Abstract. Studies in cell culture systems have indicated that oncogenic forms of Ras can affect apoptosis. Activating mutations of Ras occur in ~30% of all human tumors and 50% of colorectal carcinomas. Since these mutations appear at early or intermediate stages in multistep journeys to neoplasia, an effect on apoptosis may help determine whether initiated cells progress towards a more neoplastic state. We have tested the effects of K-rasVal12 on apoptosis in transgenic mice. A lineage-specific promoter was used to direct expression of human K-rasVal12, with or without wild-type (wt) or mutant SV-40 T antigens (TAg), in postmitotic villus enterocytes, the principal cell type of the small intestinal epithelium. Enterocytes can be induced to reenter the cell cycle by TAgWt. Reentry is dependent upon the ability of TAg to bind pRB and is associated with a p53-independent apoptosis. Analyses of K-rasVal12 × TAgWt bi-transgenic animals indicated that K-rasVal12 can enhance this apoptosis threefold but only in cycling cells; increased apoptosis does not occur when K-rasVal12 is expressed alone or with a TAg containing Glu107,Lys108 substitutions that block its ability to bind pRB. Analysis of bi-transgenic K-rasVal12 × TAgWt mice homozygous for wild-type or null p53 alleles established that the enhancement of apoptosis occurs through a p53-independent mechanism, is not attributable to augmented proliferation or to an increase in abortive cell cycle reentry (compared to TAgWt mice), and is not associated with detectable changes in the crypt–villus patterns of expression of apoptotic regulators (Bcl-2, Bcl-xL, Bak, and Bax) or mediators of epithelial cell–matrix interactions and survival (e.g., α5β1 integrin and its ligand, fibronectin). Coexpression of K-rasVal12 and TAgWt produces dysplasia. The K-rasVal12-augmented apoptosis is unrelated to this dysplasia; enhanced apoptosis is also observed in cycling nondysplastic enterocytes that produce K-rasVal12 and a TAg with a COOH-terminal truncation. The dysplastic epithelium of K-rasVal12 × TAgWt mice does not develop neoplasms. Our results are consistent with this finding: (a) When expressed in initiated enterocytes with a proliferative abnormality, K-rasVal12 facilitates progression to a dysplastic phenotype; (b) by diminishing cell survival on the villus, the oncoprotein may impede further progression; and (c) additional mutations may be needed to suppress this proapoptotic response to K-rasVal12.

Activating mutations of Ras genes are encountered in ~30% of all human tumors and in 50% of colorectal carcinomas (Bos et al., 1987; Forrester et al., 1987; Fearon and Vogelstein, 1990). These mutations have been generally described as early or intermediate events in multistep journeys to neoplasia (Barbacid, 1987; Fearon and Vogelstein, 1990). These mutations have been generally described as early or intermediate events in multistep journeys to neoplasia (Barbacid, 1987; Fearon and Vogelstein, 1990). Studies of cultured cells have shown that oncogenic forms of Ras can provoke premature senescence or influence apoptosis—two responses that can have an important impact on tumorigenesis.

1. Abbreviations used in this paper: BrdU; 5-bromo-2′-deoxyuridine; cdk, cyclin-dependent kinase; Cy3, indocarbocyanine; Fabpi, intestinal fatty acid–binding protein gene; FITC, fluorescein isothiocyanate; P, postnatal day; SV-40 TAg, simian virus 40 large T antigen; TUNEL, terminal deoxynucleotidyltransferase (TdT)-mediated, dUTP nick end labeling.

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great (Serrano et al., 1997; for review see Weinberg, 1997). p53 and/or p16 mutations are encountered in most human
tumors (Hollstein et al., 1994; Hiraoka and Koettler, 1995).

Senescence can be viewed as a protective, antineoplastic “host” response to an initial activating Ras mutation, a re-
sponse that can be overridden by subsequent inactivating
mutations of either p16 or p53, thereby allowing progress-
ion (transformation). This view fits with the multistep
model of human colorectal carcinoma, where early muta-
tion of Ras is followed by p53 inactivation (Fearon and
Vogelstein, 1990). The findings of Serrano and co-workers
also provide a rationale for two well-known phenomena:

(a) Primary cultures of embryonic cells are resistant to
transformation by onecogenic forms of Ras (Newbold and
Overell, 1983), while immortalized cells, which have often
acquired inactivating p16 or p53 mutations, are very sensi-
tive to transformation. (b) Oncogenic forms of Ras coop-
erate with viral oncoproteins, such as E1A or SV-40 large T antigen (SV-40 Tag), to transform primary cell cultures.
Both of these viral proteins bind to and disable pRB, pro-
ducing an effect functionally analogous to p16 inactiva-
tion, since pRB is the principal substrate for phosphoryla-
tion by cdk4 and cdk6 (Weinberg, 1995, 1997). In the case of
SV-40 Tag, both pRB and p53 are bound and inacti-
ated.

Apoptosis represents another response that initiated
cells may marshal to prevent their amplification. Studies of
various established cultured cell lines have revealed that
oncogenic forms of Ras can affect apoptosis. Promotion as
well as suppression of cell death has been reported (Fer-
nandez et al., 1994; Chen and Faller, 1995; Kinoshita et al.,
1995; Lin et al., 1995; Rak et al., 1995; Wang et al., 1995).
Promotion of apoptosis in initiated cells should impede
their progression, while inhibition of apoptosis would be
expected to help facilitate tumorogenesis.

The effect of Ras oncoproteins on apoptosis or senes-
cence has not been examined using genetically defined in
vivo models. The impact of a lineage’s state of differen-
tiation and/or proliferative status on Ras-modulated demise
or survival can be assessed in transgenic mice where the
activated oncoprotein can be directed to selected target
populations. The self-renewing small intestinal epithe-
lum of the adult mouse represents an attractive system for
conducting such an analysis. Its attractiveness stems from
the fact that proliferation, differentiation, and death pro-
grams are continuously expressed in spatially well-defined
domains of crypt–villus units. Normally, these decisions are
precisely coordinated to maintain cellular census. Pro-
liferation is limited to crypts of Lieberkühn. Each crypt con-
tains one or more active multipotent stem cell located near
its base (Loeffler et al., 1993). The stem cell gives rise to
four principal epithelial cell lineages that complete their
terminal differentiation during a highly organized, rapid
migration. Postmitotic absorptive enterocytes, mucus-prod-
cucing goblet cells, and enteroendocrine cells exit the
crypt and move up an adjacent villus in vertical coherent
columns (Schmidt et al., 1985; Hermiston et al., 1996). Cells
are removed from the villus tip by apoptosis and/or extru-
sion into the lumen (Hall et al., 1994). The entire sequence
is completed in 3–5 d (Cheng, 1974a; Cheng and Leblond,
1974a–c). In contrast, members of the Paneth cell lineage
undergo terminal differentiation as they move down to the
base of the crypt, where they are subsequently removed after an ~20-d residence (Cheng, 1974b). The availability of
lineage-specific promoters that function at selected lo-
cations along the crypt–villus units (e.g., Cohn et al., 1992;
Bry et al., 1994; Simon et al., 1995, 1997) makes it possible
to introduce activated Ras oncoproteins in specified cellular
contexts and explore their effects on survival or death.

In the present report, we have used transgenic mice to
examine the effects of expressing human K-ras
Val12 in postmitotic villus enterocytes. K-ras
Val12 was also coexpressed with wild-type or mutant SV-40 Tag’s that affect the prolifera-
tive status of these cells. The results disclose that
K-ras
Val12 can promote a p53-independent apoptosis that
occurs when enterocytes undergo pRB-related reentry into the cell cycle.

Materials and Methods

Production and Maintenance of Transgenic Mice

FVB/N transgenic mice containing nucleotides −1178 to +28 of the rat
intestinal fatty acid–binding protein gene (Fabpi) linked to human
K-ras
Val12, wild type (Wt) SV-40 Tag, SV-40 Tag
Val12 (mutant 3213 in Chen et al., 1992), or SV-40 Tag
Val12 to 708 (mutant d1137 in Pipas et al., 1983) were
produced as described in earlier reports (Kim et al., 1993, 1994; Chan-
drasekaran et al., 1996). Transgenics were maintained by crosses to
normal FVB/N littermates. Male and female C57Bl/6 p53+/− and p53−/−
animals were purchased from GenPharm International (Mountain View, CA).

FVB/N Fabpi–K-ras
Val12 transgenic mice were crossed to FVB/N
Fabpi–SV-40 Tag
Val12, Fabpi–SV-40 Tag
Val12, or Fabpi–SV-40 Tag
Val12 to 708
animals to generate bi-transgenic mice. Animals were genotyped using tail
DNA and PCR protocols detailed elsewhere (Kim et al., 1993, 1994;
Chandrasekaran et al., 1996).

FVB/N normal, Fabpi–K-ras
Val12, Fabpi–SV-40 Tag
Val12, and Fabpi–K-ras
Val12 × Fabpi–SV-40 Tag
Val12 transgenic mice homozygous for p53 wild-type or null alleles were generated by a series of crosses to C57Bl/6 p53+/− and C57Bl/6 p53−/−
animals.

Mice were maintained in microisolator cages under a strict light cycle
(lights on at 0600 h and off at 1800 h). Animals were given a standard irriga-
ted chow diet (Pico rodent chow 20; Purina Mills, Inc., St. Louis, MO) ad
libitum. Routine screens for Hepatitis, Minute, Lymphocytic Chorio-
meningitis, Ectromelia, Polyoma, Sendai, Pneumonia, and mouse adeno-
viruses, enteric bacterial pathogens, and parasites were negative.

Characterization of Transgenic Mice

Western Blots. Animals were sacrificed, and their gastrointestinal tracts
were removed en bloc. The small intestine was subdivided into thirds (de-
finied as duodenum, jejunum, and ileum). The distal half of the middle seg-
ment (“distal jejunum”) was snap frozen in liquid N2, lyophilized, pulver-
ized, and then resuspended in extraction buffer (50 μg lyophilized tissue/ ml buffer; extraction buffer: extraction buffer = 40 mM Tris, pH 6.8, 2% 2-mercaptoethanol, 1% SDS, 5% glycerol, 10 mM EDTA, 50 μg/ml aprotinin [Sigma Chemi-
cal Co., St. Louis, MO], 50 μg/ml leupeptin [Sigma Chemical Co.], 50 μg/ml Pefabloc [Boehringer Mannheim Corp., Indianapolis, IN], and 10 μg/ml pepstatin A [Sigma Chemical Co.]). Samples were boiled for 5 min, and
insoluble debris was removed by centrifugation for 3 min at 12,000 g. The protein concentration of the cleared supernatant was determined using the
d DC Protein Assay kit (BioRad Labs., Hercules, CA). Extracted cellu-
lar proteins (75 μg/sample) were fractionated by SDS-PAGE (Laemmli, 1970) and electrophoretically transferred to polyvinylidene difluoro-
membranes (Amersham Corp., Arlington Heights, IL). Membranes were
stained with Ponceau red to verify equivalent transfer of all samples. Blots
were pretreated in PBS-blocking buffer (1% gelatin, 0.2% Tween-20 in PBS)
for 1 h at 23°C and then incubated in blocking buffer for 2 h at 23°C with:
(a) affinity purified rabbit anti-Bcl-2 raised against a peptide span-
ings residues 4–21 of the mouse protein (Santa Cruz Biochemicals, Santa
Cruz, CA; final dilution in blocking buffer = 1:500); (b) affinity-purified
rabbit anti-Bax raised against amino acids 11–30 of the mouse protein
(Santa Cruz Biochemicals; 1:500); (c) two preparations of affinity-purified
antibodies raised in rabbits against a peptide spanning residues 43–61 of

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mouse Bax (obtained from Santa Cruz Biochemicals and from Stanley Korsmeyer, Washington University School of Medicine; 1:500); (d) affinity-purified rabbit antibodies to a peptide corresponding to amino acid residues 2–19 of mouse Bcl-x (Santa Cruz Biochemicals; 1:500); (e) affinity-purified rabbit antibodies to residues 629–648 of mouse c-Raf-1 (Santa Cruz Biochemicals; 1:500); (f) rabbit anti-mouse cdk4 (Santa Cruz Biochemicals; 1:1,000); (g) rabbit anti-mouse cdk2 (Santa Cruz Biochemicals; 1:1,000); (h) mouse SV-40 Tag mAbs (1:1,000; PharMingen, San Diego, CA); (i) rabbit anti-actin (Sigma Chemical Co.; 1:5,000); and (j) rabbit anti-rat liver fatty acid binding protein (L-Fabp; Sweetser et al., 1998; 1:7,000). Anti- gen–antibody complexes were visualized with alkaline phosphatase-conju- gated secondary antibodies and a cholineminesesue substrate using the Western Light® kit (Tropix, Inc., Bedford, MA). All experiments were performed using three to eight animals/genotype and repeated on two to six occasions.

Blocking controls were performed by preincubating antibodies with a 10-fold weight excess of their corresponding peptide antigens for 1 h at 23°C before probing the blots. In addition, the specificity of the Bax anti- body preparations was established by probing Western blots containing proteins extracted from the distal jejunum, spleen, and thymus of mice that were homozygous for wild-type or null alleles of the Bax gene (tissues obtained from S. Korsmeyer; Knudson et al., 1995).

**Immunohistochemical Analysis of the Expression of Regulators of the G1/S Transition and of Apoptosis**

Postnatal day 42 to 60 (P42–60) male and female FVB/N mice containing one or two transgenes and their normal littermates (n = 10 animals/geno- type) were given an intraperitoneal injection of 5-bromo-2-deoxyuridine (BrdU; Sigma Chemical Co.; 120 mg/kg body weight) and 5-fluoro-2-deoxy- uridine (5-FdUrd; Sigma Chemical Co.; 12 mg/kg body weight) 90 min before sacrifice to label cells in S-phase. The small intestine was removed, flushed with PBS, fixed in Bouin’s solution for 6–12 h, washed with 70% ethanol, and then divided into proximal and distal halves. Each half was opened with an incision along its cephalicaudal axis and then rolled from its proximal to distal end. Each of the resulting “Swiss rolls” was then cut in half parallel to the duodenal–ileal axis and placed in a tissue cassette with the cut edge of one half facing up and the cut edge of the other half facing up. All sections were placed in paraffin, serial 5-μm sections were deparaffinized in xylene and isopropanol and were then stained with hematoxylin and eosin or with antibodies to cyclin D1 (Upstate Biotechnology, Inc., Lake Placid, NY; 1:1,000); (a) rabbit anti–mouse cyclin D1 (Upstate Biotechnology, Inc., Lake Placid, NY; 1:1,000); (b) rabbit anti–mouse cyclin E (Santa Cruz Biochemicals; 1:500); (c) anti–mouse cdk2 (see above; 1:500); (d) mouse mAb against subunit I of human cytochrome oxidase (5D4-F5; Molecular Probes, Eugene, OR; 1:100); (e) goat anti–rabbit Ig (Cohn and Lieberman, 1984; 1:1,000); (f) mouse mAb against subunit I of human cytochrome oxidase (5D4-F5; Molecular Probes, Eugene, OR; 1:100); and (m) rabbit anti–AS-40 Tag (obtained from Doug Hanahan, University of California, San Francisco, CA; 1:1,000). Formalin-fixed tissues were used to identify Bcl-2, Bax, Bak, Bcl-x, Bcl-a2, p21, p27, CDK4, CDK2, and transgenic mice (2–4 animals/group) using the following panel of antibodies: (a) rat anti–mouse β6 integrin mAb (clone 9E7; PharMingen, San Diego, CA; diluted 1:2,000); (b) rabbit anti–mouse α6 integrin (Chemicon, Temecula, CA; 1:1,000); (c) rabbit anti–mouse α5 integrin (Chemicon; 1:2,000); (d) rat anti–human/mouse α5 mAb (clone GoH3; PharMingen; 1:4,000) (e) rabbit anti–mouse β3 mAb (clone 346-11A; PharMingen; 1:2,000); (f) rabbit anti–mouse fibronectin (Chemicon; 1:5,000); (g) rabbit anti–mouse laminin (recognizes both A and B laminin isoforms; Chemicon; 1:15,000); and (h) rabbit anti–mouse type IV collagen (Chemi- con; 1:5,000).

Immunohistochemical Analysis of the Crypt–Villus Pattern of Expression of Integrins and Their Ligands Using Tyramide Signal Amplification

The spatial distribution of these proteins was examined in P42–60 normal and transgenic mice (n = 2–4 animals/group) using the following panel of antibodies: (a) rat anti–mouse β6 integrin mAb (clone 9E7; PharMingen, San Diego, CA; diluted 1:2,000); (b) rabbit anti–mouse α6 integrin (Chemicon, Temecula, CA; 1:1,000); (c) rabbit anti–mouse α5 integrin (Chemicon; 1:2,000); (d) rat anti–human/mouse α5 mAb (clone GoH3; PharMingen; 1:4,000) (e) rabbit anti–mouse β3 mAb (clone 346-11A; PharMingen; 1:2,000); (f) rabbit anti–mouse fibronectin (Chemicon; 1:5,000); (g) rabbit anti–mouse laminin (recognizes both A and B laminin isoforms; Chemicon; 1:15,000); and (h) rabbit anti–mouse type IV collagen (Chemini- con; 1:5,000).

Mice were sacrificed, and the middle third of the small bowel was flushed with PBS and then frozen in OCT (Miles, Inc., Kanakee, IL). Serial 5-μm-thick sections were cut, fixed in methanol (–20°C for 15 min), washed three times (3 min/cycle) in PBS, and treated with PBS-blocking buffer for 15 min at room temperature. Cells with endogenous peroxidase activity were labeled by incubating the frozen sections for 10 min at room temperature with FITC-conjugated biotinyl-tyramide (New England Nu- clear Life Sciences, Boston, MA; diluted 1:100 in 1× amplification diluent from the same manufacturer). After three washes in PBS (5 min each), the sections were incubated overnight at 4°C with one of the primary antibod- ies listed above and then washed in TNT buffer (0.1 M Tris, pH 7.5; 0.15 M NaCl/0.05% Tween 20; 3 cycles; 5 min each). HRP-conjugated goat anti- rat or goat anti–rabbit Ig were added (Kirkegaard and Perry Labs, Gaith- ersburg, MD; diluted 1:100 in TNT buffer [0.1 M Tris, pH 7.5; 0.15 M NaCl/0.5% blocking reagent from New England Nuclear Life Sciences]). After a 30-min incubation at room temperature, sections were washed three times with TNT buffer (5 min each). Biotinyl-tyramide was added (diluted 1:100 in 1× amplification diluent) for 10 min. Sections were washed three times in TNT buffer (5 min/cycle), subsequently overlaid with Cy3-conju- gated streptavidin (New England Nuclear Life Sciences; diluted 1:500 in TNB) for 30 min, and then subjected to three more rinses in TNT buffer before coverslips were added.

Two controls were performed for each experiment using a given primary antiserum: direct amplification of endogenous peroxidase activity alone (see above) and omission of primary antibodies.

**Quantitation of Apoptosis**

Terminal deoxynucleotidyltransferase (TdT)-mediated, dUTP nick end labeling (TUNEL) assays were performed on sections of formalin-fixed Swiss rolls using the protocol of Gavrieli et al. (1992), except that sections were incubated with proteinase K (Boehringer Mannheim Corp.; 20 μg/ml) for 20 min at 23°C. Incorporation of digoxigenin-labeled dUTP was de- tected using peroxidase-conjugated sheep anti-digoxigenin Fab fragments (Boehringer Mannheim Corp.; diluted 1:500 in PBS-blocking buffer) and the Vector VIP kit (Vector Laboratories, Burlingame, CA). Sections were counterstained in methyl green (Zymed Labs, S. San Francisco, CA).

The number of TUNEL-positive epithelial cells with apoptotic morpho- phy (Wyllie et al., 1980, Hall et al., 1994) were scored in a blinded fashion by two observers. All intact crypt–villus units present in two non-adjacent sections of a Swiss roll prepared from the distal half of the small intestine were surveyed (n = 3–5 animals/genotype). Statistical analyses were performed using Student’s t test (SigmaPlot; Jandel Corp., San Rafael, CA).

**Quantitation of the Ratio of S-phase to M-phase Cells**

The small intestine was removed from single transgene-containing and bi- transgenic mice that had been pulse-labeled 90 min before sacrifice with BrdU. Jejunal segments were fixed in Bouin’s, embedded in paraffin, and serial 5-μm sections were stained with hematoxylin and eosin or with anti- bodies to BrdU. M-phase and S-phase cells in crypts and villi were

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counted in adjacent serial sections. 40 well-oriented crypt–villus units were scored per section (n = 2 sections/mouse; 3 mice/genotype).

Measurements of K-rasVal12 mRNA Levels

Total cellular RNA was isolated from the jejunum of single transgene-containing animals, bi-transgenic mice, and normal littermates. K-rasVal12 mRNA levels were defined using a ribonuclease protection assay (Kim et al., 1993). The 199 base cRNA yields a 148 base protected fragment when annealed to the mRNA product of Fabpi–K-rasVal12 and a 54 base protected fragment when annealed to the mRNA product of the endogenous mouse c-K-rasGly12 gene.

Results

Studies of Bi-transgenic Mice Indicate That K-rasVal12 Augments the p53-independent Apoptosis Produced by SV-40 TAgWt in Villus Enterocytes

We examined the effect of an activated Ras, human K-rasVal12, on apoptosis in villus enterocytes of transgenic mice. K-rasVal12 was selected because it is the most common activated Ras found in intestinal neoplasms (Bos et al., 1987; Forrester et al., 1987).

Studies in transgenic mice have shown that nucleotides −1178 to +28 of the rat intestinal fatty acid binding protein gene (Fabpi) can be used to restrict expression of foreign gene products to postmitotic villus enterocytes distributed along the length of the small intestine (e.g., Cohn et al., 1992; Kim et al., 1993; Hermiston et al., 1996). Transgene expression is activated just as enterocytes emerge from the crypt and is sustained as they complete their upward migration to the villus' apical extrusion zone. Fabpi–reporter transgenes are first expressed coincident with the initial cytodifferentiation of the intestinal epithelium on embryonic day 15. Expression is maintained for at least the first 2 yr of life.

Ribonuclease protection assays indicated that like other Fabpi2–1178 to 128-containing transgenes, Fabpi–K-rasVal12 is expressed at highest levels in the middle third of the small intestine (jejunum) of 6–8-wk-old FVB/N mice (data not shown).

Figure 1. Quantitative analysis of apoptosis in the jejunal villus epithelium of adult transgenic and bi-transgenic mice homozygous for p53 wild-type or null alleles. (A) Postnatal day (P) 42 p53+/+ FVB/N transgenic mice. (B) P42 p53−/− animals with equivalent genetic backgrounds. An asterisk indicates that the results obtained for animals of this genotype are significantly different from normal littermate controls (P < 0.05). Crypt apoptosis was not significantly different between normal nontransgenic animals and any of the single transgene-containing or bi-transgenic mice (data not shown).

Figure 2. Expression of cdk2 in FVB/N normal, transgenic, and bi-transgenic mice. (A) Western blot of jejunal protein extracts (75 μg/lane) prepared from P42 mice, probed with affinity-purified antibodies to cdk2. Lane 1, nontransgenic FVB/N mouse; lane 2, Fabpi–K-rasVal12 animal; lane 3, Fabpi–SV-40 TAgWt mouse; lane 4, K-rasVal12 × SV-40 TAgWt bi-transgenic animal; lane 5, K-rasVal12 × SV-40 TAgWt bi-transgenic mouse. (B–D) Expression of cdk2 along jejunal crypt–villus units of a P42 normal FVB/N mouse (B), a Fabpi–SV-40 TAgWt transgenic littermate (C), and a P42 K-rasVal12 × SV-40 TAgWt bi-transgenic mouse (D). Cdk2 was detected by incubating the tissue sections with the same rabbit anti–mouse cdk2 preparation used for the Western blotting experiment shown in A and Cy3-conjugated donkey anti–rabbit Ig. Cdk2 (red) is confined to proliferating crypt epithelial cells in the normal FVB/N mouse. (The closed arrows in B point to a crypt–villus junction; the open arrow points to a crypt base columnar epithelial cell. These cells represent proliferating crypt epithelial cells.)
ating precursors of the enterocytic lineage and are intermixed with differentiated Paneth cells at the base of crypts [Cheng and Leblond, 1974a]. SV-40 TAgWt expression is associated with an induction of cdk2 in villus enterocytes (e.g., C, arrow). A similar induction of cdk2 in villus enterocytes is evident in the bi-transgenic animal (D). (E–G) Multilabel immunohistochemical study of a jejunal crypt–villus unit from the bi-transgenic mouse shown in D, establishing that villus enterocytes have reentered the cell cycle. The mouse received an intraperitoneal injection of BrdU 1.5 h before sacrifice. The section was stained with rabbit anti–SV-40 TAg (visualized in E with Cy3-donkey anti–rabbit Ig) and goat anti-BrdU (visualized in F with FITC-conjugated donkey anti–goat Ig). G is a dual exposure of the section confirming that BrdU-positive cells distributed along the length of the villus also express SV-40 TAgWt (yellow-orange staining nuclei). The closed arrows in E–G indicate the location of the crypt–villus junction. The open arrows in G point to proliferating crypt epithelial cells that do not express SV-40 TAgWt. (Fabpi–directed expression of SV-40 TAgWt is limited to villus enterocytes.) Bar, 25 μm.
in jejunal villus enterocytes produces no statistically significant change in their basal level of apoptosis (Fig. 1 A). This is true whether animals are p53+/+ or p53−/− (Fig. 1, A and B). Apoptotic cells are limited to the tips of villi in both Fabpi−K-rasVal12 animals and their nontransgenic FVB/N cage mates.

Forced expression of K-rasVal12 has no detectable effect on the pattern of accumulation of regulators of the G1/S transition of the cell cycle. Terminal differentiation of normal FVB/N enterocytes is associated with rapid loss of cyclin D1 and cdk2 but not their partners, cdk4 and cyclin E. Cellular levels of pRB do not change as enterocytes move from the base to the tips of villi, although pRB phosphorylation appears to diminish (Chandrasekaran et al., 1996). Immunohistochemical and Western blot analyses of normal and transgenic mouse intestine revealed that K-rasVal12 does not affect expression of these cell cycle regulators (Fig. 2 A, plus data not shown), nor does it cause villus enterocytes to reenter the cell cycle, whether judged by their ability to incorporate BrdU or by the appearance of M-phase cells (data not shown).

Other markers of terminal differentiation of the enterocyte lineage, including a complex pattern of glycoconjugate production defined by in situ binding of members of a panel of lectins to fixed sections of intestine (Falk et al., 1994; Hermiston and Gordon, 1995), are unaffected by K-rasVal12 (data not shown).

Fabpi-directed expression of SV-40 TAg in villus enterocytes induces cyclin D1 and cdk2 expression and causes cell cycle reentry (Chandrasekaran et al., 1996). SV-40 TAg−/−-induced reentry is accompanied by a statistically significant threefold increase in villus epithelial cell apoptosis (P < 0.05; Fig. 1 A). Apoptotic cells are present in the lower and mid-portions of Fabpi−SV-40 TAg−/− villi, regions where apoptotic events are undetectable in normal FVB/N littermates (data not shown).

Crossing FVB/N Fabpi−K-rasVal12 mice to FVB/N Fabpi−SV-40 TAg−/− mice yields bi-transgenic animals with a proliferating and dysplastic population of enterocytes. The dysplasia is manifested in part by branched villi (Fig. 3 A). K-rasVal12 augments the apoptosis observed in SV-40 TAg−/−-positive villus enterocytes; bi-transgenic mice have a statistically significant threefold increase in apoptotic cells in jejunal villi compared to Fabpi−SV-40 TAg−/− transgenics, and a statistically significant six- to ninefold increase when compared to mice containing Fabpi−K-rasVal12 alone or no transgenes (Fig. 1 A). Apoptotic cells are evident in the lower and middle thirds of villi and occur in regions with and without dysplasia (Fig. 3, A–D).

The augmented apoptosis in bi-transgenic mice cannot be attributed to alterations in K-rasVal12 or SV-40 TAg expression. Ribonuclease protection assays indicated that K-rasVal12 mRNA levels are equivalent in jejunal RNAs isolated from animals containing Fabpi−K-rasVal12 alone or Fabpi−K-rasVal12 and Fabpi−SV-40 TAg−/−. SV-40 TAg−/− mRNA levels are also indistinguishable in Fabpi−SV-40 TAg−/− transgenic and K-rasVal12 × SV-40 TAg−/− bi-transgenic animals (data not shown).

The steady-state levels plus intracellular and crypt–villus distributions of pRB, cyclin D1, cyclin E, cdk4, and cdk2 are the same in Fabpi−SV-40 TAg−/− and K-rasVal12 × SV-40 TAg−/− animals. In addition, there is no statistically significant difference in the ratio of S-phase to M-phase cells in the jejunal villi of bi-transgenic K-rasVal12 × SV-40 TAg−/− mice compared to their Fabpi−SV-40 TAg−/− littermates (Fig. 2, A–G, plus data not shown). These results indicate that the K-rasVal12 effect on apoptosis is not due to increases in either proliferation or abortive cell cycle reentry.

K-rasVal12 Can Only Augment Apoptosis When SV-40 TAg Is Able to Produce Cell Cycle Reentry

SV-40 TAgK107/8 is a mutant with Lys for Glu substitutions at residues 107 and 108. These substitutions block the ability of TAg to bind pRB and the related pocket proteins p107 and p130 but do affect p53 binding (Fanning and Knippers, 1992). (p107 and p130 are not detectable in the small intestinal epithelium of adult FVB/N mice using commercially available antibodies and a variety of sensitive immunohistochemical detection methods [Chandrasekaran et al., 1996].) We analyzed adult (6–8-wk-old) FVB/N Fabpi−SV-40 TAgK107/8 transgenic mice with equivalent concentrations of TAg in their total jejunal protein extracts as our Fabpi−SV-40 TAg−/− animals (Chandrasekaran et al., 1996). Expression of the mutant TAg in villus enterocytes does not induce cyclin D1 or cdk2, does not cause reentry into the cell cycle, and does not cause any significant increase in apoptosis compared to normal littermate controls (Fig. 1 A, plus data not shown). Coexpression of K-rasVal12 and SV-40 TAgK107/8 in bi-transgenic mice does not change the growth-arrested state of villus enterocytes, as judged by their lack of incorporation of BrdU, by the lack of cells in M-phase, and by the lack of detectable changes in the steady-state levels or crypt–villus distributions of pRB, cyclins D1 and E, and their partners cdk4 and cdk2 (e.g., compare lanes 1 and 5 in Fig. 2 A). There are no dysplastic changes in the small intestinal epithelium of these animals (n = 15). Moreover, K-rasVal12 × SV-40 TAgK107/8 mice do not manifest any significant increases in villus apoptosis when compared to nontransgenic FVB/N animals or FVB/N mice containing either one of the transgenes alone (Fig. 1 A).

p53 Is Not Required

A comparison of the phenotypes of K-rasVal12 × SV-40 TAgK107/8 and K-rasVal12 × SV-40 TAg−/− mice establishes a linkage between pRB and the observed apoptotic response of villus enterocytes. It also indicates that functional domains of the viral oncoprotein, other than its pRB binding region, are not sufficient to induce death in villus enterocytes or to mediate the further enhancement of apoptosis by K-rasVal12. An analysis of cell death in the jejunal crypt–villus units of 6–8-wk-old p53+/+ and p53−/− K-rasVal12 × SV-40 TAg−/− mice provided definitive proof that K-rasVal12 augments apoptosis through a p53-independent pathway (Fig. 1, A and B).

The Presence or Absence of Dysplasia Has Little Effect on the Ability of K-ras to Increase Apoptosis in Cycling Villus Enterocytes

SV-40 TAgVal12 to 708 is a truncation mutant that contains the NH2-terminal 121 residues of the viral oncoprotein. It retains the ability to bind pRB and related pocket proteins
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but does not interact with p53 (Srinivasan et al., 1989). We examined members of a pedigree of FVB/N Fabpi–SV-40 TAgD122 to 708 mice whose jejunal TAg levels were equivalent to those observed in FVB/N Fabpi–SV-40 TAgWt and Fabpi–SV-40 TAgK107/8 animals. Cdk2 is induced in the villus enterocytes of Fabpi–SV-40 TAgD122 to 708 mice. Reentry into the cell cycle occurs, although the proliferative response is less than what is observed with SV-40 TAgWt (e.g., the steady-state concentrations of cdk2 in jejunal protein extracts are 50–60% of those in age-matched Fabpi–SV-40 TAgWt animals; n = 3 mice/genotype). Coexpression of K-rasVal12 and the truncation mutant does not produce dysplasia in jejunal villi (Kim et al., 1994). Nonetheless, coexpression results in a statistically significant (P < 0.05) 2.3-fold increase in apoptosis compared to age-matched Fabpi–SV-40 TAgD122 to 708 cagemates (Fig. 1 A).

Coexpression of K-rasVal12 and SV-40 TAgWt Has No Appreciable Effect on the Crypt–Villus Distribution or Steady-State Levels of Bcl-2, Bcl-xL, Bak, Bax, and c-Raf-1

Previous studies of the effect of Ras oncoproteins on apoptosis in a variety of cultured cell lines implicated Bcl-2 and/or p53 as mediators (Fernandez-Sarabia and Bischoff, 1993; Tanaka et al., 1994; Chen and Faller, 1995, 1996; Kinosita et al., 1995; Lin et al., 1995; Wang et al., 1995). Therefore, the distribution of several regulators of apoptosis were analyzed along the jejunal crypt–villus units of normal FVB/N mice and in littermates containing all combinations of the transgenes described above. Bak (bcl-2 homologous antagonist/killer) is proapoptotic (Chittenden et al., 1995; Farrow et al., 1995; Kiefer et al., 1995). Bcl-2 is
an antiapoptotic (Bakhshi et al., 1985; Cleary and Sklar, 1985; Tsujimoto et al., 1985). Alternative splicing of the Bcl-x gene gives rise to Bcl-xL, which is antiapoptotic, and Bcl-xS, which is proapoptotic (Boise et al., 1993; N.B. Western blots indicate that Bcl-xL represents >95% of total jejunal Bcl-x in normal and bi-transgenic animals; data not shown).

Differentiation of the enterocytic lineage is normally associated with an upregulation of Bcl-2 and Bcl-xL without appreciable expression of Bak. The increased apoptosis seen along the length of SV-40 TAgWt, K-rasVal12 × SV-40 TAgWt, and K-rasVal12 × SV-40 TAg Val12 to 70% jejunal villi is not associated with any remarkable change in the cellular or intracellular patterns of accumulation of these proteins.

Figure 4. Cellular and intracellular distributions of Bak, Bcl-2, Bcl-xL, and c-Raf-1 in jejunal crypt–villus units of P42 bi-transgenic K-rasVal12 × SV-40 TAgWt mice. (A) Section of a crypt stained with mouse monoclonal antibodies to Bak and Cy3-sheep anti–mouse Ig. The open arrow point to a crypt–villus junction. The apical surfaces of crypt epithelial cells are intensely stained (closed arrow). Bak falls to undetectable levels as postmitotic enterocytes emerge from the crypt and enter the lower quarter of the villus. The open arrowhead points to one of several lamina propria lymphocytes that react with secondary Cy3-conjugated sheep antibodies to mouse Ig. (B–F) Distribution of the antiapoptotic mediator Bcl-2. (B) Section of a bi-transgenic jejunal villus stained with affinity-purified rabbit anti–Bcl-2 plus Cy3-donkey anti–rabbit Ig. (C) Same section stained with mouse mAbs to subunit I of cytochrome oxidase and FluorX-conjugated donkey anti–mouse Ig. (D) Blocking control in which the primary antibody was incubated with the peptide antigen from which it was generated, before application to another jejunal crypt–villus unit. (E) Same section as shown in B and C, stained with mouse mAbs against cytochrome oxidase subunit I, a mitochondrial protein, and FluorX-conjugated donkey anti–mouse Ig. (F) A triple exposure of C and E. Bcl-2 is barely detectable in the central portion of crypts where the proliferating transit cell population resides or at the crypt base were Paneth cells are located (data not shown). Cellular levels rise markedly just as enterocytes exit the crypt. There is no appreciable change in steady-state concentration or intracellular distribution of Bcl-2 as enterocytes complete their migration up the villus. The distribution of Bcl-2 within enterocytes is similar to the distribution of the mitochondrial marker, consistent with its previously described organelle affiliation in other cell lineages (Hockenbery et al., 1990). (G–I) Antibodies to Bcl-x produce a crypt–villus and intracellular pattern of staining that mimics that of Bcl-2. G is a section of a bi-transgenic jejunal villus incubated with affinity-purified rabbit anti–Bcl-x, Cy3-donkey anti–rabbit Ig, and bis-benzimide. H shows the same section stained with mouse mAbs to subunit I of cytochrome oxidase and FluorX-conjugated donkey anti–mouse Ig. I is a triple exposure. (J) Jejunal villus from a bi-transgenic mouse stained with affinity-purified rabbit anti–c-Raf-1 and Cy3-donkey anti–rabbit Ig. c-Raf is expressed in epithelial cells distributed throughout the length of jejunal crypt–villus units. Cellular levels do not change appreciably as cells complete their migration-associated terminal differentiation. Immunoreactive protein is absent from the nucleus and present throughout the cytoplasm, where it produces a granular pattern of staining. Bars, 25 μm.
Bak remains confined to the crypt and is not induced in cycling villus enterocytes (Fig. 4 A). Bcl-2 remains undetectable or barely detectable in crypt epithelial cells, continues to be induced at the crypt–villus junction, and is not suppressed as cells move to the villus tip, even though they are dividing (Fig. 4, B–I). The crypt–villus and intracellular distributions of Bcl-xL are similar to those of Bcl-2 in all cases (e.g., compare G–I with C, E, and F in Fig. 4). Western blots of jejunal extracts prepared from transgenic and bi-transgenic mice established that expression of K-rasVal12, SV-40 TAgWt, SV-40 TAgK107/8, or SV-40 TAgA122 to 708 alone, or K-rasVal12 plus the various SV-40 TAg s, has no appreciable effects on the steady state levels of Bcl-2, Bcl-xL, or the proapoptotic mediator Bax (Olsvai et al., 1993) (Fig. 5).2

Recent studies of cultured cells suggest that c-Raf-1, a serine-threonine kinase that plays a central role in the mitogen-activated kinase signaling pathway, may also operate downstream of Ras and upstream of Bcl-2 to promote death (Wang et al., 1994; Blagosklonny et al., 1996). Coexpression of K-rasVal12 and SV-40 TAgWt does not perturb the cellular patterns of c-Raf-1 accumulation (Fig. 4 J) or the steady-state level of this protein (Fig. 5).

**Expression of SV-40 TAgWt and K-rasVal12 Does Not Have a Detectable Affect on the Crypt–Villus Patterns of Accumulation of Integrins or Their Ligands**

Depending upon the cell type, interactions with extracellular matrix (ECM) can either promote anchorage-dependent growth or signal cell cycle arrest and terminal differentiation (Lin and Bissell, 1993). Anoikis refers to the propensity of cultured (epithelial or endothelial) cells to undergo apoptosis when dissociated from extracellular matrix (Meredith et al., 1993; Frisch and Francis, 1994; for review see Ruoslahti and Reed, 1994). Integrins have been implicated as mediators of these responses to the ECM (Meredith et al., 1993; Boudreau et al., 1995; Zhang et al., 1995; Pullan et al., 1996; Wary et al., 1996; Frisch et al., 1996). A Ras-linked MAP kinase pathway has been identified recently that functions to reduce the affinity of some integrins for their ligands (Hughes et al., 1997).

It is possible that cycling SV-40 TAgWt-positive villus enterocytes or dysplastic K-rasVal12 × SV-40 TAgWt enterocytes have altered interactions with the ECM and that such alterations curtail their survival; after all, apoptosis is normally limited to the same apical region of the villus where cells detach from the matrix (and from one another) and are exfoliated into the lumen (Hall et al., 1994). Therefore, we used sensitive immunohistochemical detection methods (i.e., tyramide signal amplification; Bobrow et al., 1989; Shindler and Roth, 1996) to analyze the effects of expressing K-rasVal12, SV-40 TAgWt, or both oncoproteins on the crypt–villus distributions of integrins associated with the regulation of epithelial cell growth and survival (αβ1) and with intestinal epithelial cell adhesion to the ECM (αβ4) (Giancotti and Ruoslahti, 1990; Mortarini et al., 1992; Ruoslahti and Reed, 1994; Simon-Assmann et al., 1994; Zhang et al., 1995).

In normal FVB/N mice, α5, α6, β1, and β2 integrin subunits accumulate at the base of all epithelial cells in crypt–villus units (Fig. 6, A, E, H, L, and M). α3, another β1 partner, is limited to the basal surface of villus epithelial cells (data not shown). These patterns are not perturbed in Fabpi–K-rasVal12 or Fabpi–SV-40 TAgWt animals or in their bi-transgenic littermates (n = 2–4 animals/genotype; Fig. 6 A–O).

VL5 (αβ3) and VLA5 (αβ4) function as receptors for fibronectin. In nontransgenic FVB/N mice, fibronectin is distributed in the ECM from the crypt base to the villus tip (Fig. 6 F). Expression of K-rasVal12 or SV-40 TAgWt, alone

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2. Three different preparations of Bax antibodies, two raised against a peptide spanning residues 43–61 of the protein and the other against residues 11–30, yielded the expected patterns of reactivity with jejunal, spleen, and thymic proteins prepared from mice homozygous for wild-type or null alleles of Bax; a 21-kD protein was detected in Bax+/− tissue extracts but was absent from Bax−/− extracts. Affinity-purified polyclonal antibodies against residues 11–30 failed to detect Bax in sections of jejunal crypt–villus units prepared from Bax+/+ or Bax−/− animals. However, both preparations of antibodies to residues 43–61 produced similar patterns of staining in Bax+/− and Bax−/− animals, whether jejunal sections were fixed in Bouin’s solution or formalin and whether or not antigen masking protocols were used. In all cases, staining was blocked by preincubation of the antibody with the Bax peptide before its application to tissue sections. Immunoreactive Bax, or what is presumed to be a protein with a shared epitope, is present at low levels in epithelial cells distributed from the base to the tip of villi. The protein(s) are undetectable in the upper and middle thirds of the crypt. However, long-lived Paneth cells located at the crypt base are intensively stained (data not shown).
Figure 6. Immunohistochemical analysis of the crypt–villus pattern of accumulation of integrin subunits and their ligands in 6-wk-old FVB/N normal, transgenic, and bi-transgenic mice. Frozen sections of jejunum were processed as described in Materials and Methods. The top of each panel is labeled to show the protein that is being examined. The bottom of each panel gives the genotype of the mouse. Antigen–antibody complexes were detected using HRP-conjugated secondary antibodies, biotinyl tyramide amplification, and Cy3-streptavidin. Integrin subunits and integrin ligands appear red-orange. Cells that contain endogenous peroxidase activity were colabeled with FITC–biotinyl tyramide and Cy3-streptavidin and appear yellow-green. nl, normal (nontransgenic) FVB/N mouse; K-ras, littermate containing the Fabp1–K-rasVal12 transgene; TAg, Fabp1–SV-40 TAgWt transgenic animal; K-ras × TAg, Fabp1–K-rasVal12 × Fabp1–SV-40 TAgWt bi-transgenic mouse. (A) The α6 integrin subunit is evident at the basal surfaces of epithelial cells distributed from the base of the crypt (closed arrowhead) through the length of the villus (e.g., closed arrow). (B) Higher power view of a normal villus sectioned at its midpoint, perpendicular to the crypt–villus axis. The arrow points to immunoreactive α6 at the base of enterocytes. (C and D) The crypt–villus distribution of α6 is similar in transgenic mice and does not differ from the distribution observed in their normal littermate shown in A and B. (E–G) The β4 subunit integrin is present throughout crypt–villus units in normal, Fabp1–SV-40 TAgWt transgenic, and bi-transgenic animals. The arrows in G point to a branched (bifurcated) villus. Branching is one manifestation of the dysplasia observed in the small intestine of bi-transgenic mice. (H–K) Distribution of α5 integrin subunit. H and I show jejunal villi that have been sectioned at their midpoint, perpendicular to the crypt–villus axis. Immunoreactive protein is present at the base of villus enterocytes and in cellular components of the lamina propria. J and K show sections cut parallel to the crypt–villus axis. The insets in each of these two panels show cross sections taken at the midpoint of Fabp1–SV-40 TAgWt and bi-transgenic jejunal villi. The integrin subunit is distributed from the base to the villus tip in all animals (J and K plus data not shown). (L–O) The β1 subunit follows the distribution of its α5 partner. L and M show normal jejunal villi sectioned parallel and perpendicular to the crypt–villus axis, respectively. N illustrates results obtained from cross sections of Fabp1–K-rasVal12 villi. O contains a branched villus (arrows) from a bi-transgenic animal. (P and Q) Fibronectin shows a similar distribution along the crypt–villus axis of normal and bi-transgenic mice, as that of its receptor, α5β1 integrin. Bars, 25 μm.
or in combination, has no detectable effect on the accumulation of fibronectin (Fig. 6, P and Q) or on the crypt–villus distribution of two other integrin ligands present in the ECM: type IV collagen and laminin (data not shown).

Interactions between epithelial cells can also affect their susceptibility to anoikis and survival (Frisch and Francis, 1994). E-cadherin is the principal epithelial cadherin in the adult mouse small intestine and is present at the adherens junctions and basolateral surfaces of all villus enterocytes. Forced expression of a dominant negative N-cadherin mutant in jejunal villus enterocytes results in a loss of E-cadherin from the cell surface, a marked reduction in cellular pools of E-cadherin, disruption of cell–cell as well as cell–substratum contacts, and precocious entry into a death program (Hermiston and Gordon, 1995). In light of this previous finding, immunohistochemical surveys were conducted with a well-characterized mouse E-cadherin monoclonal antibody (Hermiston and Gordon, 1995). The results indicated that expression of K-rasVal12 and/or SV-40 TAgWt has no appreciable effects on the intracellular or crypt–villus distributions of E-cadherin (n = 2–5 mice/genotype; data not shown).

Discussion

We have used transgenic mice containing various combinations of K-rasVal12 plus wild-type or mutant SV-40 TAgS to show that K-rasVal12 augments the apoptosis that accompanies a pRB-related reentry of differentiated small intestinal enterocytes into the cell cycle.

The augmented apoptosis occurs through an as yet unknown mechanism. It does not require p53. It is not accompanied by detectable changes in accumulation of regulators of the G1/S transition (pRb, cyclins D1 and E, cdks 2 and 4) or enhanced proliferation. It is not associated with changes in the crypt–villus patterns of expression of several regulators of apoptosis (Bcl-2, Bcl-xL, Bak, Bax, and c-Raf-1). The enhancement is not accompanied by changes in the crypt–villus distribution of mediators of epithelial cell–matrix and cell–cell interactions and survival (e.g., α5β1 integrin and its ligand fibronectin or E-cadherin). In some cases our assays may be too insensitive to rule out involvement. For example, an inability to detect alterations in the crypt–villus pattern of accumulation of five integrin subunits (α5, α6, β1, and β3) or E-cadherin does not definitively prove that attachment of enterocytes to the ECM or to adjacent cells is unperturbed in bi-transgenic animals, or that integrin-activated intracellular signaling events that affect survival (e.g., Frisch et al., 1996; Wary et al., 1996) are fully functional. A recent study using CHO cells indicated that H-ras and its effector kinase, Raf-1, function to negatively regulate the ligand binding affinity of certain integrins (Hughes et al., 1997). Such a finding is consistent with the notion that an anoikis-like mechanism may contribute to the increased apoptosis seen in bi-transgenic mice.

Because K-rasVal12 is the most commonly mutated oncoprotein in colorectal cancer, our in vivo results have implications concerning the multistep journey to intestinal neoplasia. As primary cultures of human and rodent cells, villus enterocytes are not transformed through forced expression of K-rasVal12. The known collaboration that occurs in primary cell cultures between activated forms of Ras and pRB-disabling viral oncoproteins such as E1A, E7, or SV-40 TAgWt also applies to enterocytes. Bi-transgenic K-rasVal12 × SV-40 TAgWt mice exhibit dysplasia in their villus epithelium by the time that crypt–villus morphogenesis is completed between P21–28. This dysplasia is manifested in part by branched villi. Branching involves the epithelium, its underlying ECM, and the lamina propria (e.g., Fig. 6, G, O, and Q), providing an interesting model system for examining how the small intestinal epithelium is able to define overall villus structure.

Despite the early onset of this dysplasia, bi-transgenic animals do not develop small intestinal neoplasms between 2 and 18 mo of age (n = 40 animals). The lack of progression may be due to the increased apoptosis produced by K-rasVal12 in this “initiated” cell population with a proliferative abnormality. Another contributing factor may be the short residence time of migrating enterocytes on the villus; without functional anchorage, they are not able to undergo clonal expansion. As noted in the introduction, K-rasVal12 is the most commonly mutated oncoprotein in human colorectal tumors (Fearon and Vogelstein, 1990). Activating K-ras mutations are not acquired at the beginning but rather at intermediate points in the multistep journey to intestinal neoplasia (Bos et al., 1987; Forrester et al., 1987; Fearon and Vogelstein, 1990). Our results are consistent with this observation. K-rasVal12 alone has no detectable effect on noncycling enterocytes. When expressed in “initiated” enterocytes with a proliferative abnormality, K-rasVal12 can facilitate progression to a dysplastic phenotype. However, by diminishing cell survival on the villus, the oncoprotein may also impede further progression.

Intestinal neoplasia is thought to be initiated in the crypt, perhaps at the level of the functionally anchored stem cell or one of its immediate daughters (Moser et al., 1992). Although the effect of K-rasVal12 on initiated crypt epithelial cells is unknown, our studies suggest that on the villus, K-rasVal12 may serve an editing function for the host, helping to remove these cells. Additional mutations may be needed to suppress the proapoptotic response to K-rasVal12. Such mutations may affect genes encoding one or more determinants of cell survival. Since downregulation of apoptosis can contribute to the late stages of tumor progression, at least in model systems (Naik et al., 1996), it is tempting to speculate about the significance of the increased expression of antiapoptotic mediators, such as Bcl-xL or Bel-2, that has been noted during progression of human colorectal tumors (Hague et al., 1994; Bedi et al., 1995; Bronner et al., 1995; Krajewska et al., 1996). Crossing bi-transgenic K-rasVal12 × SV-40 TAgWt mice to mice that overexpress these or other mediators of cell survival in their gut epithelium provides one way of identifying molecules that may overcome the K-rasVal12–augmented apoptosis and thus promote development of gut neoplasia.

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