Possible Role of Semaphorin 3F, a Candidate Tumor Suppressor Gene at 3p21.3, in p53-Regulated Tumor Angiogenesis Suppression

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Abstract

Although the regulation of tumor angiogenesis is believed to be one of the core functions of p53, the mechanism still remains to be elucidated. Here, we report that semaphorin 3F (SEMA3F), an axon guidance molecule, is involved in p53-regulated antiangiogenesis. The expression level of SEMA3F mRNA was increased by both exogenous and endogenous p53. Chromatin immunoprecipitation assay indicated that a potent p53-binding sequence in intron 1 of SEMA3F interacts with p53 and that it has a p53-responsive transcriptional activity. Overexpression of SEMA3F inhibited in vitro cell growth of the lung cancer cell line H1299. In nude mice assay, the size of the H1299 tumors expressing SEMA3F was much smaller, and they showed lesser number of blood vessels as compared with the control tumors. Moreover, tumors derived from the p53-knockdown colorectal cancer cell line LS174T displayed a remarkable enhancement of tumor vessel formation as compared with control tumors containing normal levels of p53. The expression levels of SEMA3F and neuropilin-2 (NRP2), the functional receptor for SEMA3F, in p53-knockdown LS174T tumors were lower than those in the control tumors. Adenovirus-mediated SEMA3F gene transfer induced the remarkable in vitro growth suppression of the stable transfectant of H1299 cells, which express high levels of NRP2. These results suggest that p53 negatively regulates tumor vessel formation and cell growth via the SEMA3F-NRP2 pathway.

Introduction

The tumor suppressor p53 prevents malignant transformation when the cells suffer cellular stresses including severe DNA damage (1). Hence, p53 is known as a “guardian of human genome” (2). Indeed, p53 is mutated in >50% of all human cancers, emphasizing its essential role in tumorigenesis in any of the human cancers. Because of its importance, numerous studies on p53 have been conducted thus far, and several findings have been reported. These efforts have provided concrete evidence for the following functions of p53. The p53 gene encodes a transcription factor that binds to a specific sequence of its downstream target gene (3). Therefore, p53 exerts its functions via transcriptional activation of various target genes (3, 4). Although a considerable number of target genes have been reported, their functions display great diversity, and four major ones, including genes functioning in cell cycle arrest, apoptosis, DNA repair, and antiangiogenesis, are considered to be involved in the core mechanism of p53-regulated tumor suppression (1, 3, 4).

Because angiogenesis plays a critical role in tumor formation and progression, blocking tumor angiogenesis is one of vital antitumor functions. An inhibitor of angiogenesis was previously detected in the condition medium derived from glioblastoma cells, wherein exogenous p53 was overexpressed (5). This implies that p53 may activate the transcription of an antiangiogenic factor gene. Consistent with this assumption, thrombospondin-1 (TSP-1), which is a pivotal factor for angiogenesis, was identified as the transcriptional target gene of p53 by using cultured fibroblast cells from Li-Fraumeni patients (6). This fact confirmed that p53 might play an important role in the control of angiogenesis during tumor development. In addition, a study reported that brain-specific angiogenesis inhibitor 1 (BAI1) is the second example of p53-regulated antiangiogenic factor (7). Although these two molecules are really interesting and promising candidates, the precise mechanism of p53-regulated antiangiogenesis still remains to be elucidated.

Semaphorins are a highly conserved family of molecules that contribute to axon guidance during neural development and differentiation by repulsing axons and inhibiting growth cone extension (8–11). Based on their structure, they are classified into seven subclasses, including transmembrane proteins (classes 1, 4, 5, and 6), secreted proteins (classes 2 and 3), and proteins associated with the cell surface through glycosylphosphatidylinositol linkages (class 7; ref. 12). Semaphorin 3F (SEMA3F) belongs to the class 3 secreted type of semaphorin protein (12). During the developmental process, SEMA3F plays a critical role in axon guidance in both the peripheral nervous system and the central nervous system by interacting with its receptor, neuropilin-2 (NRP2; ref. 13). The SEMA3F-NRP2 signaling pathway guides axonal extension by means of a chemotactic repulsing effect on the axons (14–16).

Although semaphorins play a critical role as axon guidance molecules in the developing nervous system, they are expressed in a variety of adult and embryonic tissues, suggesting a broader spectrum of the functions for semaphorins. In fact, SEMA3F was initially identified as a candidate tumor suppressor gene at chromosome 3p21.3; loss of heterozygosity from 3p21.3 is frequently observed in many cases of human cancers (17–19). In particular, the region is most frequently deleted and at least one allele is lost in nearly 100% of the cases of small-cell lung cancer (20, 21). Although extensive analysis was conducted, no point mutations in SEMA3F were detected in human cancers (17–19).
Instead, the expression of \textit{SEMA3F} was frequently down-regulated epigenetically in a number of cancers, implying that alterations of \textit{SEMA3F} may be involved in tumorigenesis (22). Moreover, overexpression of \textit{SEMA3F} in cancer cells induced the growth suppression of some cancer cells (23). Despite these observations, the role of \textit{SEMA3F} in tumorigenesis remains controversial.

Here, we report yet another finding that implies that \textit{SEMA3F} is a bona fide tumor suppressor gene. In the present study, we have found that \textit{SEMA3F} is a direct target gene for tumor suppressor p53 and that \textit{SEMA3F} is likely to mediate p53-regulated antiangiogenesis during tumor development. Additionally, it is likely that NRP2 expression is also controlled by p53. Our findings suggest a novel p53regulated mechanism of antiangiogenesis and cell growth suppression via the \textit{SEMA3F-NRP2} pathway.

Materials and Methods

Cell culture. Human cancer cell lines HepG2 (hepatoblastoma), H1299 (lung cancer), T98G (Glioblastoma), LS174T (colon cancer), and MCF7 (breast cancer) were purchased from the American Type Culture Collection (Manassas, VA). Colon cancer cell lines HCT116 p53(+/+) and HCT116 p53(−/−) were a gift from Dr. Bert Vogelstein (Johns Hopkins University, Baltimore, MD). All cells were cultured under the conditions recommended by their respective depositors.

RNA interference. By using two control cell lines (HepG2 and LS174T) containing wild-type p53 (wt-p53), we established p53-knockdown cell lines (HepG2-p53-siRNA and LS174T-p53-siRNA) and control cell lines (HepG2-control and LS174T-control) as previously described (24). In brief, these cells were infected with SI-MSCV-puro-H1R-p53si retrovirus for downregulation of p53 expression and with SI-MSCV-puro-H1R retrovirus for negative control. Then, the infected cells were selected with 1 \mu M puromycin for 2 weeks and the single clones were isolated.

\textit{Northern blot analysis.} Northern blot analysis was done as previously described (25). HepG2 cells were infected with adenovirus p53 (Ad-p53) or adenoaviruses green fluorescent protein (Ad-EGFP) at 30 multiplicity of infection (MOI), or two kinds of the p53-knockdown, and the control cell lines were treated with 1 \mu M Adriamycin (doxorubicin) for 2 h. mRNAs were isolated from the cells collected at the indicated times. Probes carrying the coding sequences of \textit{SEMA3F} p21/WAFI, and \textit{\beta}-actin were labeled with [\alpha-32P]dCTP using Megaprime DNA labeling system (GE Healthcare Bio-Sciences, Piscataway, NJ). The blots were hybridized with the radioactive probes at 42°C for 16 h, washed, exposed to Phosphor Screen, and visualized with Storm 860 (Molecular Dynamics, Sunnyvale, CA).

\textit{Western blot.} Western blot analysis was carried out as previously described (26). After treatment of each cell line with 1 \mu M Adriamycin, total cell lysates were prepared in radioimmunoprecipitation assay buffer [150 mmol/L NaCl, 50 mmol/L Tris-HCl (pH 8.0), 0.1% SDS, 1% NP40, protease inhibitor cocktail]. Anti-p53 (Ab-6; Calbiochem, San Diego, CA), anti-\textit{\beta}-actin antibody (AC-74; Sigma, St. Louis, MO), polyclonal anti-NRP2 antibody (C9; Santa Cruz Biotechnology, Santa Cruz, CA), and anti-FLAG (M2) monoclonal antibody (Sigma) were used in Western blot analysis.

\textit{Chromatin immunoprecipitation assay.} Chromatin immunoprecipitation assays were done using the chromatin immunoprecipitation assay kit from Upstate Biotechnology (Lake Placid, NY) as recommended by the manufacturer and as previously described (26). The anti-p33 monoclonal antibody (Ab12; Calbiochem) for immunoprecipitation and the anti-FLAG M2 monoclonal antibody (Sigma) as a negative control were used. H1299 (1.5 × 106) or HepG2 (1.5 × 106) cells were plated onto 10-cm dish and infected with Ad-p53 or Ad-EGFP at 30 MOI HCT116 p53(+/+) and HCT116 p53(−/−) cells (3 × 104) were treated with 1 \mu M Adriamycin to activate endogenous p53. The PCRs were done with the following specific primers: SEMA3F-p53BS1, 5'-AGATCGAAGCTTCTTGGAGG-3' and 5'-CTTGGTAT-CAAGCTGGAATAT-3'; SEMA3F-p53BS2, 5'-GGGAGAAGAAGAAGAGCAT-3' and 5'-GTTCTTTGTCGCCTGTCTTTT; NRP2-p53BS1, 5'-TCTGCGCATGTGATCC-3' and 5'-CTGGAGACTTTGAGATGTTCAACTA; NRP2-p53BS2, 5'-GTTCGACGCGGAGTTTCTC-3' and 5'-TGAGCCGTGAGTCCCTATCTC-3'; and p21, 5'-ACCCACACATATTCCCTC-3' and 5'-GGCCACAGAAATAAGCCA-3'.

\textit{Luciferase assay.} The 172-bp fragment including p53BS1 and the 193-bp fragment of p53BS2, which were amplified with the same primers as in the chromatin immunoprecipitation assays, were cloned into pGL3 promoter vector (Promega, Madison, WI) and sequenced. For NRP2, the 249-bp fragment including p53BS1 was cloned into pGL3 promoter vector. Reporter assay was done as previously described (26). p53 family genes, such as p73 and p63, and four expression constructs (p8C/CMV-p73sA, p8C/CMV-p73sB, p8C/CVM-p63sA, and p8C/CVM-p63sB; refs. 27, 28) were a gift from Shuntaro Ikawa (Tohoku University, Sendai, Japan). pGL3 vector with p53BS for p21 was used for positive control.

\textit{Plasmid construction.} The entire coding region of \textit{SEMA3F} amplified by reverse transcription-PCR (RT-PCR) was cloned into pcROI and cloned into pcDNA3.1(+) (Invitrogen, Carlsbad, CA) and sequenced. The fragment containing the whole \textit{SEMA3F} sequence was digested with EcoRI and cloned into pcDNA3.1(+) (Invitrogen) to prepare sense-strand \textit{SEMA3F} (pcDNA-SEMA3F-S) and antisense-strand \textit{SEMA3F} (pcDNA-SEMA3F-AS) for colony formation assay, or into pIRE5-EGFP (BD Biosciences, San Jose, CA) for stable transformants expressing the EGFP signal that represents the expression level of \textit{SEMA3F} mRNA. The entire \textit{SEMA3F} cDNA with EcoRI and Xba1 sites was amplified by PCR, sequenced, and then cloned into p3XL-FLAG-CMV-14 expression vector (Sigma), which contains three tandems of FLAG peptides at the COOH terminus for generating Ad-FLAG-SEMA3F. For NRP2-expressing vector, the cDNA clone containing the entire coding region of NRP2 was purchased from RZPD German Resource Center for Genome Research (Berlin, Germany). The entire coding region of NRP2, amplified by PCR, was also cloned into pcROI-Bl2-TOPO vector and sequenced. The fragment containing the whole NRP2 sequence was digested with Xba1 and HindIII and cloned into pcDNA3.1(−) for stable transformants.

\textit{Construction of recombinant adenoviruses for \textit{SEMA3F}.} Replication-deficient recombinant viruses Ad-SEMA3F and Ad-3\times\textit{FLAG}-SEMA3F were generated and purified as previously described (29). In brief, blunted-ended \textit{SEMA3F} and 3\times\textit{FLAG}-tagged \textit{SEMA3F} containing 3\times\textit{FLAG} at a COOH terminus of \textit{SEMA3F} were cloned into the Smul site of the cosmid pAxCawlt (Takara, Otsu, Japan), which contains the CAG promoter and the entire genome of type 5 adenovirus except for E1 and E3 regions. BspTiOH-digested cosmids were transfected to 293 (human embryonic kidney cell line) cells. Viruses were propagated in 293 cells and purified as previously described. Expression levels of \textit{SEMA3F}-expressing adenoviruses were evaluated by Western blot for Ad-3\times\textit{FLAG}-SEMA3F or by RT-PCR for Ad-SEMA3F at 24 h postinfection at the indicated MOIs.

\textit{Cell viability assay.} To examine the role of the \textit{SEMA3F-NRP2} pathway in cell growth, we examined cell viability using trypan blue exclusion method. Parental H1299 cells and the NRP2-stable transfector cells (s-1; 1.5 × 105) were plated onto six-well culture plates and infected with either Ad-SEMA3F or Ad-3\times\textit{FLAG}-SEMA3F at 0, 10, 50, 60, and 100 MOIs. Ad-EGFP was used as a negative control. Seventy-two hours later, all the cell components were collected and viable cells were counted by hemocytometer.

\textit{Nude mice assay.} Nude mice assay was done by inoculating the \textit{SEMA3F} stable transfactors and LS174T-p53-si cells. For \textit{SEMA3F} stable transfactors, two independent clones (I-s11 and I-s17) and a control cell line (IRES1) were inoculated s.c. into the bilateral thoracic and abdominal walls (total four parts per mouse) of 7-8-week female BALB/c Ncl-nu mice (CLEA Japan, Tokyo, Japan) at 1 × 106 per mouse in a volume of 0.1 mL. LS174T-p53-si and LS174T-control cells were also inoculated at 2 × 106 per mouse. Then the mice were maintained under specific pathogen-free conditions. The tumor volume was measured with a caliper twice a week and assessed by the formula (width × width × length)/2. After 2 to 5 weeks of observation, explanted tumors were isolated and analyzed genetically and immunohistochemically. These animal experiments were repeated twice to confirm the similar results. These animal experimental protocols were approved by the Committee for Ethics of Animal Experimentation and the experiments were conducted in accordance with the Guidelines for Animal Experiments in the National Cancer Center.
Semiquantitative RT-PCR. The excised tumors were homogenized and total RNAs were extracted by TRIzol. cDNAs were synthesized from 3 μg of total RNA with SuperScript First-Strand Synthesis System (Invitrogen). The RT-PCR was run in the exponential region (19–30 cycles) to allow semiquantitative comparisons among cDNAs developed from identical reactions. Each PCR regimen involved a 94°C, 5 min initial denaturation step followed by 19 cycles ([j2-MG]), 21 cycles (SEMA3F, BAI1, and VEGF), 22 cycles (p53) and p21/WAF1), 24 cycles (TSP1), 25 cycles (NRP1), 30 cycles (NRP2), 55°C for 30 s, and 72°C for 30 s on a GeneAmp PCR system 9700 (Applied Biosystems, Foster City, CA).

Immunohistochemistry. The frozen sections (6 μm) from the resected tumors embedded into optimum cutting temperature (OCT) compound (Sakura Fine Technical, Tokyo, Japan) were stained by immunoperoxidase procedure. Briefly, after air dry, each section was fixed with cold acetone, treated with 0.3% H2O2 for 10 min, and then blocked with 10% goat serum. Antimouse CD31 (MEC13.3; BD Biosciences) was used as the first antibody (1:100) and incubated at room temperature for 1 h. Then Histofine Simplestain MAX-PO (rat; Nichirei, Tokyo, Japan) was incubated at room temperature for 30 min as the second antibody and visualized with 3-amino-9-ethylcarbazole substrate (Nichirei). The consecutive sections corresponding to mCD31 staining sections were used for H&E staining. Light images of immunofluorescent tissues were viewed under an Olympus IX71 microscope (Olympus, Tokyo, Japan) and digitized using Olympus DP70 camera and DP70-BSW software (Olympus).

Tumor vessel density. For analysis of vessel density, 8 to 10 independent tumors were randomly selected from each group (I-s11, I-s17, IRES1, LS174T-p53-siRNA, and LS174T-control). CD31-positive areas in each field were compared. Vessel densities were calculated as the ratio of the CD31-positive area divided by the total area (30, 31).

Statistical analysis. Colony numbers in colony formation assay, tumor volume, tumor weight, and tumor vessel density in nude mice assay were analyzed by Student’s t test.

Figure 1. SEMA3F as a p53-inducible gene. A, exogenous p53-dependent expression of SEMA3F. mRNAs isolated at the indicated times from the HepG2 cells infected with Ad-p53 or Ad-EGFP at an MOI of 30 were subjected to Northern blot analysis for the SEMA3F gene. p21/WAF1 and β-actin were used as positive and loading controls, respectively. B, expression of p53 protein. The cell lysates isolated at the indicated times from p53-siRNA and control cells, which were derived from either HepG2 cells or LS174T cells, were subjected to Western blot analysis for p53. β-actin was used as a loading control. C, endogenous p53-dependent expression of SEMA3F. mRNAs isolated at the indicated times from p53-siRNA and control cells (HepG2 or LS174T), which were treated with 0.2 μg/mL Adriamycin, were subjected to Northern blot analysis for the SEMA3F gene. p21/WAF1 and β-actin were used as positive and loading controls, respectively.

Results

Identification of SEMA3F as a novel p53-inducible gene. The emerging evidence suggested the role of axon guidance molecules in tumorigenesis (4, 32). In addition, semaphorin 3B (SEMA3B) was previously reported to be regulated by tumor suppressor p53 (33). Therefore, we reasoned that SEMA3F may be involved in the p53-regulated transcriptional network. To test this hypothesis, we first carried out Northern blot analysis of the SEMA3F gene. mRNAs were subjected to Northern blot analysis, which were obtained at the indicated time after infection from the HepG2 cells, a hepatocellular carcinoma cell line, infected with adenovirus p53 (Ad-p53) or adenovirus EGFP (Ad-EGFP). As shown in Fig. 1A, a 3.8-kb transcript of SEMA3F was detected, and the expression was increased in the cells infected with Ad-p53, whereas the increase was very small in the Ad-EGFP-infected cells, suggesting that exogenous p53 can induce the expression of SEMA3F. Clear induction was also observed in the case of the p21/WAF1 gene, one of the most well-known p53-inducible genes (Fig. 1A).

To further validate the p53 dependency of the SEMA3F expression, we examined whether endogenous p53 activates the transcription of SEMA3F. We established p53-knockdown cell lines derived from HepG2 cells and LS174T cells (a colorectal cancer cell line). As indicated in Fig. 1B, the expression of endogenous p53 protein was increased after treatment with 1 μg doxorubicin in HepG2-control and LS174T-control cells, whereas negligible p53 expression was detected in HepG2-p53-siRNA and LS174T-p53-siRNA cells. Similarly, the expression of SEMA3F mRNA was induced in both the control cells, but its expression was not increased in both the p53-siRNA cells (Fig. 1C). These results, when considered together, indicated that the transcription of
SEMA3F is activated by endogenous and exogenous p53 and that SEMA3F is a novel p53-inducible gene.

**SEMA3F as a direct transcriptional target of p53.** To determine whether SEMA3F is a direct target gene of p53, we investigated p53-binding sequence(s) (p53BS) in the genomic region and found two candidate sequences, BS1 and BS2, in intron 1 of SEMA3F (Fig. 2A). Therefore, we did a chromatin immunoprecipitation assay. The protein complex, including p53 and the chromatin-associated DNA fragments, was immunoprecipitated with the anti-p53 antibody and the precipitated DNA was subjected to PCR analysis with the specific primers for BS1 and BS2. As shown in Fig. 2C, the sequence of BS1, but not BS2, was clearly amplified by PCR with the protein complex immunoprecipitated from H1299 cells infected with Ad-p53 or from HCT116-p53(+/+) cells treated with 0.2 μg doxorubicin, implying that both endogenous and exogenous p53 interact with BS1.

To determine whether BS1 actually has p53-dependent transcriptional activity, we carried out a reporter assay. A 200- to 300-bp DNA fragment containing BS1 or BS2 was cloned upstream of the SV40 promoter of pGL3-promoter vector (pGL3-BS1 or pGL3-BS2). As indicated in Fig. 2, when pGL3-BS1 was cotransfected with wild-type p53-plasmid, but not mutant p53-plasmid, the luciferase activity was remarkably enhanced. In addition, it is noted that p73 also activated the transcription of pGL3-BS1 (Fig. 2D). These results suggest that SEMA3F is a bona fide target gene of p53 and possibly p73.

**The role of SEMA3F in cell growth.** To examine the role of SEMA3F in *in vitro* cell growth, we first did colony formation assay. As shown in Supplementary Fig. S1A and B, overexpression of sense SEMA3F caused significant reduction in the H1299 cell colony number as compared with overexpression of antisense SEMA3F, whereas no significant difference was observed between sense SEMA3F– and antisense SEMA3F–transfected T98G cells. The difference in the results between H1299 and T98G is likely to be due to the expression level of NRP2, which is a functional receptor for SEMA3F, because the expression level of NRP2 in H1299 was higher than that of T98G (data not shown).

Although significant growth suppression by SEMA3F was observed, the effect seemed to be mild. Therefore, we attempted to establish stable H1299 cell transformants, which consistently express high levels of SEMA3F. The sense sequence of SEMA3F was cloned into the pRES2-EGFP vector, which has the internal ribosomal site of the encephalomyocarditis virus, between the multiple cloning site and the *EGFP* gene to be translated from a single bicistronic mRNA. Therefore, it is possible to monitor the expression level of SEMA3F by examining the *EGFP* signal. Eventually, we selected two independent clones, pRES-SEMA3F-11 (l-s11) and pRES-SEMA3F-17 (l-s17). As indicated in Supplementary Fig. S2A, homogenous and high levels of green signals of *EGFP* were detected in these two clones. Consistent with the result of *EGFP*, the high expression of SEMA3F mRNA was also detected by RT-PCR (Supplementary Fig. S2B). As seen in Supplementary Fig. S1A, the high and stable expression of SEMA3F inhibited the *in vitro* cell growth to some extent, but the effect was not dramatic (Supplementary Fig. S2C).

**The role of SEMA3F in tumor growth.** To investigate the role of SEMA3F in tumor growth, three types of the established cell lines, including control, pRES-SEMA3F-11 (l-s11), and pRES-SEMA3F-17 (l-s17), were s.c. implanted on the flank of nude mice. Then, the tumor volume of each tumor was measured daily for 2 weeks until it exceeded 1500 mm³ (Fig. 2). The difference in the results between H1299 and T98G is likely to be due to the expression level of NRP2, which is a functional receptor for SEMA3F, because the expression level of NRP2 in H1299 was higher than that of T98G (data not shown).

**Figure 2.** Identification of SEMA3F as a direct target gene of p53. A, potential p53-binding sequences. Two potential p53-binding sequences, BS1 and BS2, are located in intron 1 of SEMA3F. B, binding of p53 with BS1 or BS2 of SEMA3F. Chromatin immunoprecipitation assay was done for the DNA-protein complex, which was immunoprecipitated by anti-p53 antibody from H1299 cells infected with Ad-p53 or Ad-EGFP at an MOI of 30, or from HCT116-p53(+/-) cells or HCT116-p53(–/–) cells treated with 0.2 μg/mL Adriamycin. p21/WAF1 was used as a positive control. C, p53-dependent transcriptional activity of BS1. The heterologous luciferase reporter plasmid containing BS1 or BS2 was cotransfected with the plasmid designed to express wild-type p53 (p53WT), mutant p53 (p53MT), or no p53 (mock) into H1299 cells. The luciferase activity 24 h after transfection is shown in relation to the activity of the pGL3-promoter vectors without p53BSs. D, p73-dependent transcriptional activity of BS1. The heterologous luciferase reporter plasmid containing BS1 was cotransfected with the plasmid designed to express wild-type p53, p73α, p73β, p73γ, or no p53 (mock) into H1299 cells. The luciferase activity 24 h after transfection is shown in relation to the activity of the pGL3-promoter vectors without p53BSs.
twice a week for 32 days. As clearly indicated in Fig. 3A, the tumors derived from the control cell line grew exponentially, whereas the growth of the tumors from two independent SEMA3F-expressing clones was severely impaired. Moreover, the tumors derived from pIRES-SEMA3F-11 (l-s11) and pIRES-SEMA3F-17 (l-s17) cell lines were clearly much smaller in size and weight than those derived from the control cell line (Fig. 3B and C). The comparison between the results of Supplementary Figs. S1 and S2 and those of Fig. 3 indicated that the in vivo effect of SEMA3F on tumor growth was much stronger than the in vitro effect on cell growth, implying that SEMA3F might play an additional role in vivo.

The antiangiogenic role of SEMA3F in tumor growth suppression. SEMA3F was previously reported to play a role in developmental and tumor-associated angiogenesis (34–38). Therefore, we postulated that the remarkable effect on the in vivo tumor growth might be related to the regulation of tumor angiogenesis.

Frozen sections from the tumors embedded in OCT compound were stained with mCD31, which is a specific vascular marker. As indicated in Fig. 3D, the number of tumor vessels in control tumors was much more than that in SEMA3F-expressing tumors (Fig. 3E). Moreover, the quantitative analysis of the area of tumor blood vessels stained with anti-CD31 antibody was measured in the intratumoral region. Columns, average area of tumor vessels for each group; bars, SE. P < 0.01 was deemed statistically significant. D, immunohistochemistry for CD31 expression. Frozen sections of the isolated tumors were subjected to immunohistochemical staining with anti-CD31 antibody and H&E staining. Representative result from each group (×200).

Figure 3. In vivo tumor growth suppression by SEMA3F. A, tumor volume. The tumor volume of each tumor derived from pIRES-control (IRES1), pIRES2-SEMA3F-11 (l-s11), or pIRES-SEMA3F-17 (l-s17) stable clone was measured twice every week following the establishment of xenografts in BALB/c A/Jc-nu nude mice. Bars, SE. B, macroscopic appearance. The tumors were isolated at day 32 from each group. C, tumor weight. Columns, average tumor weight for each group; bars, SE. P < 0.01 was deemed statistically significant. D, immunohistochemistry for CD31 expression. Frozen sections of the isolated tumors were subjected to immunohistochemical staining with anti-CD31 antibody and H&E staining. Representative result from each group (×200). E, measurement of the area of tumor blood vessels. The area of tumor blood vessels stained with anti-CD31 antibody was measured in the intratumoral region. Columns, average area of tumor vessels for each group; bars, SE. P < 0.01 was deemed statistically significant.

p53 negatively regulates tumor angiogenesis probably via the SEMA3F-NRP2 pathway. Although p53 has been suggested to play an important role in the inhibition of tumor angiogenesis, there exists little evidence to support this assumption. Therefore, we evaluated whether p53 is actually involved in the regulation of tumor angiogenesis. LS174T-control and LS174T-p53-siRNA cells were s.c. inoculated on the flank of nude mice and allowed
to form tumors. As shown in Fig. 4A, tumors derived from LS174T-p53-siRNA cells attained a volume of ~700 mm³ by day 11 after inoculation, which is two times greater than the tumors from LS174T-control cells. Interestingly, it is absolutely evident that the number and area of tumor vessels in p53-knockdown tumors are much more and greater than those of tumor vessels in control tumors containing normal levels of p53 (Fig. 4B and Supplementary Fig. S3). The results clearly show that p53 is definitely involved in the suppression of tumor vessel formation.

To evaluate the role of antiangiogenic p53 target genes, including TSP1, BAI1, SEMA3F, and NRP2, in this phenomenon, we examined the expression levels of these genes and several related genes in the tumors isolated at day 11. Interestingly, although the cells and tumors were not exposed to any genotoxic stress, the expression of p53 mRNA was significantly elevated in the control tumors containing wild-type p53 but not in the p53-siRNA tumors (Fig. 4C). Consistent with the expression level of p53, p21/WAF1 mRNA levels in the control tumors were higher than those in the p53-siRNA tumors. The expression levels of SEMA3F in control tumors seemed to be slightly but not remarkably higher than those in p53-knockdown tumors (Fig. 4C). On the other hand, the expression of TSP1 was not detected in either tumors, and BAI1 expression levels in control tumors were lower as compared with those in p53-knockdown tumors (Fig. 4C). Surprisingly, the expression of NRP2, the functional receptor for SEMA3F, was clearly elevated in control tumors as compared with p53-knockdown tumor (Fig. 4C).

To validate the role of SEMA3F in p53-regulated antiangiogenesis of tumors, we expressed exogenous SEMA3F in p53-knockdown tumors. As shown in Supplementary Fig. S4, p53-knockdown tumors infected with Ad-SEMA3F revealed significant reductions in tumor size and tumor vessel density. Thus, these results support the notion that SEMA3F plays an important role in p53-regulated angiogenesis suppression.

Identification of NRP2 as a direct transcriptional target of p53. Because the expression levels of NRP2 mRNA were severely impaired in p53-knockdown tumors (Fig. 4C), we speculated that NRP2 expression might be directly regulated by p53. To examine whether NRP2 is a direct transcriptional target of p53, we did several experiments (Fig. 5). As shown in Fig. 5A, the expression of NRP2 mRNA was induced in response to DNA damage in a p53-dependent manner. Furthermore, we have found a p53-binding sequence (BS1) in promoter of the NRP2 gene (Fig. 5B), and the sequence of BS1 interacted with p53 in the cells infected with Ad-p53 (Fig. 5C). Moreover, the sequence of BS1, but not BS2 (a negative control), actually had p53-dependent transcriptional activity in reporter assay (Fig. 5D). Interestingly, p63ß also activated to some extent the transcription of the luciferase-containing BS1 (Fig. 5E). These results suggest that NRP2 is a direct target gene of p53 and that p63ß may be involved in the regulation of NRP2 expression.

The role of the SEMA3F-NRP2 signaling pathway in cell growth suppression. Based on the results of Figs. 1, 2, and 5, it is likely that p53 is directly involved in the transcription of both the...
ligand SEMA3F and the receptor NRP2. Therefore, we assumed that the SEMA3F-NRP2 signaling pathway may play a role in cell growth control in vitro. To validate this assumption, we established the stable H1299 clone expressing NRP2 and prepared the adenovirus vector designed to express SEMA3F (Ad-SEMA3F) or FLAG-SEMA3F (Ad-FLAG-SEMA3F). As shown in Fig. 6A, the stable H1299 clone (s-11) highly expressed NRP2 whereas parental H1299 cells displayed low level of NRP2. Moreover, s-11 cells infected with Ad-SEMA3F or Ad-FLAG-SEMA3F revealed significant levels of SEMA3F mRNA or protein, respectively, in a dose-dependent manner (Fig. 6B).

By using these materials, we examined the effect of SEMA3F and NRP2 on in vitro cell growth rate. As expected, overexpression of EGFP did not induce any remarkable change of the cell growth rate of either parental or s-11 cells (Fig. 6C). However, the cells infected with Ad-SEMA3F or Ad-FLAG-SEMA3F clearly showed cell growth suppression in a dose-dependent manner (Fig. 6C). In addition, consistent with the result of Supplementary Fig. S1, overexpression of SEMA3F induced a maximum of 50% to 80% reduction in cell number in parental cells, whereas cell number in the s-11 clone infected with Ad-SEMA3F or Ad-FLAG-SEMA3F decreased to <10% of the control cells (without infection of Ad-SEMA3F or Ad-FLAG-SEMA3F; Fig. 6C). These results suggest that the SEMA3F-NRP2 signaling pathway may play a significant role in cell growth suppression as well as inhibition of tumor angiogenesis (Fig. 6D).

Discussion

Although SEMA3F was initially identified as a candidate tumor suppressor gene at chromosome 3p21.3, the causative point mutations in the SEMA3F gene are very rare in human cancers (17–19). This casts a doubt on the role of SEMA3F as a tumor suppressor gene. However, accumulating evidence continues to support the potential role of SEMA3F in tumorigenesis. For example, overexpression of SEMA3F in the mouse fibrosarcoma and human ovarian adenocarcinoma cell lines caused tumor growth suppression (23). This effect was confirmed by two studies. In the first study, ectopic expression of SEMA3F in the human embryonic kidney 293 cell line inhibited tumor formation in nude mice.
mice and the resulting tumors displayed reduced density of tumor vessels, implying the role of SEMA3F in antiangiogenesis during tumor formation (39). The antiangiogenic activity of SEMA3F was also confirmed by both the in vitro cell growth inhibition assay for the human umbilical vein endothelial cells and the in vivo antiangiogenesis assay in this study (39). In the second study, the enforced expression of SEMA3F in highly metastatic human melanoma cells caused the phenotypic change from a highly
metastatic phenotype to a nonmetastatic and benign tumor–like phenotype, which is characterized by a large area of apoptosis, diminished vascularity, inhibition of hyperplasia in overlying epidermal cells, and encapsulated tumor borders delineated by thick layers of fibroblasts and collagen matrix (40). These observations suggest that SEMA3F definitely plays a role in preventing tumor progression through a common mechanism of angiogenesis. In the present study, we have also shown that overexpression of SEMA3F in a lung cancer cell line, H1299, caused a remarkable tumor growth suppression in vitro, and that this effect is likely to be mediated, at least in part, by the inhibition of tumor vessel formation. Therefore, consistent with the previous observations, we assume that SEMA3F indeed plays a role as a tumor suppressor gene by the down-regulation of tumor angiogenesis.

Thus far, two important mediators for antiangiogenesis, TSP1 and BA11, were reported to be the transcriptional target genes of p53 (6, 7). In addition to these molecules, we have found that SEMA3F is the third antiangiogenic target gene. To determine whether TSP1, BA11, and SEMA3F play some role in p53-regulated antiangiogenesis, we examined the expression levels of these genes in control LS174T tumors and p53-knockdown LS174T tumors. As shown in Fig. 4C, TSP1 expression was not detected in either control or p53-knockdown tumors, and BA11 levels in control tumors were lower than those in p53-knockdown tumors. Therefore, TSP1 and BA11 were unlikely to be involved in p53-regulated antiangiogenesis in our system that used the colorectal cancer cell line LS174T. On the other hand, control LS174T tumors displayed slightly higher expression levels of SEMA3F as compared with p53-knockdown tumors. These results suggest that, at least in our system that used LS174T colorectal cancer, SEMA3F may play some role in tumor angiogenesis suppression and that SEMA3F represents a new mediator for p53-regulated antiangiogenesis.

The surprising finding is that the expression of NRP2, the functional receptor for SEMA3F, depended on the status of p53 in the tumors (Fig. 4C). Indeed, we have found that the NRP2 gene has a p53-responsive and binding sequence and that the expression of NRP2 is inducible in response to exogenous and endogenous p53, implying that NRP2 is also a p53 target gene (Fig. 5). This observation gives rise to several hypotheses. First, p53 may regulate both SEMA3F and its receptor NRP2, thereby the p53-activated SEMA3F is secreted and binds to NRP2 present on the cell secreting SEMA3F, leading to cell growth suppression in an autocrine fashion. Second, the secreted SEMA3F binds to NRP2 on vascular endothelial cells that express NRP2 on their surface, leading to inhibition of tumor vessel formation in a paracrine fashion. Third, the secreted SEMA3F binds to NRP2 on the surface of neighboring tumor cells, leading to growth suppression of tumor cells. Fourth, p53 may function as a bystander to express and secrete SEMA3F, which binds to NRP2 on the surface of the target cancer cell and induces growth suppression of the target cells. Fifth, again p53 may function as a bystander to express NRP2 on the surface of vascular endothelial cells that bind to the tumor-secreted SEMA3F, leading to antiangiogenesis under the control of p53. Several other hypotheses may arise as a result of further studies. Consistent with these assumptions, the SEMA-NRP2 pathway, in fact, controlled not only tumor angiogenesis but also the in vitro cell growth of cancer cells (Fig. 6C). In any case, p53 would regulate both cell growth itself and tumor angiogenesis in an autocrine or paracrine fashion (Fig. 6D). The role of p53 as a bystander in the surrounding cells including cancer cells, endothelial cells, fibroblast cells, immune cells, etc., would be considerably important for the prevention of tumor progression via the SEMA3F-NRP2 pathway.

In our in vivo assay, SEMA3F revealed a more remarkable inhibitory effect on tumor growth compared with the in vitro assay. According to the in vitro result on the same H1299 cells (Supplementary Figs. S1 and S2), the effect was likely mediated by the communication between the secreted SEMA3F protein and its receptor NRP2 expressed in the surrounding cells, including epithelial cells and fibroblasts. Consistent with this notion, the parental H1299 cells do not express high levels of NRP2 protein (Fig. 6A). Because we used the human SEMA3F protein in the mouse system, this will raise one important question about whether human SEMA3F can interact with mouse NRP2 receptor. However, the sequence homology reveals >96% identity between human and mouse SEMA3F protein. In addition, it shows >94% identity between human and mouse NRP2 protein. Based on the very high homology of SEMA3F or NRP2 between human and mouse, we assume that human SEMA3F probably interacts with mouse NRP2 in our assay. Thus, we think that the results in the in vivo assay support the paracrine model for the SEMA3F-NRP2 pathway (Fig. 6D).

In contrast to SEMA3B, which is the neighboring SEMA member at 3p21.3 and whose function is involved in apoptosis (33, 41), many lung cancers continue to express SEMA3F in spite of p53 mutations (18). This seems to be the long-lasting issue for the role of SEMA3F in the 3p21.3 tumor suppressor genes. However, our two findings provide some clue to address this question. First, the expression of SEMA3F is likely regulated not only by p53 but also by p73, the function of which is usually normal in human lung cancers. This implies that inactivation of p53 does not necessarily lead to the total down-regulation of SEMA3F expression. Second, it seems that the expression of NRP2 is very critical for the SEMA3F-NRP2 autocrine pathway (Fig. 6D). In our data, the expression of NRP2 was more dependent on p53 in the tumors (Fig. 4C) and NRP2 was also a direct target gene of p53 (Fig. 5). Therefore, not SEMA3F but NRP2 might be the key target for lung cancer tumorigenesis via the p53-SEMA3F-NRP2 pathway. To clarify the precise regulatory mechanism for the SEMA3F-NRP2 pathway, further investigation is needed for the role of NRP2 in human primary lung cancers.

How does SEMA3F suppress tumor progression and metastasis? The precise mechanism for the SEMA3F-dependent pathway still remains largely unclear. However, SEMA3F may be involved in the regulation of angiogenesis and metastasis. In SEMA3F-regulated antiangiogenesis, the target cell that expresses NRP2 is likely to play a critical role in the antiangiogenic activity. Thus far, the expression of NRP2 has been found in vascular endothelial cells (42), lymphatic endothelial cells (43), and possibly cancer cells. Therefore, vascular endothelial cells or lymphatic endothelial cells may be SEMA3F target cells. However, recent observations have provided evidence that the recruitment of bone marrow–derived endothelial precursor cells contributes to tumor vasculature (44–46). Although it is still unclear whether bone marrow–derived endothelial precursor cells express the SEMA3F receptor(s), including NRP2, it will be interesting to investigate whether SEMA3F suppresses the recruitment of bone marrow–derived endothelial precursor cells to the site of tumor vessel formation. Moreover, in cancer metastasis, the previous reports clearly showed that SEMA3F prevents in vivo tumor metastasis (35, 40) and that SEMA3F inhibits in vitro cell
attachment and motility (47, 48). Very recently, a study reported that the premetastatic niche initiated by vascular endothelial growth factor receptor 1–positive hematopoietic bone marrow progenitor cells (VHBMPC) is very critical for the establishment of cancer metastasis at the metastatic site (49). We speculate that SEMA3F might prevent cancer metastasis by inhibiting the recruitment of VHBMPCs at the premetastatic niche. Therefore, it is also interesting to investigate whether VHBMPCs express the functional SEMA3F receptor(s) and whether SEMA3F actually inhibits cell attachment, motility, and proliferation of VHBMPCs.

We believe that further investigation on the SEMA3F-NRP2 pathway will clarify the mechanism for the p53-regulated communication between cancer cells and the surrounding cells and the mechanism for the SEMA3F-regulated pathways of antiangiogenesis and antimetastasis. In addition, application of the SEMA3F pathway will definitely initiate the development of novel strategies for cancer therapy.

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