (Fig 4). Also, MEK induced p53 (about threefold induction), p21 (about fivefold induction), and p16 (about sixfold induction) (Fig. 4). Finally, MEK1^{Q56P}-expressing cells exhibited high levels of SA-β-gal activity, with a percentage of positive cells similar to that observed in Ras-expressing or late passage IMR90 cells (Fig. 5A,B). Of note, these effects require constitutive MEK activation, because cells overexpressing a wild-type MEK allele grow normally, display normal levels of p53, p21, p16, and are negative for SA-β-gal activity (data not shown).

These data indicate that the cell-cycle arrest produced by oncogenic Ras and activated MEK is virtually identical and is characteristic of cellular senescence. Indeed, the only difference we observed between these populations is their morphology: Whereas Ras-arrested cells became large and flat, MEK-arrested cells were small and refractile (Fig. 5A). Nevertheless, MEK-expressing cells remain arrested in a metabolically active state for as long as we have observed them (>2 weeks). Because Rho family members mediate changes in cell morphology in other settings (for review, see Van Aelst and D’Souza-Schorey 1997), they may contribute to the morphological changes observed in Ras-arrested cells. Nevertheless, cell morphology has been dissociated from senescence in other settings (Wistrom and Villeponteau 1990). Thus, activation of the MAPK cascade is sufficient to promote premature senescence.

MEK activation is required for premature senescence

To determine whether MEK activity was required for senescence, we took advantage of PD98059, a highly spe-
specific noncompetitive MEK inhibitor that prevents its activation by Raf (Alessi et al. 1995; Dudley et al. 1995). IMR90 cells were infected with control or oncogenic Ras-expressing retroviruses, and treated daily with 50 µM PD98059 prior to the onset of cell-cycle arrest and the appearance of senescence-related markers (3 days postinfection). Notably, this drug concentration has been found previously to be compatible with the proliferation of fibroblasts albeit at slower rates (Pumiglia and Decker 1997). All cultures were monitored for proliferation and senescence as described above.

Remarkably, PD98059 prevented the onset of Ras-induced arrest but had no effect on the accumulation of vector-containing cells (Fig. 6A). Although the accumulation rate of Ras-expressing cells treated with PD98059 did not achieve that of untreated cells harboring an empty vector, they displayed an approximately sixfold increase in BrdU incorporation compared to Ras-expressing cells treated with solvent alone (Fig. 6B). Similarly, treatment with PD98059 prevented the appearance of the senescence-like morphology and the upregulation of SA-β-galactosidase in Ras-expressing cells (Fig. 6C).

Moreover, PD98059 had no effect on the detection of SA-β-gal activity in late passage (senescent) cells, demonstrating that PD98059 does not directly inhibit this enzyme. Thus, these data imply that MEK—and hence the MAPK cascade—is required for premature senescence induced by oncogenic Ras.

MEK arrests primary murine fibroblasts but transforms fibroblasts lacking p53 or p16INK4a

Premature senescence may act as a fail-safe mechanism that suppresses transformation by Ras (Serrano et al. 1997). Thus, whereas oncogenic Ras arrests primary rodent fibroblasts, it readily transforms primary fibroblasts lacking p53 or INK4a (Tanaka et al. 1994; Serrano et al. 1996). Because MEK is both necessary and sufficient for Ras-induced senescence in human fibroblasts, we asked whether inactivation of p53 or INK4a would overcome MEK-induced arrest in primary rodent cells. Therefore, wild-type, p53-null, and INK4a-null primary mouse embryo fibroblasts (MEFs) were infected with control vector-, Ras-, and MEK1Q56P-expressing retroviruses, and the rate of cell accumulation was examined in the resulting populations. Of note, our INK4a-null cells have deletions in INK4a exon 2 that may also inactivate p19ARF (Haber 1997, see discussion); hence, these cells are referred to as INK4aex2−/− for clarity.

As in human fibroblasts, MEF promoted cell-cycle arrest in wild-type MEFs (Fig. 7A). However, unlike in human cells, MEK-arrested MEFs acquired the enlarged and flat senescent morphology characteristic of Ras-expressing MEFs (data not shown). Remarkably, when MEK1Q56P was expressed at comparable levels in primary p53−/− and INK4aex2−/− MEFs (Fig. 7B), the resulting populations grew at rates similar to parallel cultures expressing oncogenic Ras, with no indication of a growth delay or arrest. Moreover, MEK-expressing p53−/− and INK4aex2−/− MEFs appeared transformed morphologically (data not shown) and continued to proliferate in low serum (Fig. 7C). Therefore, deletion of either p53 or INK4aex2−/− was sufficient to alter the cellular response to MEK from permanent arrest to uncontrolled mitogenesis.

The transformation properties of virus-infected populations were assessed by their ability to grow in suspension and form tumors in immunocompromised mice. In soft-agar assays, MEK-expressing populations grown minimally in culture derived from p53−/− and INK4aex2−/− cells formed colonies at high frequency (Fig. 7D). Similarly, MEK-expressing populations derived from p53−/− and INK4aex2−/− MEFs formed tumors at high frequency when injected subcutaneously into athymic nude mice (Table 1). Importantly, the transforma-

Figure 6. MEK is required for Ras-induced cell-cycle arrest and premature senescence. (A) Representative growth curves documenting the effects of PD98059 on Ras-induced growth arrest. IMR90 cells were infected with an empty vector- or an H-RasV12-expressing retrovirus. Immediately after selection, cell populations were treated daily with medium containing 50 µM PD98059 or 0.25% DMSO (carrier). Each value was determined in triplicate and normalized to the cell number at day 0. (B) Cell populations containing an empty vector (V) or expressing H-RasV12 (R) were treated daily with PD98059 (PD). On day 6 postselection, cell cultures were pulsed with BrdU and analyzed for BrdU incorporation by flow cytometry. The calculated percentage of BrdU-positive cells is shown. (C) Photomicrographs of control and PD98059-treated cell populations stained for SA-β-gal activity on day 6 postselection. The percentage of positive cells in each population was 9% (vector); 12.5% (vector + PD98059); 80% (Ras); 15.4% (Ras + PD98059); 90% (late passage); 91% (late passage + PD98059).
tion properties of MEK-expressing cells required cooperation between MEK and loss of either p53 (see also Fukasawa and Vande Woude 1997) or INK4aex2, because p53−/− and INK4aex2−/− MEFs harboring an inert vector neither grew in soft agar nor formed tumors in nude mice. However, MEK1Q56P was not as potent as oncogenic Ras in transforming p53−/− and INK4aex2−/− MEFs, because both soft-agar colonies and tumors expressing MEK were typically smaller than those expressing oncogenic Ras (Fig. 7D; Table 1). The inability of activated MEK to recapitulate the potency of oncogenic Ras in transformation assays is consistent with previous studies showing that multiple effector pathways contribute to Ras-mediated transformation (White et al. 1995).

These data demonstrate that activated MEK efficiently transforms primary MEFs lacking p53 or INK4aex2, cells otherwise predisposed to immortalization. Because these MEK-expressing cell populations were derived from primary fibroblasts grown minimally in culture, it seems highly unlikely that additional factors contribute to the transformed phenotype of these cells. Similarly, we show that human fibroblasts expressing an immortalizing oncogene (i.e., E1A) overcome MEK-induced arrest. Together, these data demonstrate that constitutive activation of the MEK/MAPK cascade can produce either premature senescence or forced mitogenesis depending on the integrity of a senescence program controlled by p53 and p16. The precisely opposite response of primary and immortal cells to MEK activation implies that premature senescence is a cellular fail-safe mechanism that suppresses the transforming potential of aberrant Ras mitogenic signaling.

### Discussion

Oncogenic Ras transforms immortal rodent fibroblasts, in part, by constitutively transmitting mitogenic signals through the MAPK cascade. By marked contrast, oncogenic Ras acutely activates senescence in primary fibroblasts through the p53 and p16 tumor suppressors (Serrano et al. 1997). Here we demonstrate that Ras signals premature senescence through activation of the MAPK cascade. In normal diploid human fibroblasts, Ras effector loop mutants that retain their ability to bind Raf-1 promoted premature senescence and, among a series of Ras downstream components examined, only activated MEK was capable of inducing p53, p16, and features of...
senescence. Moreover, a MEK inhibitor prevented Ras-induced cell-cycle arrest and senescence. In primary murine fibroblasts, activated MEK arrested wild-type MEFs but forced uncontrollable mitogenesis and transformation when expressed at comparable levels in p53−/− or Ink4a−/− mice. Thus, oncogenic Ras arrests normal cells through the primary mechanism whereby it transforms immortal cells or primary fibroblasts defective in p53 or p16 function.

Role of the MAPK cascade in cell-cycle arrest

Signaling through the MAPK cascade is relevant to several physiological processes, including mitogenesis, differentiation, and even apoptosis, depending on the cellular context (for review, see Marshall 1998). In some settings, the difference between proliferation and arrest is related to the intensity or duration of the signal (Sewing et al. 1997; Woods et al. 1997; Kerkhoff and Rapp 1998). We show that oncogenic Ras initially forces uncontrollable proliferation, and only later does cell arrest. Importantly, the cell-cycle arrest induced by expression of Ras or MEK is abolished in cells expressing E1A (IM R90), or lacking p53 or p16 (MEFs). In contrast to our observations, others have found that high levels of Raf activity can produce an immediate cell-cycle arrest that occurs independently of p53 or p16 (Sewing et al. 1997; Woods et al. 1997). Perhaps the difference between these studies and ours reflects the fact that other Ras effector pathways can attenuate Raf-induced arrest and impinge upon its ability to activate components of the arrest machinery (Olson et al. 1998); alternatively, they may arise from different levels and kinetics of activation of the Raf/MAPK cascade. Nevertheless, the results presented here demonstrate clearly that Ras-mediated MAPK activation can produce two precisely opposite outcomes—cell-cycle arrest or forced mitogenesis—depending on the integrity of the senescence program controlled by p53 and p16.

Other mitogenic oncogenes can display antiproliferative potential. For example, the c-myc and E1A oncogenes induce proliferation, but both activate p53 to promote apoptosis (for review, see Weinberg 1997). The Myc functions involved in both proliferation and apoptosis are related (Evan et al. 1992; Amati et al. 1993) and, at least in primary cells, the E1A domains required for p53 accumulation, apoptosis, and oncogenic transformation are inseparable (Samuelson and Lowe 1997). The fact that oncogenic Ras uses the same signal-transduction pathway to promote both arrest and forced mitogenesis reinforces the view that normal cells counter malignant transformation by actively responding to hyperproliferative signals; in this case, by sensing excessive MEK/MAPK activity and activating senescence. The necessity to overcome these fail-safe mechanisms can explain the phenomenon of oncogene cooperation and may be important during multistep carcinogenesis.

The MAPK cascade and senescence

The cell-cycle arrest induced by deregulated MEK activity in primary fibroblasts reflects bona fide cellular senescence and not an unusual form of quiescence or differentiation. First, unlike quiescence, both oncogenic Ras and MEK arrest cells at subconfluent densities in the presence of mitogenic growth factors. This arrest is permanent: whereas the MEK inhibitor PD98059 prevents Ras-induced arrest when added prior to the appearance of senescence markers, it cannot reverse the arrest at later times (data not shown). Second, although p53 and p16 play fundamental roles in Ras-induced arrest, neither have essential roles in differentiation. Indeed, most p53-and Ink4a−/−-deficient mice develop normally (Donehower et al. 1992; Serrano et al. 1996). Cell-cycle arrest induced by Ras and MEK is accompanied by accumulation of p53, p21, p16, and SA-β-gal activity. Importantly, these markers together are associated with cellular senescence in both fibroblasts and epithelial cells but rarely, if ever, with quiescence or differentiation (Atadja et al. 1995; Dimri et al. 1995; Palmero et al. 1997). Finally, cDNA array analysis of ras-arrested fibroblasts reveals global changes in gene expression consistent with that reported for senescent cells (G. Ferbeye and S.W. Lowe, unpubl.).

The characterization of Ras-induced arrest in primary fibroblasts as cellular senescence, and the involvement of the MEK/MAPK cascade in this process, have important implications for our understanding of senescence biology. Senescence was defined originally by the observation that primary cells have a genetically determined limit to their proliferative potential in cell culture, after which they arrest permanently with characteristic features (Hayflick 1965). Owing to the ‘end-replication problem’, telomeres shorten during each cell division unless telomerase is expressed, and it has been proposed that some aspect of excessive telomere shortening activates cell-cycle arrest and other characteristics of senescence (for review, see Greider 1998). However, a number of exogenous stimuli can also produce a phenotype suggestive of senescence, including DNA damage, DNA-demethylating drugs, ceramide, or inhibitors of histone deacetylation (Holliday 1986; Venable et al. 1995; Ogryzko et al. 1996; Linke et al. 1997), raising the possibility that telomere shortening is not the only signal capable of activating the senescence program.

Our data demonstrate that senescence can be activated in response to deregulated MAPK activation through MEK. Consistent with these results, forced Raf-1 expression also activates senescence in normal human fibroblasts (see Fig. 2B; Zhu et al. 1998). Whereas it remains possible that telomere shortening or some other cell-doubling-sensitive mechanism activates senescence through Ras/MAPK signaling, our hypothesis is that senescence can be activated by diverse stimuli leading to engagement of a common cell-cycle arrest program. In this view, the process of senescence is related conceptually to apoptosis, a genetically controlled program carried out by a common cell-death machinery. Diverse stimuli and signaling pathways feed into this machinery, allowing regulation of cell death to be highly versatile. Although less understood, the possibility that multiple
 signaling pathways induce senescence is consistent with cell fusion studies (for review, see Voigt and Barrett 1995). Consequently, the biological roles of cellular senescence may go well beyond the control of cellular or organismal aging, acting as a global anti-proliferative response to a variety of cellular stresses.

The role of p16 and p53 in premature senescence

The precise role of p16 in tumor suppression is confounded by the complexity of the INK4a locus, which encodes a second tumor suppressor translated in an alternative reading frame designated p19ARF (Quelle et al. 1995). Mutations at the INK4a locus often disrupt both p16 and p19ARF (for review, see Haber 1997) and mice lacking p19ARF develop tumors (Kamijo et al. 1997). p19ARF also promotes cell-cycle arrest and primary cells derived from ARF-deficient mice can be transformed by oncogenic Ras alone (Kamijo et al. 1997). Because the mutation in our INK4aex2−/− MEFs may disrupt p19ARF function, it is possible that p19ARF is the sole product of the INK4a/ARF locus responsible for senescence in response to oncogenic Ras. Oncogenes can signal p53 through p19ARF (Zindy et al. 1998; de Stanchina et al. 1998); moreover, oncogenic Ras and MEK induce ARF message and protein and ARF-null MEFs are defective in Ras-induced arrest (Palmero et al. 1998; G. Ferbeyre, A.W. Lin, F. Zindy, M. Roussel, C.J. Sherr, and S.W. Lowe, unpubl.). Nevertheless, p16 contributes to Ras-induced senescence. For example, p16 accumulates in response to oncogenic Ras and MEK, and expression of a p16-insensitive Cdk4 mutant (R24C) (Wolfel et al. 1995) counters Ras-induced growth arrest (Serrano et al. 1997). Furthermore, in human cells, p16 can directly induce senescence (Uhrbom et al. 1997; M. McConnell et al. 1998) and can be inactivated spontaneously during the immortalization process (Reznikoff et al. 1996; Foster et al. 1998). Regardless of the relative contribution of p16 and p19ARF to this process, the INK4aex2−/− cells used accurately recapitulate mutations in human tumors.

In rodent cells, the cell-cycle arrest induced by activated MEK is abrogated by inactivation of either p53 or p16 (Fig. 7; see also Serrano et al. 1997). Nevertheless, p53 and p16 do not function in a simple linear pathway. p16 levels increase in response to Ras in p53−/− MEFs and, conversely, p53 levels increase in INK4aex2−/− MEFs (data not shown). Introduction of oncogenic Ras or MEK into INK4aex2−/−p53−/− double-mutant MEFs produces more transformed foci than in either single mutant (A.W. Lin and S.W. Lowe, unpubl.). Thus, p53 and p16 (and perhaps p19ARF) act cooperatively to promote Ras-induced arrest, and disruption of either p53 or p16 prevents arrest and is sufficient for transformation. In human cells, p53 and p16 also appear to cooperate to promote arrest, although inactivation of both pathways appears necessary to escape senescence (see discussion in Serrano et al. 1997). Consistent with this view, ectopic expression of p21 (a p53 target) and p16 can induce senescence in human cells (McConnell et al. 1998; Vogt et al. 1998).

The activation of p53 and p16 in response to deregulated MEK/MAPK signaling provides a relevant setting in which these tumor suppressors act to limit malignant transformation. In this view, mutations that deregulate the MAPK cascade provide an initial proliferative advantage but also accelerate senescence. However, cells acquiring mutations in p53 or at the INK4a/ARF locus escape senescence, thereby revealing the full mitogenic potential of the MAPK cascade. Consistent with this scenario, ras mutations often precede mutations in p53 or INK4a during tumor development (Fearon and Vogelstein 1990; Burns et al. 1991; Linardopoulos et al. 1995).

Premature senescence and tumor suppression

In summary, our data demonstrate that oncogenic Ras promotes premature senescence through activation of the MAPK cascade. Remarkably, this is the same pathway whereby Ras induces mitogenesis in immortal cells. Escape from Ras-induced senescence allows inappropriate proliferation and, in rodent cells, oncogenic transformation. Consequently, the biological outcome of constitutive MAPK activation—cell-cycle arrest or forced mitogenesis—is largely context-dependent and determined by the integrity of the senescence machinery. These data strongly support the notion that normal cells possess fail-safe mechanisms that limit the consequences of Ras mitogenic signaling. Because these safeguards involve p53 and p16, the activation of premature senescence in response to forced mitogenic signaling may be an important mechanism of tumor suppression.

Materials and methods

Cell culture

Normal diploid human IMR90 fibroblasts (early-mid passages) expressed the murine ecotropic receptor (Serrano et al. 1997), and primary mouse embryo fibroblasts derived from wild-type, p53−/− and INK4aex2−/− day 13.5 embryos were prepared as described previously (Jacks et al. 1994; Serrano et al. 1997). All cultures were maintained in Dulbecco’s modified Eagle medium (DMEM; Gibco) supplemented with 10% fetal bovine serum (FBS; Sigma) and 1% penicillin G/streptomycin sulfate (Sigma).

Retroviral vectors and gene transfer

Oncogenic Ras (H–RasV12) (Serrano et al. 1997) and Ras effector loop mutants (White et al. 1995; Jonesson et al. 1996) were expressed using the pBabe-Puro vector (Morgenstern and Land 1990). cDNA encoding Rac1 (N115I), RalA, RhoA (Q63L), R–Ras (G38V), TC21 (Q72L), PI(3)Kp110α (K227E), PI(3)Kp110–CAAX, wild-type Akt, or E1A12S, respectively, were cloned into a pLPC retroviral vector. Raf-1–CAAX was cloned into a Babe-Puro retroviral vector. MEK1356P (Bottorff et al. 1995) was expressed using Babc–Puro- or WZL–Hygro-based retroviral vectors (Ariad Pharmaceuticals, unpubl.). Retroviral gene transfer was performed as described (Serrano et al. 1997) using high-titer retroviral stocks generated by transient transfection of the Phoenix ecotropic packaging line (G. Nolan, Stanford University, CA). Infected cell populations were selected by culture in puromycin (2.5 µg/ml, 3 days) or in hygromycin (200 µg/ml, 5 days) to eliminate uninfected cells. Day 4 postinfection was...
designated as day 0 (Serrano et al. 1997). In all experiments, cells infected with an empty vector were used as control.

Analysis of cell proliferation

Cell proliferation was assessed by three criteria: growth curves, [3H]thymidine incorporation and flow cytometry of BrdU-labeled cells. For growth curves, cells were plated at a density of 2.5 x 10⁴ per well in 12-well-plates following drug selection. Cells were fixed in 10% formalin in the indicated times according to a time scheme and method described previously (Serrano et al. 1997). Crystal violet (0.1%; Sigma) was used to stain cells.

Cell-associated dye was extracted in 2 ml of 10% acetic acid and the optical density was measured at 590 nm with each value normalized to day 0. Each point was done in triplicate, each growth curve was performed at least twice. For experiments using PD98059 (Calbiochem), treatment was initiated 2 days postinfection. Cells were fed daily with fresh medium containing 50 µM PD98059 containing 0.25% DMSO. Solvent-containing medium was used to treat control cultures in parallel.

For [3H]thymidine incorporation assay, 2 x 10⁴ cells per well in 12-well-plates were plated in triplicate. Cells were pulsed for 20-24 hr or 4 hr with 5 µCi/ml methyl-3H-thymidine (Amersham, 2 Ci/m mole). After washing with PBS, cells were trypsinized and transferred to glass fiber filters (Whatman) followed by scintillation counting. For BrdU incorporation and DNA-content analysis, subconfluent cultures were labeled with 10 µM BrdU (Amersham) for 4 hr. Cells were trypsinized and fixed in 75% ethanol for 30 min. Cells were washed in PBS and were treated subsequently with 2 M HCl for 20 min at room temperature. After washing in washing buffer (0.1% BSA/PBS), cells were neutralized in 0.1 M sodium borate for 2 min followed by washing in washing buffer. Cells were incubated subsequently with FITC-conjugated anti-BrdU antibody (Pharmingen) for 1 hr at room temperature or overnight at 4°C. Finally, cells were washed in washing buffer three times followed by treatment with propidium iodide (10 µg/ml) and RNase A (100 µg/ml) at 37°C for 30 min. Samples were analyzed by two-dimensional flow cytometry to detect both fluorescein and propidium iodide.

Protein expression and activity

Western blot analysis was carried out as described previously with minor modifications (Serrano et al. 1997). Whole-cell lysates were derived by lysis of cell pellets in Laemmli SDS sample buffer. Samples corresponding to 30 µg of protein were resolved on SDS-PAGE gels and transferred to immobilone-P membranes (Millipore). The following antibodies were used: CM1 (Novocastra) at 1:1000 dilution; Santa Cruz) for 3 hr at 4°C. Afterwards, 25 µl of protein A–Sepharose beads (50%) were added to the mixture and were incubated for another 3 hr at 4°C. Samples were washed subsequently and resuspended in 20 µl of kinase buffer (25 mM Tris-HCl at pH 7.5, 10 mM MgCl₂, 5 mM β-glycerolphosphate, 2 mM DTT, 0.1 mM Na₃VO₄) supplemented with 10 µCi [γ-³²P]ATP (3000 Ci/m mole) and 5 µg of myelin basic protein (MBP; Sigma) and the reaction mixtures were incubated at 30°C for 30 min. The samples were resolved on 12% SDS-polyacrylamide gels and exposed to Kodak X-OMAT AR.

SA-β-galactosidase activity was detected as described previously (Dimri et al. 1995) with minor modifications. Cells were washed with PBS, fixed in 0.5% glutaraldehyde, and were incubated with staining solution (1 mg/ml 5-bromo-4-chloro-3-indolyl β-galactoside (X-gal), 5 mM potassium ferrocyanide, 5 mM potassium ferricyanide, 1 mM MgCl₂ in PBS at pH 6.0) for 12 hr at 37°C.

Transformation assays

For soft-agar assays, cells were resuspended in 0.3% Noble agar (in DMEM supplemented with 10% FBS) at a density of 2 x 10⁴ cells per well (in six-well plates), and were plated onto solidified bottom layer medium. Cultures were fed weekly and photomicrographs of colonies were taken 20–24 hr or 4 hr with 5 µCi/ml [methyl-3H]thymidine (Amersham). After staining with PBS, cells were trypsinized and transferred to glass fiber filters (Whatman) followed by scintillation counting. For BrdU incorporation and DNA-content analysis, subconfluent cultures were labeled with 10 µM BrdU (Amersham) for 4 hr. Cells were trypsinized and fixed in 75% ethanol for 30 min. Cells were washed in PBS and were treated subsequently with 2 M HCl for 20 min at room temperature. After washing in washing buffer (0.1% BSA/PBS), cells were neutralized in 0.1 M sodium borate for 2 min followed by washing in washing buffer. Cells were incubated subsequently with FITC-conjugated anti-BrdU antibody (Pharmingen) for 1 hr at room temperature or overnight at 4°C. Finally, cells were washed in washing buffer three times followed by treatment with propidium iodide (10 µg/ml) and RNase A (100 µg/ml) at 37°C for 30 min. Samples were analyzed by two-dimensional flow cytometry to detect both fluorescein and propidium iodide.

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For in vivo tumorigenicity assays, 10⁶ cells/0.25 ml of PBS were injected subcutaneously into male NSW athymic nude mice (Tatonic Farms). Each animal received two injections, one on each rear flank. Virus-infected MEFs from two different embryo preparations were used for each genotype, and at least two animals were examined for each cell population (i.e., four tumors). Tumor formation was monitored three times per week by palpation at the sites of injection. The length (L) and width (W) of tumors were measured using a caliper and the volume of tumor formed was determined using the formula V = (L x W²)/2.

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