Sprouty2 Association with B-Raf Is Regulated by Phosphorylation and Kinase Conformation

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Abstract

Sprouty2 is a feedback regulator that controls the Ras/Raf/MEK/extracellular signal-regulated kinase mitogen-activated protein kinase (MAPK) pathway at multiple levels, one way being through direct interaction with Raf kinases. Consistent with a role as a tumor suppressor, Sprouty2 expression is often down-regulated in human cancers. However, Sprouty2 is up-regulated in some cancers, suggesting the existence of posttranscriptional mechanisms that permit evasion of Sprouty2-mediated antitumorigenic properties. We report that MAPK activation induces Sprouty2 phosphorylation on six serine residues, which reduced Sprouty2 association with wild-type B-Raf. Mutation of these six serines to nonphosphorylatable alanines increased the ability of Sprouty2 to inhibit growth factor–induced MAPK activation. Oncogenic B-Raf mutants such as B-Raf V600E did not associate with Sprouty2, but this resistance to Sprouty2 binding was not due to phosphorylation. Instead, the active kinase conformation induced by oncogenic mutation prevents Sprouty2 binding. These results reveal a dual mechanism that affects the Sprouty2/B-Raf interaction: Sprouty phosphorylation and B-Raf conformation. [Cancer Res 2009;69(17):6773–81]

Introduction

Sprouty was identified in a Drosophila screen for modifiers of airway branching induced by fibroblast growth factor (FGF) signaling (1). Four mammalian Sprouty orthologues (Sprouty1–4; Sprouty2 being the closest to Drosophila Sprouty; refs. 1–4) comprise a family of receptor tyrosine kinase feedback regulators that modulate the Ras/Raf/MEK/extracellular signal-regulated kinase (ERK) mitogen-activated protein kinase (MAPK) pathway (5). As a consequence, they regulate processes such as proliferation, survival, and motility in response to receptor tyrosine kinase activation and influence the development of many tissues (6).

Clinical and experimental evidence is consistent with Sprouty proteins being tumor suppressors. When overexpressed, Sprouty proteins suppressed proliferation of cultured tumor cells (7–9) and urethane-induced lung cancer in mice (10). Conversely, small interfering (siRNA)–mediated Sprouty2 knockdown induced melanocyte proliferation (11), whereas Sprouty2 deletion enhanced K-Ras–induced lung cancer in mice (12). If Sprouty genes were tumor suppressors, expression would be expected to be repressed or lost in cancers, which has been observed in a variety of tumors (13–19). However, Sprouty levels are elevated in melanoma cells expressing oncogenic B-Raf (11, 20, 21) or N-Ras (20) and in gastrointestinal stromal tumors expressing activated c-Kit (22, 23), suggesting that there may be posttranscriptional mechanisms to evade its tumor-suppressive effects.

Sprouty proteins are composed of a variable NH2 terminus and a highly conserved cysteine-rich COOH terminus (24), which regulate MAPK signaling at multiple levels depending on cell context and growth factor receptor activated. One way they inhibit MAPK activation is by directly interacting with and inhibiting activation of Raf kinases (11, 25–28). We reported previously that Sprouty2 did not associate with oncogenic B-Raf mutants, which allows for increased active ERK levels in melanoma cell lines expressing mutant B-Raf despite elevated Sprouty2 (11). In this study, we investigated the Sprouty2/B-Raf interaction and propose two factors that influence this association: Sprouty2 phosphorylation at specific sites and the B-Raf kinase domain conformation.

Materials and Methods

Plasmids and proteins. pEF Myc-B-Raf constructs and pcDNA3 FLAG-Sprouty2 were described previously (11). pcDNA3 HA B-Raf was from W. Kolch (Beatson Institute). pEX MEKEE was from C.J. Marshall (Institute for Cancer Research). For site-directed mutagenesis, FLAG-Sprouty2 was subcloned into the Gateway vector system (Invitrogen) and point mutations were introduced using the QuikChange kit (Stratagene) according to the manufacturer’s protocol.

Recombinant human wild-type and V600E glutathione-S-transferase (GST)–B-Raf catalytic domains were from Cell Signaling Technology or Millipore. FLAG-Sprouty2 was cloned into pGEX-4T3 (GE Healthcare) and expressed and purified using standard methods (29).

Alternatively, FLAG-Sprouty2 was subcloned into pGEX-6P3 (GE Healthcare) and expressed and purified as described above. Flag Sprouty2 was cleaved from GST by incubation with 80 units PreScission Protease (GE Healthcare) in cleavage buffer [50 mmol/L Tris (pH 7.5), 150 mmol/L NaCl, 1 mmol/L EDTA, 1 mmol/L DTT] for 4 h at 4°C.

In vitro transcribed and translated [35S]methionine-labeled HA B-Raf was generated from pcDNA3 HA B-Raf using the TNT T7 Quick Coupled Transcription Translation kit (Promega) according to the manufacturer’s protocol.

Cell culture and transfections. HEK293 cells were cultured in DMEM supplemented with 10% (v/v) fetal bovine serum, 100 units/ml penicillin, and 100 μg/ml streptomycin (Invitrogen) at 37°C and 5% CO2. Cells were transfected with FuGENE6 (Roche) according to the manufacturer’s protocol and harvested 24 h later. Amounts of wild-type and phosphorylation site mutant Sprouty2 plasmids transfected were adjusted to equilibrate protein levels.

Parental NIH3T3 cells were maintained in DMEM supplemented with 10% (v/v) donor calf serum (Invitrogen). Details of generation of Tet-Off Sprouty2 cell lines are in Supplementary Methods. Mouse embryo fibroblasts were maintained in DMEM with 10% (v/v) fetal bovine serum,
100 units/mL penicillin, and 100 μg/mL streptomycin at 37°C and 10% CO₂, and cell lysates were prepared as described in ref. 30.

**Cell extraction and immunoblotting.** Whole-cell lysates (WCL) were prepared and Western blotted as described previously (11). Primary antibodies used were as follows: murine anti-FLAG (Sigma-Aldrich), rabbit anti-Myc 9B11 (Cell Signaling Technology), rabbit anti-pERK1/2 (Cell Signaling Technology), rabbit anti-ERK1/2 (Millipore), rabbit anti-GST (Cell Signaling Technology), rabbit anti-Sprouty (Upstate Biotechnology), and rabbit anti-MEK (Cell Signaling Technology). Alexa Fluor 680 (Molecular Probes) or IRDye800 (Rockland)–conjugated secondary antibodies were detected by infrared imaging (Li-Cor Odyssey).

**Immunoprecipitations.** Myc-B-Raf complexes were immunoprecipitated from HEK293 WCL and analyzed for associated FLAG-Sprouty2 as described previously (11). Detailed methods can be found in Supplementary Methods.

**In vivo cell labeling.** HEK293 cells transiently expressing FLAG-Sprouty2 and Myc-B-Raf V600E were cultured overnight in phosphate-free DMEM supplemented with 0.2 mCi/mL 32P (Amersham Biosciences). Cells were lysed in Tris lysis buffer [50 mmol/L Tris (pH 7.5), 150 mmol/L NaCl, 1% Triton, 1 mmol/L Na₃VO₄, 50 mmol/L NaF, 2 mmol/L β-glycerophosphate, 2 mmol/L EDTA, 1 mmol/L DTT, 1 mmol/L phenylmethylsulfonyl fluoride, and 1× Complete protease inhibitor cocktail (Roche)]. FLAG-Sprouty2 was immunoprecipitated from WCL and analyzed by SDS-PAGE followed by Coomassie staining with Simply Blue SafeStain (Invitrogen) and autoradiography.

**Identification of Sprouty2 phosphorylation sites by mass spectrometry.** FLAG-Sprouty2 was immunoprecipitated from HEK293 cells, eluted with Laemmli buffer, separated by SDS-PAGE, and detected by staining with Brilliant Blue G Colloidal Coomassie (Sigma). Sprouty2 bands were excised, destained with successive ammonium bicarbonate and acetone nitrite washes, and digested with trypsin as described previously (31). Extracted tryptic peptides were analyzed by liquid chromatography-mass...
spectrometry with precursor of 79 scanning as described previously (32). Detailed methods can be found in Supplementary Methods.

**Phosphorylation of recombinant Sprouty2 in B-Raf V600E cell lysate.** GST-FLAG-Sprouty2 (2 µg) immobilized on glutathione beads was incubated with 0.74 MBq [γ-32P]ATP, 200 µmol/L ATP, 10 µg/mL creatine kinase, and 5 mmol/L creatine phosphate in 2.5 mg WCL from HEK293 cells transiently expressing Myc-B-RafV600E at 30°C for 2 h. Beads were washed 5× in Tris/NP-40 [50 mmol/L Tris (pH 8), 150 mmol/L NaCl, 1.0% (v/v) NP-40, 1 mmol/L phenylmethylsulfonyl fluoride, 1 mmol/L DTT, and Complete protease inhibitor cocktail] and GST-FLAG-Sprouty2 was eluted in Laemmli buffer, separated by SDS-PAGE, and analyzed by Coomassie staining and autoradiography.

**Peptide arrays.** The 23-mer peptides offset by 2 amino acids that covered the mouse Sprouty2 sequence were arrayed onto membranes (Cancer Research UK). Arrays were blocked in 5% (w/v) bovine serum albumin/TBS for 1 h at room temperature before incubation with 50 µL [35S]methionine-labeled in vitro transcribed and translated HA B-Raf in 5% (v/v) bovine serum albumin/TBS for 1 h at room temperature. Membranes were washed three times in TBS/0.1% (v/v) Tween, allowed to dry, and exposed to film.

**Pull-down of FLAG-Sprouty2 with GST-B-Raf.** Recombinant soluble GST or GST-B-Raf catalytic domain (0.5 µg) was incubated with 1 µg recombinant FLAG-Sprouty2 in 50 µL Tris/NP-40 at 30°C for 30 min. The reaction was diluted with 250 µL Tris/NP-40 and GST-B-Raf complexes were affinity purified on glutathione Sepharose 4B beads at 4°C for 2 h. Beads were washed 5× in Tris/NP-40 and complexes were eluted in Laemmli buffer and analyzed by SDS-PAGE as described above.

**B-Raf siRNA.** NIH3T3 cells were transfected with siRNA against B-Raf (siRNA1 Invitrogen R-300183; siRNA4 Dharmacon L-094802-00) or mock transfected without RNA duplex. Further details can be found in Supplementary Methods.

**Figure 2.** B-Raf induces Sprouty2 phosphorylation on multiple serine residues. A, liquid chromatography-mass spectrometry analysis of Sprouty2 tryptic peptides with precursor 79 scanning on a 4000 Q-Trap mass spectrometer. Extracted ion chromatograms for the phosphopeptides detected from immunoprecipitated Sprouty2 without (left) or with (right) coexpressed B-Raf V600E, identities of each phosphorylation site shown. Ser111 phosphopeptide ion was detected as the M-2H (1256.7) and M-3H (844.5) ions, whereas the other three phosphopeptides were detected as M-2H ions. Schematic representation showing Sprouty2 phosphorylation sites identified by mass spectrometry in bold. Sequence alignment compares Sprouty2 homology surrounding these sites (highlighted by boxes) in selected species. Conserved Sprouty2 cysteine-rich domain and previously identified sites Ser111 and Ser120 are shown. B and C, wild-type or phosphorylation site mutant FLAG-Sprouty2 coexpressed with B-Raf V600E were Western blotted for FLAG and Myc. D, recombinant GST-FLAG-Sprouty2 wild-type or 6A were incubated with WCL-expressing Myc-B-Raf V600E in the presence of [γ-32P]ATP and analyzed for 32P incorporation by autoradiography (top) and Coomassie staining (bottom).
Results

Sprouty2 phosphorylation regulates the association with B-Raf. When WCL from HEK293 cells transfected with FLAG-Tagged Sprouty2 and either wild-type or an activated oncogenic form of myc-tagged B-Raf (V600E) were run on SDS-PAGE beside anti-Myc immunoprecipitations, only the fastest-migrating Sprouty2 electrophoretic variant copurified with wild-type B-Raf (Fig. 1A). Although the distribution of Sprouty2 electrophoretic forms was relatively equal when cotransfected with wild-type B-Raf, there was a shift toward the slower forms in cells expressing B-Raf V600E, with markedly less of the fastest-migrating form that had associated with wild-type B-Raf (Fig. 1A). The shift from fast to slow electrophoretic variants was observed when Sprouty2 was cotransfected with B-Raf V600E or constitutively active MEK1 (MEK:EE; ref. 33), accompanied by increased ERK1 and ERK2 phosphorylation (pERK1/2), suggesting that the mobility shift resulted from MAPK activation (Fig. 1B). It should be noted that typically only two electrophoretic forms are consistently well-separated on SDS-PAGE (e.g., Fig. 1B); separation into three forms is variable likely due to variations in electrophoresis conditions. Consistent with previous results (34), the slower-migrating electrophoretic variants was observed when Sprouty2 was cotransfected with B-Raf V600E or constitutively active MEK1 (MEK:EE; ref. 33), accompanied by increased ERK1 and ERK2 phosphorylation (pERK1/2), suggesting that the mobility shift resulted from MAPK activation (Fig. 1B). It should be noted that typically only two electrophoretic forms are consistently well-separated on SDS-PAGE (e.g., Fig. 1B); separation into three forms is variable likely due to variations in electrophoresis conditions. Consistent with previous results (34), the slower-migrating Sprouty2 electrophoretic form was shifted to the faster form when immunoprecipitated protein was incubated with calf intestinal phosphatase (Fig. 1C, left). Furthermore, in cells metabolically labeled with [32P]orthophosphate, coexpression of B-Raf V600E resulted in increased Sprouty2 phosphorylation (Fig. 1C, right, top). The [32P]-labeled Sprouty2 band comigrated with the slower-migrating Coomassie-stained band, consistent with the mobility shift resulting from phosphorylation (Fig. 1C, right, bottom). These results indicate that B-Raf preferentially associates with faster mobility hypophosphorylated Sprouty2 and suggest that MAPK activation leads to Sprouty2 phosphorylation and reduced B-Raf association. In support of this conclusion, we determined using quantitative direct scanning of Western blots and near-infrared fluorophore-conjugated secondary antibodies that binding of recombinant GST-B-Raf catalytic domain was reduced by 40% if immunoprecipitated Sprouty2 had been coexpressed with active MEK1, which was reversed by phosphatase pretreatment of the immunoprecipitated Sprouty2 (Fig. 1D). The reduced B-Raf binding was associated with a MEK1-induced Sprouty2 mobility shift, which also was reversed by phosphatase (Fig. 1D), consistent with Sprouty2 phosphorylation negatively regulating B-Raf binding. Interestingly, B-Raf association was higher (150%) and the proportion of fast-migrating Sprouty2 was greater following phosphatase treatment than for Sprouty2 immunoprecipitated from cells not expressing MEK1, suggesting that basal Sprouty2 phosphorylation influences B-Raf binding and electrophoretic mobility (Fig. 1D). As a result, there was >2-fold difference in B-Raf binding of Sprouty2 coexpressed with active MEK without or with phosphatase treatment, indicating the contribution of phosphorylation in modifying this interaction. These results indicate that MAPK activation by oncogenic B-Raf results in a higher proportion and/or stoichiometry of Sprouty2 phosphorylation that antagonizes B-Raf binding.

Sprouty2 is phosphorylated on multiple serine residues. To identify B-Raf–induced Sprouty2 phosphorylation sites, we purified Sprouty2 expressed with or without B-Raf V600E and analyzed tryptic fragments by mass spectrometry using a precursor ion scan of m/z -79 on a triple-quadrupole instrument in negative ion mode followed by an ion-trap high-resolution scan, which produces a mass spectrum that contains only the molecular ions of the
phosphopeptides present in the sample (32). Although no basal Sprouty2 phosphorylation was detected (Fig. 2A, left), four sites were identified (Ser7, Ser42, Ser140, and Ser167; numbering = mouse Sprouty2) when coexpressed with B-Raf V600E (Fig. 2A, right). All lie within the Sprouty2 NH2-terminal half and are highly conserved in vertebrate species (Fig. 2A, bottom).

Individual mutations to nonphosphorylatable alanines (S7A, S42A, S140A, and S167A) did not significantly affect the B-Raf V600E-induced Sprouty2 mobility shift (Fig. 2B). It was reported previously that epidermal growth factor–induced phosphorylation of two sites on human Sprouty2 corresponding to Ser111 and Ser120 resulted in slower Sprouty2 electrophoretic mobility (35). We found that mutation of either site to alanine (S111A and S120A) reduced but did not completely reverse the B-Raf V600E-induced mobility shift (Fig. 2B), suggesting that these sites were also modified downstream of B-Raf. Mutation of the four phosphorylation sites identified by mass spectrometry (4A) did not affect Sprouty2 mobility (Fig. 2C) but did result in a loss in Sprouty2 phosphorylation detectable by mass spectrometry (Supplementary Fig. S1B). Mutation of both S111 and S120 to alanines (2A) completely abolished the B-Raf V600E-induced mobility shift (Fig. 2C). Furthermore, 32P incorporation on Escherichia coli–expressed GST-Sprouty2 induced by incubation with WCL from B-Raf V600E-expressing HEK293 cells was markedly diminished in the 6A mutant compared with wild-type Sprouty2 (Fig. 2D). Based on the direct and indirect evidence (identification by mass spectrometry, electrophoretic mobility shift, and 32P incorporation), we have identified six serine residues that appear to be the principal phosphorylation sites modified in response to MAPK activation, with the caveat that there may be additional minor phosphorylation sites not detected by mass spectrometry or that may not influence Sprouty2 electrophoretic mobility.

Sprouty2 phosphorylations cooperate to regulate B-Raf binding. Having identified B-Raf–induced Sprouty2 phosphorylations, we next determined where B-Raf bound to Sprouty2 and whether specific phosphorylations affected B-Raf binding. A peptide array consisting of 23-mers, each consecutively shifted by two residues and spanning the entire Sprouty2 protein, was incubated with [35S]methionine-labeled wild-type B-Raf produced in vitro transcription and translation. The pattern of 35S labeling revealed direct interactions between B-Raf and three distinct
Sprouty2 regions that we named Raf binding domains (RBD) 1-3 (Fig. 3A). Interaction with RBD1 was relatively weak, whereas regions of strong binding were detected within RBD2 and RBD3 (Fig. 3A, red). Four of the six Sprouty2 phosphorylation sites modified in the presence of B-Raf (S42, S111, S120, and S167) lie within these binding domains (Fig. 3A). B-Raf binding to peptides with corresponding phosphoserine residues was reduced for pS111 and pS120 but not for pS42 or pS167 (Fig. 3B). Given that in the initial phosphopeptide array many peptides containing pS111 also contained pS120, we examined whether phosphorylation of either site individually would inhibit B-Raf binding. Individual substitution of either S111 or S120 with phosphoserine inhibited B-Raf binding and no binding was observed when both sites were substituted (Fig. 3C). Similarly, phosphomimetic aspartate or glutamate substitutions at either site inhibited B-Raf binding. Finally, substitution of S111 or S120 with alanine had no effect on B-Raf association, indicating that alanines were equivalent to nonphosphorylated serines (Fig. 3C; summarized in Supplementary Table S1). Collectively, these results indicate that B-Raf directly binds Sprouty2 at three domains and that phosphorylation of Ser111 or Ser120 within RBD2 inhibits this interaction. A similar pattern of phosphorylation-regulated interaction was observed with Raf-1, indicating that it is likely a general phenomenon for Raf kinases (data not shown).

We next investigated how Sprouty2 phosphorylation affected B-Raf binding in cells. Individual mutation of S7, S42, S140, or S167 to alanines had no effect on Sprouty2 binding to wild-type
B-Raf (Fig. 4A, left; Supplementary Fig. S2A). Although individual mutation to S111A or S120A had no significant effect on B-Raf binding, mutation of both increased binding almost 2-fold (Fig. 4A, middle; Supplementary Fig. S2B). Intriguingly, when all six sites were mutated to alanines there was a significant ~3.5-fold increase in B-Raf binding (Fig. 4A, right; Supplementary Fig. S2C). Therefore, although mutation of S7, S42, S140, or S167 individually had little effect on B-Raf association, collective phosphorylation of these sites likely cooperates with S111 and S120 phosphorylation within RBDB to inhibit B-Raf binding.

Oncogenic B-Raf mutants do not bind Sprouty2 independent of phosphorylation status. We found previously that Sprouty2 did not bind to B-Raf V600E (11) and postulated that this could be due to B-Raf–induced Sprouty2 phosphorylation (Fig. 1A). Because the Sprouty2 6A mutant displayed enhanced binding to wild-type B-Raf (Fig. 4A; Supplementary Fig. S2C), we examined whether inhibiting phosphorylation on these sites would restore B-Raf V600E binding. Surprisingly, Sprouty2 association with B-Raf V600E was unaffected by the 6A mutations (Fig. 4B; Supplementary Fig. S3A). Similarly, although recombinant nonphosphorylated FLAG-Sprouty2 associated with recombinant GST-B-Raf wild-type catalytic domain in vitro, it did not associate with the V600E mutant (Fig. 4C). These results indicate that although Sprouty2 phosphorylation inhibits wild-type B-Raf binding, the V600E B-Raf mutant resists Sprouty2 binding independent of phosphorylation status. One possible explanation is that B-Raf mutations that induce active kinase conformations prevent Sprouty2 binding. We therefore tested three B-Raf mutants that lie within the glycine-rich P-loop that either modestly increase (G466A) or impair (G466E and G466V) kinase activity but induce an active kinase conformation in each case (36). Similar to B-Raf V600E, they did not significantly bind Sprouty2 relative to wild-type B-Raf (Fig. 4D; Supplementary Fig. S3B), consistent with our previous observation that the oncogenic activation loop B-Raf mutants L587V, V600D, and K601E did not bind Sprouty2 (11). These results indicate that there are two independent factors influencing the B-Raf/Sprouty2 interaction: Sprouty2 phosphorylation and B-Raf kinase domain conformation.

Sprouty2 phosphorylation influences MAPK activation. Given that Sprouty2 phosphorylation affects wild-type B-Raf binding, we tested whether the 6A mutant would have an enhanced ability to repress MAPK activation by FGF. We generated NIH3T3 cells expressing tetracycline-regulated (Tet-Off) wild-type or 6A mutant Sprouty2. Three wild-type and three 6A Sprouty2 cell lines were selected for their relatively comparable Sprouty2 expression following doxycycline withdrawal (Supplementary Fig. S4). A control nonexpressing clone was also established using the empty tetracycline-responsive vector. FGF treatment of the control cells resulted in increased phosphorylated ERK1 and ERK2 that were sustained over 90 min, which was unaffected by doxycycline withdrawal (Fig. 5A, top). In contrast, ERK activation was somewhat inhibited when wild-type Sprouty2 was induced following doxycycline removal, consistent with Sprouty2 being an inhibitor of FGF-induced MAPK activation (ref. 37; Fig. 5A, middle). Inhibition of ERK activation was more pronounced in 6A Sprouty2-expressing cells relative to wild-type Sprouty2 (Fig. 5A, bottom). When the areas under the curve for active ERK levels over time were calculated as a measure of signal output (38), 6A Sprouty induction significantly inhibited MAPK signal output relative to wild-type Sprouty2 (Fig. 5B). Although ERK activation in 6A Sprouty2-expressing cells was lower at every time point, the differences between wild-type and 6A Sprouty2 did not achieve statistical significance until 90 min (Fig. 5C). One interpretation is that wild-type Sprouty2 inhibits FGF-induced MAPK activation, with Sprouty2 phosphorylation at later time points reducing this inhibitory effect. In contrast, the 6A Sprouty2 mutant is not phosphorylated on critical sites and MAPK inhibition persists, decreasing MAPK signal output. Therefore, Sprouty2 phosphorylation may act as an additional regulatory mechanism to control MAPK signaling. Consistent with Sprouty2 inhibition of B-Raf being the cause of reduced MAPK activation, B-Raf knockdown with two independent siRNA duplexes (Fig. 5D, left) resulted in proportionate and significant reduction in MAPK activation (Fig. 5D, right).

Endogenous Sprouty2 phosphorylation downstream of endogenous V600E B-Raf. We examined how endogenous oncogenic V600E B-Raf affected the mobility of endogenous Sprouty2 in mouse embryonic fibroblasts (30). Mouse embryonic fibroblasts from mice that expressed a conditionally activated Cre-estrogen receptor (Cre:ER) alone or in combination with a heterozygous knock-in V600E B-Raf mutation preceded by a Lox-Stop-Lox (LSL) cassette were either left untreated or treated with 50 nmol/L 4-hydroxytamoxifen (4HT) for 96 h to induce Cre-mediated recombination. Sprouty2, pERK1/2, and ERK2 levels were determined by quantitative Western blotting.

Discussion

Sprouty regulation of receptor tyrosine kinase signaling is complex, because it acts at multiple levels and may have apparently
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positive and negative actions within the same pathway. That being said, one way Sprouty proteins regulate MAPK signaling is through direct association with Raf kinases (11, 25–28). We observed that B-Raf binding is negatively regulated by Sprouty2 phosphorylation on six serine residues that act in concert; two sites (Ser111 and Ser120) directly affect binding, whereas the remaining four appear to contribute indirectly, possibly by affecting protein conformation. Furthermore, Sprouty2 was unable to associate with B-Raf mutants that adopt active kinase conformations. Therefore, we propose that there are two independent factors that affect the Sprouty/B-Raf interaction: Sprouty2 phosphorylation and B-Raf kinase conformation. These findings have important implications for MAPK signaling in normal and cancer cells.

A significant observation from this study is that activated B-Raf mutants do not bind Sprouty2 regardless of its phosphorylation status. These findings are consistent with, and provide some mechanistic detail for, a recent publication reporting that B-Raf V600E evades a feedback inhibition response that includes elevated Sprouty2 (21).

Sprouty gene transcription is regulated by the MAPK pathway (40, 41), consistent with their role as negative feedback regulators, and oncogenic mutations in melanoma and gastrointestinal stromal tumors that activate MAPK also lead to elevated Sprouty levels (11, 20–23). At the same time, Sprouty2 phosphorylation is increased in response to MAPK activation, thereby reducing the ability of Sprouty2 to bind B-Raf and to inhibit FGF activation of MAPK (Figs. 4 and 5). Therefore, it appears that the MAPK pathway both transcriptionally up-regulates Sprouty2 and posttranscriptionally attenuates its ability to inhibit MAPK signaling. One possible reason for this arrangement is that Sprouty may be regulated via transcription, but once protein levels have been elevated, a further level of acute control may be exerted by varying the phosphorylation-dephosphorylation status. This possibility suggests that Sprouty proteins may not act as simple on/off switches but might be tunable regulators that influence MAPK signaling kinetics, duration, magnitude, and signal output. If phosphorylation inhibits the ability of Sprouty2 to repress B-Raf, then it would be predicted that dephosphorylation should increase Sprouty2-mediated MAPK inhibition. Consistent with this possibility, we found that the 6A mutant was significantly better at suppressing FGF-induced MAPK activation at later time points (Fig. 5), whereas it was recently reported that Xenopus laevis Sprouty2 does not influence acute ERK activation by FGF but does regulate the duration of ERK activity (42).

In conclusion, we have established that the ability of Sprouty2 to act as an inhibitor of B-Raf activity and the MAPK pathway is under dual control by two independent mechanisms: posttranslational phosphorylation of Sprouty2 and B-Raf kinase conformation. The phosphorylation status of specific residues on Sprouty2 influences its ability to interact with B-Raf and to modulate MAPK activation in response to growth factors, whereas evidence suggests that the active B-Raf kinase conformation independently prevents association with Sprouty2. Delineation of the precise signaling pathway leading to Sprouty phosphorylation is an important future goal in light of the important implications these findings have for MAPK signaling in normal and cancer cells.

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

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