Srcasm Inhibits Fyn-Induced Cutaneous Carcinogenesis with Modulation of Notch1 and p53

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Abstract
Src family tyrosine kinases (SFK) regulate cell proliferation, and increased SFK activity is common in human carcinomas, including cutaneous squamous cell carcinomas (SCC) and its precursors. The elevated SFK activity in cutaneous SCCs was modeled using K14-Fyn Y528F transgenic mice, which spontaneously form punctate keratotic lesions, scaly plaques, and large tumors resembling actinic keratoses, SCC in situ, and SCCs, respectively. Lesional tissue showed increased levels of activated SFKs, PDK1, STAT3, and ERK1/2, whereas Notch1/ NICD protein and transcript levels were decreased. p53 levels also were decreased in SCC in situ and SCCs. Increasing Srcasm levels using a K14-Fyn Y528F/K14-Srcasm double transgenic model markedly inhibited cutaneous neoplasia. In contrast, increased expression of a nonphosphorylatable Srcasm mutant maintained the neoplastic phenotype. Increasing Srcasm levels decreased levels of Fyn, activated SFKs, ERK1/2, PDK1, and phospho-STAT3, and increased Notch1/ NICD and p53 levels. Analysis of human specimens revealed that levels of Fyn and activated SFKs were elevated in SCCs compared with adjacent nonlesional epidermis. In addition, Notch1 and Srcasm protein and transcript levels were decreased in human SCCs compared with nonlesional epidermis. Therefore, the SCCs produced by the Fyn Y528F mice resemble their human counterparts at the molecular level. K14-Fyn Y528F mice represent a robust model of cutaneous carcinogenesis that manifests precancerous lesions and SCCs resembling their human counterparts at the molecular level.

Introduction
Cutaneous squamous cell carcinoma (SCC) is the second most common form of cancer with >250,000 cases annually in the United States leading to ~2,500 deaths3,4 (1). Many cutaneous SCCs arise from a precursor lesion termed an actinic keratosis, and ~60% of all individuals ages >40 years will develop an actinic keratosis requiring treatment during their lifetime, making actinic keratosis the most common precancerous lesion in the United States (2). Such data demonstrate that actinic keratoses and cutaneous SCCs are a significant health problem and show a need for in vivo model resembling this human disease.

Src family tyrosine kinases (SFK) are known oncogenes and promote neoplasia in many tissues (3, 4). The majority of human carcinomas, including colonic, breast, pancreatic, and cutaneous squamous cells, show elevated SFK activity compared with corresponding nonneoplastic epithelium (5–9). However, the cellular mechanisms promoting increased SFK activity in human carcinomas remain unclear. Mutation of a COOH-terminal regulatory tyrosine can lead to increased SFK activity in tumors (3). However, such activating mutations in SFKs are rare in human carcinomas (10–13). These observations raise the hypothesis that additional cellular mechanisms could account for the elevated SFK activity in carcinomas, such as impaired negative regulation of SFKs.

Keratinocytes express three SFKs: Src, Fyn, and Yes (14). Fyn is a dually acylated kinase containing covalently linked myristoyl and palmitoyl moieties on its NH2-terminal SH4 domain (15). Palmitoylation targets SFKs to lipid rafts (16, 17), and Fyn localizes to lipid rafts and caveolae, subcellular domains associated with epidermal growth factor receptor signaling and caveolae/raft-dependent endocytosis (18–20). These characteristics suggest that Fyn may play a significant role in regulating keratinocyte proliferation.

Srcasm (Src activating and signaling molecule) is a SFK substrate that, when phosphorylated, engages SFKs and downregulates them through a lysosomal-dependent mechanism (18, 21, 22). Srcasm localizes to the multivesicular body that is important for targeting endosomal proteins for lysosomal degradation (23, 24). Srcasm also contains two conserved Tsg101 tetrapeptide binding motifs (25); Tsg101 is a component of the ESCRT-1 complex and a regulator of the multivesicular body (23, 26). Srcasm lies at a signaling nexus between SFKs, Tsg101, and lysosomal protein degradation.

Human cutaneous SCC in situ (SCIS) and SCCs show decreased Srcasm levels compared with unremarkable epidermis (18). SFK activity is elevated in actinic keratoses, SCIS, and SCC in human biopsies compared with adjacent nonlesional epidermis (6). Together, these data suggest an inverse relationship between SFK activity and Srcasm levels in human skin neoplasia.

Therefore, we hypothesized that increasing Fyn activity in the epidermal keratinocytes would promote cutaneous neoplasia and increasing Srcasm levels would inhibit Fyn-induced neoplasia.

Note: Supplementary data for this article are available at Cancer Research Online (http://cancerres.aacrjournals.org/).

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These hypotheses were tested using transgenic mice expressing Fyn Y528F, Srcasm, and SrcasmDN, a nonphosphorylatable Srcasm mutant. Increased Fyn Y528F expression induced the spontaneous formation of precancerous lesions and SCCs within 5 weeks. Increased expression of Srcasm, but not SrcasmDN, inhibited the formation of precancerous and cancerous lesions. Elevated Fyn levels promoted ERK1/2, PDK1, and STAT3 activation and downregulated Notch1 and p53. Increased Srcasm lowered Fyn levels in vivo and normalized the activity and levels of these molecules.

Together, these data identify a mechanism of carcinogenesis in which the level of Fyn activity is inversely related to Srcasm levels; the data also show an important regulatory relationship between Fyn and Srcasm that involves Notch1 and p53. The relationship between SFKs, Srcasm, Notch1, and p53 may play an important role in regulating carcinogenesis.

Materials and Methods

Generation and characterization of transgenic mice. Murine HA-tagged Srcasm and HA-tagged SrcasmDN transgenic mice were generated previously (21). FynY528F cDNA was cloned into a vector driven by the human K14 promoter (21). Transgene cassettes were excised and purified via Tris-acetate-EDTA-agarose electrophoresis. C57BL/6 × CBA-fertilized oocytes were microinjected with the transgene cassettes using standard protocols at the University of Pennsylvania Transgenic Core Facility in accordance with Institutional Animal Care and Use Committee proposal 801519. Two independent K14-Fyn Y528F lines with hyperkeratotic plaques and tumor phenotype were derived. Founders were crossed with C57BL/6 mice to generate the C57BL/6 lines. The C57BL/6 lines were crossed with FVB/N and the F1 hybrids were characterized in this study. Up to three additional backcrosses of the Fyn Y528F transgene onto the FVB/N background yielded an identical phenotype. A two-sided Fisher’s exact test was used to determine the statistical significance of differences in phenotype incidence.

Antibodies. Activated Src family kinase, phospho-ERK1/2, Fyn, keratin 6, phospho-STAT3, phospho-PDK1, β-actin, and α-Srcasm antibodies were used as described previously (25). For Western blotting, α-p21 mouse monoclonal antibody (F-5; 1:500; Santa Cruz Biotechnology), α-MDM2 mouse monoclonal antibody (2A10; 2 μg/mL; Calbiochem), α-p53 mouse monoclonal antibody (1C12; 1:1,000; Cell Signaling), and α-Notch1/NICD (C4H11; 1:1,000; Cell Signaling).

Histologic and immunohistochemical analysis. Skin samples were fixed in 10% neutral-buffered formalin and subjected to standard processing and staining with H&E. Tissue sections were subjected to immunohistochemistry as reported (6). Photomicrographs were obtained using a Leica DC300 digital camera coupled to a Zeiss Axioshot microscope; photos were obtained under identical conditions at the indicated magnifications as high-resolution JPEG files.

Immunoblotting. Cell and tissue lysates were prepared and analyzed as described (22).

Quantitative reverse transcription-PCR. Total RNA was isolated from epidermis and tumor samples using an OMNI homogenizer and the NEasy Fibrous Tissue kit (Qagen) according to the manufacturer’s instructions. RNA quantity was assessed using a NanoDrop spectrophotometer (ThermoScientific). Total RNA was reverse-transcribed with random hexamers using High-Capacity RNA-to-cDNA kit (Applied Biosystems) following the manufacturer’s protocols. Equivalent amounts cDNA were subjected to quantitative PCR using the Power SYBR Green PCR master mix (Applied Biosystems) on an ABI 7000 instrument. Samples were run in triplicate on 96-well reaction plates with appropriate species-specific primer pairs. Primers sets for each target gene were designed and purchased from Integrated DNA Technology. The comparative Ct method was used to determine the level of the target gene mRNA in tissue samples (Applied Biosystems User Bulletin #2, October 2001). Human samples were standardized to β-actin and murine samples were standardized to glyceraldehyde-3-phosphate dehydrogenase. Statistical analysis was done using the independent groups t test for means.

Results

C57BL/6 K14-Fyn Y528F transgenic mice exhibit hyperkeratotic plaques and spontaneous tumor formation. The effect of the K14-Fyn Y528F transgene was evaluated in the C57BL/6 genetic background, a tumor-resistant genetic background (27). Increased epidermal expression of Fyn Y528F induces the development of multiple hyperkeratotic plaques in the epidermis, whereas control mice did not exhibit this phenotype (P < 1.2 × 10−19; Fig. 1A; Supplementary Table S1). The hyperkeratotic plaques are detectable at ages 3 to 4 days and persist until ages 3 to 4 weeks. Approximately 21% of the K14-Fyn Y528F C57BL/6 mice that exhibited hyperkeratotic plaques developed spontaneous keratotic tumors resembling cutaneous SCCs between ages 3 and 7 months, whereas littermate controls mice do not (P = 0.001; Fig. 1B). Spontaneous SCCs arising in the C57BL/6 background is unusual and suggests that Fyn Y528F functions as an oncogene in epidermal keratinocytes (27).

To test the effects of increasing Srcasm levels on the K14-Fyn Y528F phenotype, K14-Fyn Y528F/K14-Srcasm double transgenic mice were generated; these double transgenic mice exhibited a significantly lower incidence of hyperkeratotic plaques compared with the parental line (P = 0.003; Supplementary Table S1). In addition, no keratotic tumors formed in the K14-Fyn Y528F/K14-Srcasm double transgenics; these data show that the Srcasm transgene inhibits tumor formation in C57BL/6 K14-Fyn Y528F mice (P = 0.04).

F1 hybrids between C57BL/6 K14-Fyn Y528F and FVB/N develop multiple spontaneous SCCs and precancerous lesions. To characterize the effect of the Fyn Y528F transgene in a standard tumor-permissive genetic background, the C57BL/6 × K14-Fyn Y528F line was crossed with FVB/N to generate F1 progeny. These F1 hybrid progeny showed hyperkeratotic plaques within 4 days postnatal. The C57BL/6 × FVB/N F1 K14-Fyn Y528F hybrids developed tumors sooner than the C57BL/6 parental line, typically manifesting multiple tumors between 5 and 8 weeks. The formation of hyperkeratotic plaques in the K14-Fyn Y528F F1 hybrids was statistically significant compared with littermate controls (P < 4.5 × 10−96; Supplementary Table S2). Spontaneous SCC formation was observed in 36% of K14-Fyn Y528F mice with none arising in littermate controls (Fig. 1C; P = 4.6 × 10−23). K14-Fyn Y528F transgenic mice derived from backcrossing the F1 hybrids one (n = 20) or two (n = 10) additional times onto the FVB/N background yielded a phenotype indistinguishable from the F1 hybrid.

The incidence of hyperkeratotic plaque formation in K14-Fyn Y528F/K14-Srcasm double transgenic mice (41%) was decreased compared with the parental line (81%; P < 1.4 × 10−4; Supplementary Table S2). Tumor formation was almost completely inhibited in the K14-Fyn Y528F/K14-Srcasm double transgenic mice (P < 1.5 × 10−3), suggesting that Srcasm can function as an anti-oncogene. The incidence of tumor formation was similar between K14-Fyn Y528F mice and K14-Fyn Y528F/K14-SrcasmDN double transgenic mice that
express a nonphosphorylatable form of Srcasm (\(P = 1\)). Therefore, the Srcasm\(\text{DN}\) transgene does not inhibit SCC formation and does not function as an anti-oncogene. The incidence of hyperkeratotic plaque and SCC formation was lower in the K14-Fyn Y528F/K14-Srcasm\(\text{DN}\) line compared with the K14-Fyn Y528F/K14-Srcasm\(\text{DN}\) lines with \(P < 5.3 \times 10^{-3}\) and 0.01, respectively.

In addition to keratotic tumors, the C57BL/6 K14-Fyn Y528F FVB/N F1 hybrids develop punctate (1-3 mm) hyperkeratotic lesions at 4 to 5 weeks that clinically resemble human precancerous lesions termed actinic keratoses (Fig. 1D). Histologic analysis shows a precancerous lesion exhibiting hyperplasia, parakeratosis (retained nuclei in the stratum corneum), and keratinocyte atypia (Fig. 1D).

The K14-Fyn Y528F FVB/N mice with tumors were followed for at least 6 months and did not show evidence of metastasis, which mirrors the low metastatic potential seen with human cutaneous SCCs.

**Histologic analysis of the hyperkeratotic plaques and tumors shows SCIS and SCC.** Histologic analysis of the hyperkeratotic plaques that develop during the first week postnatal in the K14-Fyn Y528F C57BL/6 and C57BL/6-FVB/N F1 mice shows a markedly thickened epidermis with increased mitotic activity, cytologic atypia, and architectural disorganization compared with adjacent nonlesional epidermis (Fig. 2A). These histologic features are consistent with SCIS and resemble human SCIS (Fig. 2A).

The C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F mice exhibited small hyperkeratotic lesions ranging in size from 1 to 3 mm (Fig. 2B); these lesions show focal epidermal hyperplasia, keratinocyte dysplasia, and hyperkeratosis, resembling a human actinic keratosis (Fig. 2B).

The spontaneous skin tumors that developed in both genetic backgrounds carrying the K14-Fyn Y528F transgene manifested enlarged keratinocytes with nuclear atypia, mitotic activity, dermal invasion, and focal keratinization (Fig. 2C). The cytologic and histologic features exhibited by these murine lesions are consistent with well-differentiated SCCs and closely resemble corresponding human lesions (28, 29). These data show that the Fyn Y528F transgene induces cutaneous lesions that are histologically similar to human actinic keratoses, SCIS, and SCCs.

**Immunohistochemical analysis of precancerous lesions and cutaneous tumors in C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F mice.** Immunohistochemical analysis of small precancerous lesions from a 5-week-old C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F mouse showed increased levels of activated SFKs, keratin 6, phospho-STAT3, phospho-ERK1/2, and phospho-PDK1 (Fig. 3A). Analysis of cutaneous SCCs from 5-week-old C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F mice showed a pattern similar to that seen in the precancerous lesions with markedly increased levels of Fyn, activated SFKs, and keratin 6 compared with nonlesional epidermis (Fig. 3B). Increased staining for phospho-STAT3,
phospho-PDK1, and phospho-p44/42 also was seen in the SCCs. Together, the data suggest that neoplastic lesions secondary to elevated Fyn activity also exhibit increased activation of STAT3, PDK1, and ERK1/2. Similar staining was seen in SCIS lesions (data not shown) and SCIS from K14-Fyn Y528F C57BL/6 mice (Supplementary Fig. S1).

**Increased Fyn activity decreases Notch1/NICD transcript levels.** The formation of precancerous lesions and SCCs in C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F mouse phenocopies SM22α-DNMAML mice, which have decreased epidermal Notch1 signaling (30). Therefore, a link between increased Fyn activity and decreased Notch1 signaling was assessed. Quantitative reverse transcription-PCR (qRT-PCR) for Notch1 transcript was done on total RNA from unremarkable skin, SCIS lesions, and SCCs from K14-Fyn Y528F mice and littermate controls. Skin from FVB mice showed relatively high levels of Notch1 transcript (Fig. 4A). K14-Fyn Y528F SCIS lesions showed ∼2% of the Notch1 transcript levels seen in control mice (P < 0.01). SCCs from K14-Fyn Y528F and double transgenic expressing SrcasmDN exhibited only 1% of the control Notch1 transcript levels (P < 0.01). These data show that the cutaneous lesions associated with increased Fyn activity contain lower levels of Notch1 transcript.

**Increased Fyn activity decreases Srcasm transcript levels.** Srcasm is a negative regulator of activated SFKs, including Fyn; therefore, the levels of Srcasm transcript were assessed in control skin, SCIS-like lesions, and SCCs (21). Control skin contained relatively high levels of Srcasm transcript, whereas SCIS lesions contained only 13% (P = 0.041) of control levels (Fig. 4B). SCCs from K14-Fyn Y528F mice contained only 1% of control levels of Srcasm transcript (P = 0.029). As expected, SCCs from the K14-Fyn Y528F/SrcasmDN mice contained ∼20 times the control level of Srcasm transcript because of the SrcasmDN transgene (data not shown).

**Increased Fyn activity decreases Notch1/NICD levels and promotes STAT3 phosphorylation.** To determine if the Fyn-induced decrease in Notch1 transcript levels correlated with a decrease in Notch1/NICD protein levels, Western blot analysis was done on lysates of control skin, SCIS, and SCCs. Increased Fyn levels and activated SFK levels in SCIS and SCCs correlated with a dramatic drop in Notch1/NICD protein levels (Fig. 4C). K14-Fyn Y528F/Srcasm double transgenic mice showing a weak phenotype exhibited increased Srcasm levels associated with partially normalized Fyn and Notch1/NICD levels (Fig. 4C). K14-Fyn Y528F/Srcasm double transgenic mice exhibiting a corrected phenotype showed increased Srcasm levels associated with normalized levels of Fyn, activated SFKs, and Notch1/NICD.

In oral SCCs, the level of phospho-STAT3 is tightly correlated with the levels of SFK activity (31). Likewise, in the K14-Fyn Y528F skin lesions, the level of phospho-STAT3 is proportional to the levels of Fyn and activated SFKs (Fig. 4C). Together, these data show that increased levels of activated Fyn enhance phospho-STAT3 levels and downregulate Notch1 transcript, Notch1 protein, and Srcasm transcript levels. In K14-Fyn Y528F/K14-Srcasm mice, increasing Srcasm levels decreases Fyn and phospho-STAT3 levels while restoring Notch1/NICD levels. Together, Fyn and Srcasm can alter Notch1 levels while promoting or inhibiting neoplasia, respectively.

**Increased SrcasmDN expression does not downregulate levels of Fyn Y528F in transgenic skin.** SrcasmDN has its known Fyn phosphorylation sites mutated to phenylalanines, and this mutant cannot be phosphorylated by Fyn nor downregulate native Fyn.

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**Figure 2.** K14-Fyn Y528F lesions histologically mimic human lesions. A, human SCIS shows full-thickness keratinocyte atypia, epidermal hyperplasia, and hyperkeratosis without evidence of dermal invasion. Similar findings are seen in hyperkeratotic plaques taken from the back skin of K14-Fyn Y528F 1-week-old mice of the indicated genetic backgrounds. Bar, 200 μm; magnification, ×125. B, C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F mice show precancerous lesions that resemble human actinic keratoses. AK, actinic keratosis; Pre, precancerous lesion from the ear of a 5-week-old mouse. Bar, 200 μm; magnification, ×125. C, biopsies of the spontaneous cutaneous tumors from a 5-week-old FVB/N F1 hybrid and an 8-week-old C57BL/6 mouse show SCCs resembling human cutaneous SCC. Bar, 800 μm; magnification, ×31.25 (low). Bar, 100 μm; magnification, ×250 (high). Representative of three independent mice.
or activated Fyn in vitro (21). To better understand why SrcasmDN does not inhibit SCC formation, protein lysates from littermate control skin, K14-Fyn Y528F SCCs, and K14-Fyn Y528F/SrcasmDN SCCs were subjected to Western blotting. Western blot analysis shows that the SrcasmDN molecule does not downregulate Fyn Y528F like native Srcasm (Fig. 4D); therefore, levels of activated SFKs remain elevated and Notch1/NICD levels remain low. These data confirm that SrcasmDN cannot downregulate Fyn Y528F in vivo.

SFK activity is increased and Notch1 and Srcasm levels are decreased in human SCC. The relationship between Fyn, SFK activity, Srcasm, and Notch1 has not been evaluated in genetically matched nonlesional human skin and SCCs (6, 18, 30, 32). To address this question, protein lysates of human SCCs and adjacent nontumorigenic epidermis from two independent patients were subjected to Western blot analysis for activated SFKs, Fyn, NICD, and Srcasm (Fig. 5A). Human SCCs show increased levels of activated SFKs and Fyn compared with unremarkable skin. Human SCCs also show decreased levels of NICD and Srcasm compared with unremarkable skin. These data show that the increased Fyn and SFK activity in human SCCs are associated with decreased NICD and Srcasm levels.

To determine if the decreased NICD and Srcasm levels were associated with lower mRNA levels, qRT-PCR was done on mRNA isolated from three randomly selected human SCCs and three independent unremarkable skin samples. Both NICD and Srcasm transcript levels were decreased in the SCCs compared with unremarkable skin (Fig. 5B). These data suggest that human cutaneous SCCs manifest decreased Notch1 and Srcasm transcript and protein levels. Together, these data show that human SCCs and the K14-Fyn Y528F SCCs exhibit similarities regarding the levels of Fyn, SFK activity, Srcasm, and Notch1.

**Fyn and Srcasm modulate p53 levels.** p53 is an important regulator of cutaneous carcinogenesis in humans, and impaired p53 function is associated with the formation of actinic keratoses and SCCs (32, 33). Therefore, we examined the levels of p53 protein and mRNA in skin samples from our transgenic lines. In C57BL/6-FVB/N F1 hybrid K14-Fyn Y528F SCCs, increased Fyn levels were associated with decreased p53 levels compared with age-matched nontransgenic controls (Fig. 6A). Western blot analysis shows that increasing Srcasm levels elevates p53 levels in K14-Fyn Y528F/K14-Srcasm mice compared with nontransgenic controls and K14-Fyn Y528F mice (Fig. 6I). In contrast, increased expression of SrcasmDN lowers p53 levels in SCIS and SCCs in K14-Fyn Y528F/SrcasmDN double transgenic mice (Fig. 6A and B). In SCCs, as in SCIS lesions, increased Fyn expression was associated with lower p53 levels compared with controls (Fig. 6B).

In all samples, p21 levels closely paralleled p53 levels, implying that p53 is transcriptionally active (Fig. 6A and B; ref. 34). MDM2 levels remained constant in all tissue samples, except in SCCs from K14-Fyn Y528F mice; therefore, fluctuations in MDM2 levels do not correlate with the significant changes in p53 levels (Fig. 6A and B).

qRT-PCR analysis of RNA from these samples for p53 transcript shows that p53 transcript levels were lower in mice harboring a Fyn Y528F transgene (Fig. 6C). However, the fluctuations in p53 protein levels among the various Fyn transgenic lines did not correlate with changes in p53 transcript levels (Fig. 6C). The p53 transcript level is significantly lower in the Fyn transgenic lines together compared with controls (P < 0.02). In the Y528F transgenic lines, increasing Srcasm levels lower Fyn levels and elevate p53, whereas increasing levels of SrcasmDN decrease p53 levels compared with controls. These in vivo data suggest that Fyn downregulates p53 transcript levels, whereas Srcasm can modulate p53 protein levels independent of transcript levels. Therefore, changes in Fyn and Srcasm levels could alter cellular p53 levels and influence keratinocyte susceptibility to genotoxic stress.
Discussion

K14-Fyn Y528F transgenic mice represent a robust model of cutaneous carcinogenesis that spontaneously form neoplastic lesions resembling those seen in human cutaneous neoplasia. Within the first week postnatal, K14-Fyn Y528F transgenic mice exhibit hyperkeratotic plaques resembling human SCIS at the histologic and molecular levels. These SCIS lesions show increased activity of SFKs, the Ras/mitogen-activated protein kinase pathway, and the phosphoinositide 3-kinase/PDK1/Akt kinase pathway, similar to human lesions and other murine models (6, 35–38).

At 4 to 5 weeks, the K14-Fyn Y528F FVB/N F1 hybrid transgenic mice spontaneously form punctate hyperkeratotic lesions resembling human actinic keratoses; both human actinic keratoses and these murine precancerous lesions are associated with increased levels of activated SFKs (6). Spontaneous generation of precancerous lesions within a short time makes the K14-Fyn Y528F model well suited for screening topical agents that may be efficacious in treating actinic keratoses.

At 5 weeks, K14-Fyn Y528F mice spontaneously develop cutaneous SCCs that have not metastasized within a 6-month follow-up period, which is consistent with the low metastatic incidence of human cutaneous SCCs, ~4% in tumors 2 to 6 mm thick (39).

Transgenic mice expressing Src or activated mutant Src in the epidermis have been reported to develop SCCs spontaneously after 3 months, have an increased risk for developing papillomas secondary to two-stage chemical carcinogenesis, or develop SCCs in
25% of mice on the edges of healing wounds (40–42). However, these Src transgenic models take longer to manifest tumors and actinic keratosis–like lesions have not been described. The K14-Fyn Y528F lines develop actinic keratosis–like lesions and spontaneous SCCs in a 4- to 5-week time frame showing that Fyn is a potent oncogene in keratinocytes.

The K14-Fyn Y528F transgenic lines phenocopy the SM22α-DNMAML mouse, although the SM22α-DNMAML line typically takes 6 months to manifest lesions (30). Western blot and qRT-PCR analysis of skin, SCIS, and SCCs from the various K14-Fyn Y528F lines showed that increased levels of Fyn and activated SFKs are associated with decreased levels of Notch1 mRNA, Notch1 protein, and NICD. These data link increased SFK activity in keratinocytes with Notch1 downregulation in vivo. This observation correlates with prior observations showing that SFKs activate MEK1/ERK1 downstream of epidermal growth factor receptor and that increased epidermal growth factor receptor signaling downregulates Notch1 transcription through a MEK1/ERK1-dependent pathway (18, 32). Notch1 downregulation by Fyn may be important for promoting neoplasia; it will be interesting to determine if increased NICD expression can inhibit Fyn-dependent SCC induction.

Figure 5. Elevated SFK levels correlate with decreased Notch1 and Srcasm levels in human SCC. A, levels of activated SFKs are inversely related to NICD and Srcasm levels. Western blot analysis of lysates from paired nonlesional human skin and adjacent SCC using the indicated antibodies. Corresponding samples from two patients. B, Notch1 and Srcasm transcripts are downregulated in human SCC. qRT-PCR was done on mRNA derived from independent unremarkable human skin samples and SCCs to determine Notch1 and Srcasm transcript levels. Normalized mean and SD are shown. Three independent unremarkable skin and SCC samples were analyzed. *, P = 0.025; **, P = 0.064.

Figure 6. Fyn and Srcasm modulate p53 levels. A, analysis of SCIS lesions and age-matched controls. Protein lysates from SCIS lesions of C57BL/6-FVB/N F1 hybrid transgenic lines or controls were subjected to Western blot analysis for p53, p21, MDM2, and β-actin. Representative of two independent sets of mice. B, analysis of SCC lesions and age-matched controls. Protein lysates from SCCs of the indicated lines or controls were subjected to Western blot analysis as in A. Representative of two independent sets of mice. C, qRT-PCR analysis for p53 transcript. mRNA from the indicated lesions and K14-Fyn Y528F FVB/N F1 hybrid transgenic lines or controls were subjected to qRT-PCR for p53. Data derived from two independent sets of samples. D, Fyn/Srcasm signaling nexus. Arrows, positive regulatory relationships; T bars, negative regulatory relationships. Fyn increases levels of PDK1, ERK1/2, and STAT3 activation. Fyn lowers Notch1 transcript and protein levels perhaps through p53-dependent and p53-independent mechanisms. Fyn lowers Srcasm and p53 transcript levels. Srcasm lowers Fyn protein levels.
Supraphysiologic SFK levels and subphysiologic Srcasm levels are common in human actinic keratoses, SCIS, and cutaneous SCCs (6). These findings suggest that the inverse relationship between SFK activity and Srcasm levels may represent a common mechanism of carcinogenesis. Supporting this hypothesis, Srcasm levels are decreased in human basal cell carcinomas and esophageal SCC tumor cell lines compared with nonneoplastic keratinocyte lines (data not shown; ref. 43).

The data presented show that epidermal neoplasia develops when keratinocytic SFK activity exceeds the negative regulatory capacity of endogenous Srcasm (Fig. 6D). However, the pro- oncogenic effect of Fyn can be inhibited by increasing the Srcasm level such that its SFK-downregulatory capacity restores Fyn levels to near physiologic. The data presented suggest that one mechanism of SFK-dependent neoplasia would be to disrupt the balance between SFK activity and Srcasm-dependent SFK downregulation so that it elevates SFK activity. Such an event could result by decreasing intracellular Srcasm levels. Further characterization of the mechanisms that regulate Srcasm levels may provide insights into SFK-dependent carcinogenesis.

The ability of Srcasm to inhibit Fyn-induced carcinogenesis suggests that Srcasm can function as an anti-oncogene. As such, Srcasm appears to represent a novel class of anti-oncogene that targets activated SFKs for degradation in a lysosomal-dependent manner (21). Such a pathway may be important for limiting SFK signaling and maintaining cell homeostasis. Manipulation of Srcasm levels may provide a tool for modulating the level of SFK signaling and a means of promoting or inhibiting epithelial cell growth.

Disclosures of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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