**Molecular and Cellular Pathobiology**

**Alternative Splicing of Caspase 9 Is Modulated by the Phosphoinositide 3-Kinase/Akt Pathway via Phosphorylation of SRp30a**

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**Abstract**

Increasing evidence points to the functional importance of alternative splice variations in cancer pathophysiology. Two splice variants are derived from the CASP9 gene via the inclusion (Casp9a) or exclusion (Casp9b) of a four-exon cassette. Here we show that alternative splicing of Casp9 is dysregulated in non−small cell lung cancers (NSCLC) regardless of their pathologic classification. Based on these findings we hypothesized that survival pathways activated by oncogenic mutation regulated this mechanism. In contrast to K-RasV12 expression, epidermal growth factor receptor (EGFR) overexpression or mutation dramatically lowered the Casp9a/9b splice isoform ratio. Moreover, Casp9b downregulation blocked the ability of EGFR mutations to induce anchorage-independent growth. Furthermore, Casp9b expression blocked inhibition of clonogenic colony formation by erlotinib. Interrogation of oncogenic signaling pathways showed that inhibition of phosphoinositide 3-kinase or Akt dramatically increased the Casp9a/9b ratio in NSCLC cells. Finally, Akt was found to mediate exclusion of the exon 3,4,5,6 cassette of Casp9 via the phosphorylation state of the RNA splicing factor SRp30a via serines 199, 201, 227, and 234. Taken together, our findings show that oncogenic factors activating the phosphoinositide 3-kinase/Akt pathway can regulate alternative splicing of Casp9 via a coordinated mechanism involving the phosphorylation of SRp30a. Cancer Res; 70(22); 9185–96. ©2010 AACR.

**Introduction**

Non−small cell lung cancer (NSCLC) accounts for >80% of all lung cancers diagnosed. Thus, there is an urgent need for further understanding of the molecular mechanisms of NSCLC, thereby defining new molecular targets for the treatment of this disease (1). In NSCLC, inappropriate activation of the phosphoinositide 3-kinase (PI3K) pathway by direct oncogenic mutation or from a K-Ras or epidermal growth factor receptor (EGFR) mutation has been associated with the development of NSCLC (2–4). A critical component of the PI3K nexus is the proto-oncogene Akt, a serine/threonine kinase. Akt is activated by recruitment to membranes through direct contact of its PH-domain with PtdIns (3,4,5) P₃ and subsequent phosphorylation on two key residues, threonine 308 and serine 473. Previously, several laboratories reported that Akt is constitutively active in NSCLC, and experimental models have shown that Akt activation contributes to the resistance of NSCLC to chemotherapy.

An important player in the contrasting signaling cascade of apoptosis, with roles in both the sensitivity of NSCLC to chemotherapy and radiation, is Casp9. Casp9 is a member of the caspase family of proteases and is activated upon the formation of trimeric complex with cytochrome c and APAF-1, termed the apoptosome. Casp9 has roles in both the response of cells to undergo chemotherapy-induced cell death and the suppression of oncogenic transformation (5–15). The CASP9 gene produces two antagonistic isoforms, the proapoptotic Casp9a and the prosurvival Casp9b, via the inclusion/exclusion of an exon 3, 4, 5, 6 cassette (7, 12). The Casp9b isoform (exon exclusion) lacks catalytic activity while retaining key interacting domains (e.g., CARD; refs. 7, 12). Casp9b acts as an endogenous inhibitor of Casp9a by competing with the full-length Casp9a for binding to the apoptosome (7, 12). Casp9b has also been surmised to directly interact with Casp9a blocking the autoproteolysis of the enzyme (7). In this study, Casp9 splicing was shown to be dysregulated in NSCLC tumors and cell lines, and regulated by the PI3K/Akt pathway. Furthermore, this study shows that Akt exerts its effects via the phospho-status of the RNA trans-factor SRp30a.

**Materials and Methods**

**Cell culture**

A549, H2347, H358, H226, H2170, H596, H460, H1792, H1299, H520, H1703, H2030, H838, HCC827, and H292 cells were
obtained from the American Type Culture Collection and were grown in 50% RPMI 1640 and 50% DMEM supplemented with L-glutamine, 100 units/mL penicillin G sodium, and 100 µg/mL streptomycin sulfate. NHBE cells, from Clonetics (Cambrex Bio Science), were maintained in basal bronchial epithelial growth media (BEGM) supplemented with SingleQuot Kits-growth factors, cytokines, and supplements (Cambrex). HBEC-3KTs, HBEC-3KT K-RasV12, wild-type EGFR, and mutant EGFR HBEC-3KT cell lines were previously created and characterized (16). HBEC-3KT (E746-A750 del EGFR, K-RasV12) cell lines stably expressing control or Casp9b short hairpin RNA (shRNA) were created using lentivirus followed by selection with 20 µg/mL hygromycin (Lentigen). To generate A549, H838, and HCC827 cell lines stably expressing Casp9b cDNA, 1.5 × 10⁶ cells were transfected with pcDNA3.1(-)/Casp9b using Effectene followed by selection with zeocin. For all competition studies between cell lines, cells were plated in keratinocyte serum-free medium without supplements overnight prior to analysis.

Quantitative reverse transcriptase-PCR

**Competitive.** Total RNA (1 µg) was reverse-transcribed using Superscript III and oligo (dT) and analyzed for Casp9, Bcl-x, caspase 8, and caspase 2 splice variants as previously described (17).

**Real-time.** Total RNA was used for real-time PCR for Casp9a, Casp9b, and 18s using TaqMan PCR master mix and the Applied Biosystems 7500 Real-Time PCR System. Casp9a and 18s Q-PCR primers were ID numbers Hs00154261_m1 and Hs99999901_s1. The Q-PCR primers for Casp9b were 5′-TTGGTGATGTCGGAGCA-3′ (forward) and 5′-GATT-TGGTGATGTCGGAGCA-3′ (reverse) and 5′-TTCCCTGAGA-GACGACTCCCTGG-3′ (probe).

Inhibitor studies/adenviral infection

Cells (1.2 × 10⁶) were infected two hours with constitutively active Akt2, PTEN, or Null adenovirus (50MOI). After 48 hours, total RNA and protein were isolated.

Clonogenic and soft agar assays

For clonogenic assays, 150 viable cells were seeded into 6-well tissue culture dishes with complete growth media. Two hours postplating, cells were treated with erlotinib. After 24 hours, cells were rinsed and placed in complete media for 12 days. Colonies were counted following fixation and 0.1% crystal violet. For soft agar assays, cells (10³) were prepared as previously described (16).

Phosphorylation assays

Twenty-four hours after transfection, cells were scraped into 150 µL of NP-40 lysis buffer [50 mmol/L Tris-HCl (pH 8.0), 150 mmol/L NaCl, 1% NP-40,100 µg of phenylmethylsulfonyl fluoride/mL, and protease inhibitor cocktail]. The lysates were incubated with 500 U of alkaline phosphatase for 1 hour at 37°C. Wild-type and mutant SRp30a proteins were detected using anti-T7 tag antibody.

Western immunoblotting

Immunoblotting was accomplished as previously described (18) using primary antibodies, anti-Casp9 (Assay Designs), anti-Akt1 (Santa Cruz), anti-Akt2 (Santa Cruz), anti-Akt phospho-S473 (Cell Signaling), anti-Akt pan (Cell Signaling), anti-ERK1/2 (Santa Cruz), anti-ERK1/2 phospho-Thr202/204 (Cell Signaling), anti-PDK1 (Santa Cruz), anti-T7 tag (Novagen), and anti-f-actin (Sigma-Aldrich). Secondary antibodies were horseradish peroxidase–conjugated goat anti-mouse or anti-rabbit (Sigma-Aldrich).

Transfections

The mammalian expression vectors pCGT7-SRp30a-WT and SRp30a-RD have been previously described (19). To generate SRp30a phospho-mutants, site-directed mutagenesis was performed using QuikChange XL Site-Directed Mutagenesis protocol. To analyze Casp9 splicing, 1.5 × 10⁶ cells were transfected with 0.5 µg of pCGT7-SRp30a ± 0.2 µg of the Casp9 minigene using Effectene as previously described (17). After 24 hours, RNA and protein were isolated. For downregulation of Akt1, Akt2, and PDK1, cell lines were transfected with either Akt1, Akt2, or PDK1 SMARTpool multiplex siRNA or control siRNA (Dharmacon) using DharmaFECT 1 as previously described (18). After 48 hours, RNA and protein were isolated.

Statistical analysis

When appropriate, the data are presented as mean ± SE. Data points were compared using a two-tailed, unpaired Student’s t test, and the P values were calculated. P values < 0.05 were considered significant.

Results and Discussion

Casp9 RNA splicing is dysregulated in NSCLC tumors and cell lines

In this study, we examined the hypothesis that Casp9 RNA splicing was dysregulated in all pathologies of NSCLC. Utilizing total RNA from pathologist-verified human NSCLC samples, quantitative/competitive reverse transcriptase-PCR (RT-PCR) analysis was performed to determine the degree of dysregulation in the Casp9a/9b ratio as compared with matched, normal lung tissue controls (Supplementary Table S1; ref. 17). Tumor samples were categorized into three separate groups: normal, a Casp9a/9b mRNA ratio of >3; moderately dysregulated, a Casp9a/9b mRNA ratio of 2.3–3.3; and highly dysregulated, a Casp9a/9b mRNA ratio <2.3 (Fig. 1A and B). The normal group corresponds to the normal ratio of Casp9a/9b mRNA observed in nontransformed cells; the moderately dysregulated group corresponds to a ratio of Casp9a/9b reported to have a significant, but minor effect on Casp9 activity (15, 20); and the highly dysregulated group corresponds to a ratio of Casp9a/9b reported to significantly reduce Casp9 activity and inhibit the association of Casp9a with APAF-1 (15, 17, 20). Analysis (18, 21–23) of Casp9 splice variants showed that 36% of NSCLCs examined presented a moderately dysregulated Casp9a/9b mRNA ratio (n = 149). Importantly, 42% of tumors showed a >50% decrease in the Casp9a/9b ratio. Thus, the ratio of Casp9a/9b mRNA is significantly lower in a high percentage of NSCLC tumors irrespective of NSCLC subtype.
We then examined a pure population of nontransformed lung epithelial cells, specifically primary human bronchial epithelial cells (NHBE) and immortalized HBEC-3KT cells, for the ratio of Casp9a/9b in comparison with the transformed lung epithelial cell lines A549, H838, H2347, H358, H2030, H226, H2170, H596, H1792, H1299, H520, H1703, and H292 (Supplementary Table S2). HBEC-3KT cells presented with a normal Casp9a/9b ratio of 4.02 ± 0.15, as did NHBE cells (4.15 ± 0.23; Fig. 1C). In contrast, 8 of 11 transformed lung epithelial cell lines grown under the exact same culture conditions showed a dysregulated Casp9a/9b ratio.

Figure 1. The Casp9a/9b mRNA ratio is dysregulated in NSCLC tumors and transformed lung epithelial cells. cDNAs from pathologist-verified lung adenocarcinomas, and squamous and large cell carcinomas (Origene) underwent quantitative/competitive PCR for expression of Casp9 splice variants. A, representation of matched pair analysis used for the degree of Casp9a/9b dysregulation in NSCLC tumors. N, normal; T, tumor. B, RT-PCR analysis of Casp9 splice variants showing 36% of NSCLC tumors are moderately dysregulated (C9a/9b ratio of 2.2:3.3) and 42% of NSCLC tumors are highly dysregulated (C9a/9b ratio < 2.2; n = 149). C, RT-PCR analysis of Casp9 splice variants from NSCLC and HBEC-3KT cells. *, dysregulated Casp9a/9b mRNA ratio; **, moderately dysregulated Casp9a/9b mRNA ratio. D, in parallel, protein expression of Casp9 splice variants.
conditions showed a significant decrease in the Casp9a/9b mRNA ratio. Importantly, the disproportionate ratio of Casp9a/9b mRNA observed in the transformed lung epithelial cell lines translated to a disproportionate ratio of Casp9a/9b protein expression (Fig. 1D). We further validated the decrease in the Casp9a/9b mRNA ratio of A549s in comparison with HBEC-3KTs via quantitative PCR (Q-PCR; Supplementary Fig. S1), reconfirming the quantitative nature of the assay as also previously shown by ribonuclease protection assay (21). These data indicate that a significant portion of NSCLC tumors and transformed lung epithelial cells show severe dysregulation of the alternative splicing of Casp9 to favor a prosurvival/pro-oncogenic phenotype.

The EGF pathway regulates Casp9 RNA splicing in a pro-oncogenic fashion

In essentially all epithelial cancers, including NSCLC, one or more members of the family of EGFR genes are either overexpressed or mutated (4). As a large percentage of NSCLC tumors and cell lines showed a dysregulated ratio of Casp9a/9b, we next examined whether this common oncogene in NSCLC affected Casp9 RNA splicing in HBEC-3KT cells. Whereas low expression of K-RasV12 (Supplementary Fig. S2A, B) in HBEC-3KT cell lines (16) had no discernable effect on the ratio of Casp9a/9b mRNA, the overexpression of wild-type EGFR, the expression of L858R mutation in EGFR, and the expression of the del E746-A750 EGFR mutant induced a significant reduction in the Casp9a/9b ratio (Fig. 2A). These HBEC-3KT cell lines expressing either wild-type or mutant EGFR have previously been reported to exhibit enhanced phosphorylation of EGFR in the absence of EGF, in contrast to HBEC-3KT cells expressing K-RasV12 (16). We also reconfirmed these previous findings showing increased phosphorylation of AKT in HBEC-3KT cells expressing wild-type or mutant EGFR, as well as showing an increase in the phosphorylation of extracellular signal-regulated kinase 1/2 (ERK1/2) in HBEC-3KT cells expressing K-RasV12 (Supplementary Fig. S2A, B). Importantly, the del E746-A750 mutation of EGFR induced the most significant decrease in the Casp9a/9b ratio correlating with the ability of this EGFR mutant to induce anchorage-independent growth (AIG; ref. 16). Therefore, we hypothesized that increased expression of Casp9b was a major mechanism in conferring AIG of HBEC-3KT ΔE746-A750 del cells. To test this hypothesis, an E746-A750 del EGFR clonal cell line (HBEC-3KT Δ746/C9b-dr) stably expressing Casp9b shRNA was produced. The expression of Casp9b in the HBEC-3KT Δ746/C9b-dr clonal cell line was reduced to normal immunoreactive levels, and presented with a normal Casp9a/Casp9b ratio (Fig. 2B). Importantly, the HBEC-3KT Δ746/C9b-dr cells showed a complete loss of AIG compared with E746-A750 del EGFR cells stably expressing control shRNA (Fig. 2C). These effects did not require stable expression and were not due to integration artifacts as short-term/transient downregulation of Casp9b also inhibited the AIG of E746-A750 del EGFR cells (Fig. 2D and E). Furthermore, K-RasV12-expressing HBEC-3KT cells with Casp9b downregulated exhibited no difference in their AIG capacity, showing specificity for oncogenic EGFR (Supplementary Fig. S3).

Thus, the distal mechanism of Casp9 splicing plays a major and specific role in the ability of EGFR signaling to confer AIG. The significance of this finding stems from the knowledge that EGFR mutation occurs in a large percentage of NSCLCs, roughly 5% to 10%, and overexpressed EGFR and its ligands occur in approximately 70% of NSCLCs. Interestingly, EGFR mutation/overexpression is also considered an early event in NSCLC because the mutation is also found in normal epithelial cells prior to metaplasia and adenoma formation, rationalizing the hypothesis that the alternative splicing of Casp9 plays a role in early events leading to the formation of NSCLC.

Modulation of Casp9 RNA splicing regulates the ability of erlotinib to inhibit the colony formation of NSCLC cells

Based on the above data, we were prompted to investigate the effect of erlotinib, a clinically relevant inhibitor of the human EGFR, on the Casp9a/9b ratio. Treatment of A549, H838, H460, and HCC827 cells with erlotinib led to a dose-dependent increase in the Casp9a/9b ratio (Fig. 3A). We further validated the increase in the Casp9a/9b ratio via Q-PCR (Supplementary Fig. S4A and B). Additionally, the effectiveness of erlotinib treatment was shown by loss of Akt phosphorylation (Supplementary Fig. S4C). Next, the effect of lowering the Casp9a/9b ratio on the sensitivity of A549 cells to erlotinib was examined. Low ectopic expression of Casp9b dramatically inhibited the ability of erlotinib to suppress cell survival (e.g., induced a resistant phenotype; Fig. 3B). In contrast, downregulation of Casp9b sensitized A549 cells to erlotinib as shown by a dramatic reduction in the IC50 (14.9 μmol/L for control shRNA to 4.8 μmol/L for C9b shRNA; Fig. 3B). To determine translatable, the effect of lowering the Casp9a/9b ratio on the sensitivity of H838 and HCC827 cells to erlotinib was also examined. H838 cells ectopically expressing Casp9b cDNA significantly inhibited the ability of erlotinib to suppress cell survival increasing the IC50 (8.5 μmol/L for vector control to 22.2 μmol/L for C9b cDNA; Fig. 3C). Conversely, downregulation of Casp9b sensitized H838s to erlotinib, reducing the IC50 to 2.9 μmol/L for C9b shRNA cells (Fig. 3C). Additionally, these same effects translated to HCC827 cells, a cell line that has been reported to be highly sensitive to erlotinib-induced growth inhibition (24). Specifically, HCC827 cells ectopically expressing Casp9b cDNA showed an increase in the IC50 of erlotinib (22.2 nmol/L for vector control to 89.9 nmol/L for C9b cDNA; Fig. 3D). Conversely, downregulation of Casp9b reduced the IC50 to 11.7 nmol/L for C9b shRNA cells (Fig. 3D). Thus, Casp9 RNA splicing plays a major role in the sensitivity of NSCLC cells to erlotinib. As erlotinib has shown promise in clinical trials, significantly improving the survival rate of NSCLC patients (25, 26), examining the alternative splicing of Casp9 may have future predictive/prognostic value for a subset of patients or allow for determination of erlotinib responsiveness in NSCLC tumors.

The PI3K/PDK1/Akt pathway regulates Casp9 RNA splicing

We next hypothesized that a major mitogenic signaling pathway activated by EGFR regulates Casp9 RNA splicing
to favor the production of Casp9b. To investigate this hypothesis, mitogenic pathways were examined for effects on the ratio of Casp9a/9b utilizing small-molecule inhibitors at doses/times reported in the scientific literature for A549 cells (27–33). Only treatment with the PI3K inhibitor LY 294002 (50 μmol/L) resulted in a significant increase in the ratio of Casp9a/9b, compared with the inactive, structurally related compound LY303511 (Fig. 4A, Supplementary Table S3). Specifically, the Casp9a/9b ratio increased from 2.10 ± 0.18 for control samples to 3.91 ± 0.08 for inhibitor-treated samples (P < 0.0008; n = 6). The inhibition of PI3K effectively returned the Casp9a/9b ratio to the ratio observed in nontransformed lung epithelial cells. Importantly, this effect on the Casp9a/9b ratio translated to the protein level (Fig. 4A).

To determine translatability, H838, H2030, H358, H292, and H520 cells were also treated with LY294002 (Fig. 4B). Again, the Casp9a/9b ratio was dramatically increased (H838 cells from 1.24 ± 0.05 to 3.29 ± 0.19; H2030 cells from 2.30 ± 0.08 to 3.74 ± 0.08; H358 cells from 3.11 ± 0.09 to 4.13 ± 0.12; H292 cells from 2.01 ± 0.09 to 4.11 ± 0.11; and H520 cells from 2.35 ± 0.08 to 3.91 ± 0.13). Therefore, the ability of the PI3K pathway to regulate the alternative splicing of Casp9 translates to multiple NSCLC cell lines.

Akt/PKB, SGK, PKCξ, and PKCδ are downstream of PI3K (34). To investigate the downstream effector of PI3K responsible for regulating Casp9a RNA splicing, small-molecule inhibitors in conjunction with RNA interference technology was employed. Treatment of A549 cells with the PKC inhibitors GÖ6976 (10 μmol/L) and GÖ6983 (10 μmol/L; Supplementary Fig. S5A and B, Supplementary Table S3) resulted in no significant change in the Casp9a/b ratio. Conversely, treatment of A549 cells with the Akt inhibitor Akt VIII (25 μmol/L) and the
Figure 3. Casp9 RNA splicing regulates the ability of erlotinib to inhibit colony formation. A, RT-PCR analysis of Casp9 splice variants from A549, H838, H460, and HCC827 cells treated with erlotinib. Clonogenic plate assays of A549 cells (B), H838 cells (C), and HCC827 cells (D) expressing ectopic Casp9b cDNA, vector control (vec con), Casp9b shRNA, or control (con) shRNA after 24 hours in the presence of erlotinib. Data are mean ± SE represented as percent control.
phosphatidylinositol analog SH-5 (10 μM) resulted in an increased Casp9a/9b ratio to the same extent as PI3K inhibition, from 2.10 ± 0.11 for DMSO to 4.03 ± 0.13 for Akt VIII (n = 6; P < 0.01) and from 2.10 ± 0.08 for DMSO to 3.73 ± 0.13 for SH-5 (n = 4; P < 0.01; Fig. 4C and D). In addition, treatment of cells with the PDK1 inhibitor OSU03012 (10 μM) or siRNA downregulation of PDK1 (Fig. 4E and F) resulted in a significantly increased Casp9a/9b ratio, from 2.25 ± 0.13 for control to 4.05 ± 0.22 for OSU03012 (n = 4; P < 0.01) and from 2.13 ± 0.09 for siRNA controls to 3.78 ± 0.21 for siPDK1 (n = 4; P < 0.01). To verify appropriate inhibition of downstream signaling pathways, the phosphorylation status of Akt was examined via Western immunoblotting (Supplementary Fig. S6A–C).

These data show that Akt regulates Casp9 RNA splicing in an antiapoptotic/prosurvival manner. This effect was specific for Casp9 RNA splicing, and not a generalized effect on constitutive RNA splicing, as no effect on the alternative splicing of caspase 8, caspase 2, and Bcl-x pre-mRNA was observed with Akt VIII treatment (Fig. 4G). Furthermore, cotreatment of A549 cells with both Akt VIII and LY294002 could not further increase the Casp9a/9b ratio, showing a linear pathway.
with Akt as the downstream effector of PI3K (Supplementary Fig. S7).

To examine the Akt isoform responsible for regulating the alternative splicing, siRNA was again utilized. Downregulation of Akt1 using multiplex siRNA resulted in no significant change in the Casp9a/b ratio ($P > 0.34$; Fig. 5A). In contrast, downregulation of Akt2 resulted in a dramatic increase in the Casp9a/b ratio from $2.10 \pm 0.05$ to $4.09 \pm 0.25$ ($P < 0.01$; $n = 5$; Fig. 5B). Again, this effect translated to the protein level (Fig. 5C).

To further validate a role for Akt2, expression of constitutively active Akt2 (always membrane bound via myristoylation) induced a significant decrease in the Casp9a/9b ratio (Fig. 5D). Furthermore, ectopic expression of PTEN induced the contrasting effect (Fig. 5E). Therefore, Akt2 regulates Casp9 RNA splicing via PIP$_3$ membrane interactions in A549 cells, and also requires PDK1 phosphorylation.

To determine translatability, H2030, H292, H358, H520, and HCC827 cells were treated with Akt VIII. Indeed, all cell lines also showed a dramatic increase in the Casp9a/9b ratio (Fig. 5F). Additionally, Akt1 and Akt2 were downregulated in H358 cells to determine the Akt isoform responsible for regulating Casp9 RNA splicing. In contrast to A549s, downregulation of Akt1, but not Akt2, induced a significant increase in the Casp9a/9b ratio (Fig. 5G). Therefore, multiple Akt isoforms have the potential to regulate Casp9 RNA splicing in NSCLC cells.

Lastly, as with the NSCLC cell lines, the ability of del 746-750 EGFR mutation to induce a lowering of the Casp9a/9b ratio in HBEC-3KT cells was dependent on PI3K and Akt.

Figure 5. Akt regulates Casp9 RNA splicing via PIP$_3$ membrane interactions. A and B, RT-PCR analysis of Casp9 splice variants from A549 cells transfected with control siRNA or Akt1 siRNA (A), and control siRNA or Akt2 siRNA (B). C, Western blot analysis for Casp9. D and E, RT-PCR analysis of Casp9 splice variants from A549s infected with null or constitutively active Akt2 (D), or null and PTEN adenovirus (E). F, H2030, H358, H292, H520, and HCC827 cells were treated with 0.1% DMSO or Akt inhibitor VIII (25 μmol/L) for 24 hours. Total RNA was isolated and RT-PCR performed for Casp9 splice variants. G, RT-PCR analysis of Casp9 splice variants from H358s transfected with control siRNA or Akt1 siRNA. Data are expressed as means ± SE.
as treatment with LY294002 and Akt VIII increased the ratio of Casp9a/Casp9b (Fig. 6A and B). These data suggest that EGFR overexpression/mutation correlates with Akt activation affecting the alternative splicing of Casp9. We further extended these results by examining the Akt isoform responsible for regulating Casp9 RNA splicing in HBEC-3KT (del E746-A750 EGFR) cells. Interestingly, siRNA against both Akt1 and Akt2 (Fig. 6C) resulted in a significantly increased Casp9a/Casp9b ratio as compared with control siRNA, from 2.65 ± 0.05 for siRNA control to 4.01 ± 0.13 for siAkt1 (α = 3; P < 0.01) and to 3.72 ± 0.11 for siAkt2 (α = 3; P < 0.01). Therefore, multiple Akt isoforms are responsible for regulating the alternative splicing of Casp9. The finding that the Akt pathway regulates this splicing mechanism important for EGFR conferring AIG makes logical sense in relation to cellular transformation, as this pathway is found constitutively active in approximately 58% of NSCLC cell lines and tumors (35, 36). This pathway is also linked to constitutive EGFR activity, NF-κB activation, and the ability of oncogenic Ras to transform various cell types (37–39). Thus, the Akt pathway activated by EGFR is extremely important in a therapeutic sense, and this study suggests that the alternative splicing of Casp9 is a key distal mechanism in the biological role of this pathway in NSCLC development/maintenance.

How Casp9b acts to drive AIG is more of an enigma. As we have shown, removal of Casp9b blocked the ability of EGFR (del E746-A750) to induce AIG. This is likely not attributed to the blockade of an initiator caspase such as Casp9a, and suggests a role in cell signaling for Casp9b. In this regard, the possibility that Casp9b acts as a signaling molecule has been reported by Latchman and coworkers (5). Specifically, this laboratory group showed that ectopic expression of Casp9b induced the activation of NF-κB irrespective of caspase activation (5). Activation of NF-κB by Casp9b expression also “fits” well with cooperation with K-Ras mutations for the induction of cellular transformation based on the findings of Ma and Baldwin (40, 41). These laboratory groups showed in several different ways that NF-κB activation enhances the ability of oncogenic ras to induce cellular transformation. Coupled with the knowledge that EGFR overexpression/mutation leads to both cooperation with oncogenic ras in cellular transformation and NF-κB activation, a role for Casp9b in these pathways important for cellular transformation is logical. Thus, Casp9b may act as a scaffolding protein to elicit downstream signaling events with roles outside the simple inactivation of Casp9a. This possibility is far from inconceivable as the initiator caspase, caspase 8, is reported to recruit cell survival factors such as PI3K subunits (42–44). Although this caspase is an initiator of extrinsic pathways of apoptosis, Casp9b may play an analogous role in survival signaling as an initiator of the intrinsic pathway of apoptosis.

The phospho-status of SRp30a modulates the effect of Akt signaling on Casp9 RNA splicing

Our laboratory reported that SRp30a was a required enhancer factor for the inclusion of the exon 3,4,5,6 cassette of Casp9 (18). Furthermore, Srp30a has been shown to be a specific target of Akt in vitro (45, 46). Therefore, we hypothesized that the phospho-status of SRp30a regulates the inclusion of the exon 3,4,5,6 cassette of Casp9, downstream of Akt activation. To investigate this hypothesis, we employed an established phospho-mimic of SRp30a, Srp30a-RD, in which the majority of serine residues in the RS domain were mutated to aspartic acid (19). Coexpression of Srp30a-RD with a functional Casp9 minigene induced a significant decrease in the Casp9a/9b ratio compared with wild-type Srp30a and empty vector controls (Fig. 7A). Importantly, expression of Srp30a-RD also induced a significant decrease in the endogenous Casp9a/9b ratio as compared with wild-type Srp30a and empty vector controls (Fig. 7B).

To determine the serine residue/residues of Srp30a required for regulating the alternative splicing of Casp9, site-directed replacement mutagenesis was utilized. Multiple serine residues in the RS domain of Srp30a contain recognized motifs for Akt phosphorylation and numerous residues have been validated by mass spectrometric analysis (Supplementary Table S4; ref. 45). These serine residues were individually mutated into aspartic acid to produce phospho-mimics (Supplementary Table S4). Coexpression of only the Srp30a-S199D, Srp30a-S201D, Srp30a-S227D, and Srp30a-S234D mutants with a functional Casp9 minigene decreased the
Figure 7. The phospho-status of SRp30a regulates Casp9 RNA splicing. A, B, C, and D, RT-PCR analysis of Casp9 minigene-derived transcripts (A and C) or endogenous Casp9 splice variants (B and D) from A549 cells transfected with the Casp9 minigene and SRp30a-WT or SRp30a-RD (A); SRp30a-WT or SRp30a-RD (B); and the indicated SRp30a phospho-mutants (C and D). E, F, G, and H, Western blot analysis of the phosphorylation state of SRp30a. Lysates were analyzed for the expression/migration of T7-tagged SRp30a. E, A549 cells were transfected with the indicated T7-tagged SRp30a constructs and total cell lysates were incubated in the presence of either active or denatured AP. F, HBEC-3KT and A549 cells transfected with the indicated T7-tagged SRp30a constructs and total cell lysates were incubated in the presence of either active or denatured AP. G, endogenous SRp30a from A549 cells treated with 0.1% DMSO or erlotinib (50 μmol/L) for 48 hours. Total protein lysates were incubated in the presence of either active or denatured AP. H, endogenous SRp30a from A549 cells treated with 0.1% DMSO or Akt VIII (50 μmol/L). I, RT-PCR analysis of endogenous Casp9 splice variants from A549s transfected with SRp30a-WT or SRp30a-QD ± Akt inhibitor Akt VIII (25 μmol/L). Data are expressed as means ± SE.
Casp9a/9b ratio compared with wild-type SRp30a control (Fig. 7C).

We extended these results to produce a SRp30a double phospho-mutant (SRp30a-S199,201,234D), a SRp30a triple phospho-mutant (SRp30a-S199,201,234D), and a SRp30a quadruple (SRp30a-QD) mutant harboring serine to aspartic acid mutations at residues 199, 201, 227, and 234. Expression of SRp30a double and triple phospho-mutants with the Casp9 minigene further decreased the Casp9a/9b ratio (Supplementary Fig. S8). Ectopic expression of SRp30a-QD induced a reduction in the Casp9a/9b ratio comparable with the SRp30a-RD mutant. Conversely, a quadruple dephospho-mimic of SRp30a (SRp30a-QA) induced the opposite effect in A549 cells (Fig. 7D). These effects could not be due to localization issues as both SRp30a quadruple mutants are localized in the nucleus and were expressed in equivalent amounts (Supplementary Fig. S9).

To show that these phospho-sites are hyperphosphorylated in NSCLC, the phosphorylation status of the transiently expressed SRp30a was analyzed by comparing the electrophoretic migration profiles in the presence of either alkaline phosphatase (AP) or denatured AP. A dramatic increase in the migration of SRp30a-WT was observed after treatment with AP, indicating that this protein is phosphorylated in A549 cells. In contrast, the migration of SRp30a-QD and SRp30a-QA after treatment with AP was significantly decreased in comparison with the migration of SRp30a-WT, indicating that these proteins are phosphorylated to a significantly lesser extent (Fig. 7E). These data show that serines 199, 201, 227, and 234 exist in a phosphorylated state in A549 cells. Moreover, the electrophoretic migration profiles of A549s and HBEC3-KT cells after transfection with SRp30a-WT and SRp30a-QA indicate that SRp30a exists in a decreased phosphorylated state in nontransformed cells versus NSCLC cells (Fig. 7F).

We extended these results to determine the effect of inhibiting the EGFR/Akt pathway on the phosphorylation status of SRp30a in A549 cells. As predicted, an increase in the migration of endogenous SRp30a was observed after treatment with erlotinib (Fig. 7G), as well as after treatment with the Akt inhibitor Akt VIII (Fig. 7H). To determine whether the PI3K/Akt pathway regulates Casp9 RNA splicing in a phospho-SRp30a-dependent manner, SRp30a-QD was expressed in the presence or absence of Akt VIII. In the presence of SRp30a-QD, Akt VIII was unable to increase the ratio of the Casp9a/9b to the same extent as compared with wild-type SRp30a (Fig. 7I). Therefore, the Akt pathway regulates the alternative splicing of Casp9 at least partially via the phospho-state of SRp30a on serines 199, 201, 227, and 234.

These data solidify a role for phosphorylation of SRp30a in regulating the alternative splicing of Casp9, but also suggest additional regulating mechanisms. In this regard, our laboratory recently found that the RNA trans-factor hnRNP L acts as a repressor for the inclusion of the exon 3,4,5,6 cassette of Casp9, and its repressor activity is regulated by the phosphorylation status of Ser52 (17). Thus, we hypothesize that the EGFR/PI3K/Akt pathway may also regulate the phosphorylation status of hnRNPL at Ser52, suggesting a coordinated interplay between these two trans-factors in regulating the alternative splicing of Casp9. This possibility is logical, as Lynch and coworkers showed the ability of SRp30a and hnRNPL to directly compete for binding to the exon 5 regulatory sequence of CD45 and that this interplay between SRp30a and hnRNPL influences the extent of exon inclusion (47).

The phospho-state of SRp30a regulating the inclusion of the exon 3,4,5,6 cassette also fits with our previous findings that ceramide induced both the dephosphorylation of SRp30a and the inclusion of the Casp9 exon cassette. SRp30a was also required for ceramide effects on the inclusion of the exonic cassette of Casp9. Thus, the regulation of SRp30a phosphorylation and the alternative splicing of Casp9 may be a key distal point by which ceramide acts as a tumor suppressing/cell senescence agent as the ceramide signaling and PI3K/Akt pathway are well established to antagonize each other (Supplementary Fig. S10).

In conclusion, the presented study reports several major findings taking a comprehensive approach. First, the deregulation of the alternative splicing of Casp9 toward a prosurvival phenotype was shown in NSCLC. Second, a survival/mitogenic/oncogenic pathway involving EGFR, PI3K, and Akt was shown to regulate this splicing mechanism. Lastly, the phospho-state of SRp30a was shown to regulate this distal mechanism via Akt signaling. Therefore, the present study shows a novel and key distal mechanism in NSCLC and provides new target mechanisms for the development of therapeutics to combat the cancer with the highest mortality rate.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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