Abstract

Cables1 is a candidate tumor suppressor that negatively regulates cell growth by inhibiting cyclin-dependent kinases. Cables1 expression is lost frequently in human cancer but little is known about its regulation. Here, we report that Cables1 levels are controlled by a phosphorylation and 14-3-3–dependent mechanism. Mutagenic analyses identified two residues, T44 and T150, that are specifically critical for 14-3-3 binding and that serve as substrates for phosphorylation by the cell survival kinase Akt, which by binding directly to Cables1 recruits 14-3-3 to the complex. In cells, Cables1 overexpression induced apoptosis and inhibited cell growth in part by stabilizing p21 and decreasing Cdk2 kinase activity. Ectopic expression of activated Akt (AKT1) prevented Cables1-induced apoptosis. Clinically, levels of phosphorylated Cables1 and phosphorylated Akt correlated with each other in human lung cancer specimens, consistent with pathophysiologic significance. Together, our results illuminated a dynamic regulatory system through which activated Akt and 14-3-3 work directly together to neutralize a potent tumor suppressor function of Cables1. Cancer Res; 75(1); 1–12. © 2014 AACR.

Introduction

Cables1 (Cdk5 and Abl enzyme substrate 1) is a novel Cdk2, Cdk3, and Cdk5-binding protein, which acts as a link between the Cdk5 and nonreceptor tyrosine kinases and regulates the activity of Cdks by enhancing their Y15 phosphorylation (1, 2). In neurons, Cables1 promotes C-Abl to phosphorylate Cdk5 at Y15, resulting in increased kinase activity, and is believed to positively regulate neurite outgrowth. However, in proliferating cells, Cables1 connects Cdk2 and Wee1, which results in increased phosphorylation of Cdk2 at Y15, decreased kinase activity, and reduced cell proliferation. Cables1 interacts with p53 and p73, resulting in the induction of cell death (3), and also binds to TAp63α to protect it from proteasomal degradation to ensure deletion of cells after genotoxic stress (4). Compared with Cables1+/− MEFs, Cables1−/− MEFs exhibit an increased growth rate, delayed senescence, and decreased serum dependence (5). Furthermore, Cables1−/− mice have an increased incidence of endometrial cancer and a reduced survival rate in response to unopposed estrogen and colorectal cancer caused by 1,2-dimethylhydrazine (6, 7). Loss of Cables1 expression is observed with high frequency in human colon, lung, ovarian, and endometrial cancers (6), (8–10), and also enhances tumor progression in the ApcMin/− mouse model and activates the Wnt/β-catenin signaling pathway (11). Together, these observations suggest that Cables1 may function as a tumor suppressor. However, little is known about the regulation of Cables1 itself. It remains to be established how the growth suppressive function of Cables1 is coupled to cell survival and proliferative mechanisms. Our work revealed a signaling network interface by which Cables1 is complexed with a phospho-Set/Thr-recognition protein, 14-3-3, and its upstream kinase.

The 14-3-3 proteins are a highly conserved family of regulatory proteins expressed in all eukaryotic cells (12–16). In mammals, there are seven 14-3-3 isoforms (β, η, ε, σ, ζ, γ, τ) encoded by distinct genes. 14-3-3 proteins function as dimers to bind to functionally diverse target proteins, including kinases, phosphatases, receptors, and molecular adaptors. 14-3-3 proteins regulate target proteins by cytoplasmic sequestration, occupation of interaction domains, prevention of degradation, activation/repression of enzymatic activity, and facilitation of protein modifications (12, 13), (15–18). Binding of 14-3-3s with target proteins is tightly regulated and the major mode of regulation is through reversible phosphorylation of target proteins within a defined motif. Two canonical 14-3-3–binding motifs have been identified as RSXp/TXP (model I) and RXFXp/TXP (model II), and a third C-terminal motif, pS/TX1-2-COOH (model III), has been defined (14, 19, 20). Within these motifs, phosphorylation of a specific serine (S) or threonine (T) residue is necessary for binding with 14-3-3. However, many target proteins do not contain sequences that accord precisely with these motifs, and some target...
proteins bind to 14-3-3 in a phosphorylation-independent manner. Interestingly, the consensus phosphorylation motif of the serine/threonine kinase Akt, RXRXpS/T, partially overlaps with the sequences of mode I and II 14-3-3–binding motifs. Indeed, Akt phosphorylates many substrates within phosphorylation motifs, which recruits 14-3-3 binding. Therefore, 14-3-3 binds to a number of Akt substrates and regulates various cell biological functions, including cell survival, proliferation, and metabolism. For example, Akt directly phosphorylates the Bcl-2 family member Bad on residue S136 and this creates a binding site for 14-3-3 proteins, which triggers release of Bad from its target proteins and inhibits the proapoptotic function of Bad (21–23). The FOXO transcription factors are also phosphorylated by Akt, which then recruits 14-3-3 binding and promotes their cytoplasmic retention. In this way, Akt prevents FOXO-induced target gene transcription that promotes apoptosis, cell-cycle arrest, and metastatic processes (24, 25). Thus, the identification and characterization of new protein targets that act downstream of Akt with coupled 14-3-3 binding may have significant biologic and therapeutic implications.

Here, we present data to suggest a novel signaling mechanism by which Cables1 is suppressed by the combined actions of the Ser/Thr kinase, Akt, and the adaptor protein 14-3-3. Akt phosphorylation-mediated 14-3-3 binding prevents the apoptosis-inducing function of Cables1. Together, our data offer a new mechanism through which Cables1/Akt/14-3-3 interactions couple survival signaling to cell death.

**Materials and Methods**

**Cells and reagents**

COS7 and HEK293T cells were purchased from ATCC and maintained in DMEM with 10% FBS and 100 U penicillin–streptomycin in a humidified atmosphere of 5% CO₂. Low-serum DMEM (1% FBS) was used in all experiments. Plasmids were transfected into cell lines using the QuikChange kit, following the manufacturer’s protocol (Stratagene). Transfections were performed using FuGene HD (Roche).

**Plasmids and transfection**

Cables1 cDNAs were amplified by PCR and cloned into Gateway expression vectors (Invitrogen). Site-directed mutagenesis was performed using the QuikChange kit, following the manufacturer’s protocol (Stratagene). Transfections were performed using FuGene HD (Roche).

**Protein interaction assays**

**HexaHistidine (His)-affinity pulldown assay.** Cells were lysed in Hi-pulldown lysis buffer (1% Nonidet P-40, 137 mmol/L NaCl, 1 mmol/L MgCl₂, 40 mmol/L Tris-Cl, 60 mmol/L imidazole, 5 mmol/L Na₂HPO₄, 5 mmol/L NaF, 2 mmol/L Na₃VO₄, 1 mmol/L phenylmethylsulfonyl fluoride, 10 mg/mL aprotinin, 10 mg/mL leupeptin). Lysates were cleared by centrifugation at 4°C. The clarified cell lysate was incubated with nickel-charged hexaHis resin for 2 hours at 4°C. The resin was washed two times with washing buffer (500 mmol/L NaCl, 20 mmol/L Tris-Cl, 60 mmol/L imidazole) and once with binding buffer (500 mmol/L NaCl, 20 mmol/L Tris-Cl, 5 mmol/L imidazole). Bound proteins were released from the resin by boiling in 6× SDS sample buffer for Western blot analysis.

**GST pull-down assay.** Cells were lysed in GST pulldown lysis buffer (1% Nonidet P-40, 150 mmol/L NaCl, 100 mmol/L Hepes, 5 mmol/L Na₂HPO₄, 5 mmol/L NaF, 2 mmol/L Na₃VO₄, 1 mmol/L phenylmethylsulfonyl fluoride, 10 mg/mL aprotinin, 10 mg/mL leupeptin). Cleared cell lysates were incubated with glutathione-conjugated sepharose (GE Healthcare) for 2 hours at 4°C. Then the resin was washed three times with GST pulldown lysis buffer and boiled in 6× SDS sample buffer for Western blot analysis.

**Coimmunoprecipitation assay.** Cells were lysed in coimmunoprecipitation lysis buffer (1% Nonidet P-40, 500 mmol/L NaCl, 100 mmol/L Hepes, 5 mmol/L Na₂HPO₄, 5 mmol/L NaF, 2 mmol/L Na₃VO₄, 1 mmol/L phenylmethylsulfonyl fluoride, 10 mg/mL aprotinin, 10 mg/mL leupeptin). Cleared cell lysates were incubated with Protein A or G conjugated sepharose (GE Healthcare) and the appropriate antibody for 2 hours to overnight at 4°C. Following incubation, the resin was washed three times with coimmunoprecipitation lysis buffer and protein samples were eluted by boiling in 6× SDS sample buffer for Western blot analysis.

**Akt1 and Cdk2 kinase assays**

Recombinant active Akt1 (100 ng) was incubated with 10 µCi of [γ-³²P]ATP and 10 µg of recombinant Cables1 in 30 µL of kinase buffer (25 mmol/L Hepes, 25 mmol/L β-glycerophosphate, 25 mmol/L MgCl₂, 2 mmol/L dithiothreitol, 0.1 mmol/L NaVO₃). To examine Cdk2 activity, Cdk2 isolated from lysates was incubated with 10 µCi of [γ-³²P]ATP and 5 µg of histone H1 in 30 µL of kinase buffer (50 mmol/L Hepes, 5 mmol/L MgCl₂, 10 mmol/L dithiothreitol). All reactions were incubated at 30°C for 30 minutes and terminated by addition of 6× sample buffer. Proteins were separated by 10% SDS-PAGE, and phosphorylation was visualized by autoradiography.

**Time resolved–Förster resonance energy transfer assays**

Our published protocols for the time resolved–Förster resonance energy transfer (TR-FRET) assay were followed (26, 27). FITC-conjugated Cables1 T44 (FITC-Ahx-ENAPLRRCRTLSGSPR), T150 (FITC-Ahx-ENAPGARRNTIDSTSS), pT44 (FITC-Ahx-ENAPLRRC(pT) LSQSPR), and pT150 (FITC-Ahx-ENAPGARRN(pT) IDSTSS) peptides were synthesized by Peptide 2.0 Inc (>80% purity). Bad pS136 was generated as described previously (28). Purified 6×His tagged 14-3-3 proteins were indirectly labeled with terbium (Tb) fluorophore as a TR-FRET donor through a Tb-conjugated anti-6×His antibody (Cisbio Biosays). The TR-FRET assay was performed in 384-well plates (30 µL/well). All assay components were diluted in assay buffer containing 20 mmol/L Tris buffer, pH 7.5, 50 mmol/L NaCl, and 0.01% Nonidet P-40. Briefly, increasing amounts of 14-3-3 proteins were mixed with Flu-labeled pT44, T44, pT150, T150 peptide, or pBad and incubated with anti-His-Tb antibody (50 ng/mL). After incubation at room temperature for 2 hours, the TR-FRET signal was detected using an Envision Multilabel plate reader (PerkinElmer Life Sciences) with laser excitation at 337 nm, emissions at 486 nm and 520 nm, with a dual dichroic mirror (400/505 nm). The delay time was set at 50 µs. The TR-
FRET signal is expressed as the TR-FRET signal ratio: F520 nm/F486 nm×107, where F520 nm and F486 nm are fluorescence counts at 520 nm and 486 nm for fluorescein and Tb, respectively. The TR-FRET signal window was calculated as the difference between the TR-FRET signal values for bound Flu-peptide in the presence of 14-3-3 protein and values for unbound Flu-peptide in the absence of 14-3-3 protein. All experimental data were analyzed using Prism 5.0 software (GraphPad Software).

14-3-3y affinity chromatography for identification of 14-3-3 binding partners
14-3-3–binding protein identification from A549 lung cancer cells, including the discovery of Cables1 as a novel 14-3-3 partner, is described in the Supplementary Materials section.

Western blot analysis
Proteins were separated on 12.5% SDS-PAGE gels and transferred to polyvinylidene difluoride membranes. Membranes were blocked with 5% BSA and incubated with the indicated primary antibodies. Corresponding horseradish peroxidase-conjugated secondary antibodies (Santa Cruz Biotechnology) were used against each primary antibody. Proteins were detected using West-Pico or West-Dura enhanced chemiluminescent detection reagents (Pierce) and a Kodak imaging system or films.

Immunohistochemistry assay
Formalin-fixed, paraffin-embedded human lung cancer tissue array slides (ABXIS and Biochain) were stained with anti-pCables1 T44, T150 (21st Century), and pAkt S473 (Epitomics) antibodies using a microwave-enhanced avidin-biotin staining method. For quantitation of protein expression, the following formula was used: immunohistochemistry (IHC) score = % positive cells × intensity score. The intensity was scored as follows: 0, negative; 1, weak; 2, moderate; and 3, intense. An IHC score of 100 or greater was considered positive.

Statistical analysis
A Student t test was used to compare individual data points among each group. Correlation was analyzed using Fisher exact test. A P value of less than 0.05 was set as the criterion for statistical significance.

Results
Cables1 interacts with 14-3-3
To discover critical signaling nodes at the junction of cell survival and death, we utilized 14-3-3 protein as a molecular probe in an affinity capture-based proteomics study to explore novel 14-3-3–binding proteins and their regulation (Supplementary Materials, ref. 29). Our proteomics analysis in A549 lung cancer cells identified known 14-3-3–binding partners such as keratins, various 14-3-3 isoforms, and MEKI, validating the employed approach (Supplementary Table S1). This approach also revealed a number of potential novel 14-3-3–binding proteins, including Cables1. Cables1 attracted our attention due to its demonstrated role in the regulation of cell-cycle progression, although, the precise mechanisms by which Cables1 is regulated remain unclear. To validate whether Cables1 indeed interacts with 14-3-3y, we cotransfected GST or GST-Cables1 and His-14-3-3y into COS7 cells and performed a His pulldown assay. As shown in Fig. 1A, only GST–Cables1, but not GST, was detectable in His–14-3-3y complexes as analyzed by Western blot analysis. We also conducted the reverse GST pulldown assay and found the presence of His-14-3-3y in GST–Cables1 complexes, but not in GST complexes (Fig. 1B). There are seven 14-3-3 mammalian isoforms (β, η, ε, σ, γ, τ), which often share many binding partners, but also demonstrate isoform-specific binding to some proteins. To determine whether Cables1 has any isoform selectivity in its interaction with 14-3-3, GST–Cables1 along with the seven different His-14-3-3 isoforms were overexpressed in COS7 cells and the cell lysates were subjected to His pulldown and Western blot analysis. As shown in Fig. 1C, Cables1 preferentially bound to η, σ, γ, and τ, but not to the β and ε isoforms of 14-3-3. As expected, the negative control 14-3-3y/K50E, which reduces the association of 14-3-3 with most ligands due to a mutated ligand-recognition site in 14-3-3, did not bind to Cables1, supporting 14-3-3–binding specificity (30). To further confirm the interaction of Cables1 with 14-3-3 under physiologic conditions, we investigated their endogenous interaction. Endogenous Cables1 and 14-3-3y were separately immunoprecipitated from PC12 cells in both directions with Cables1 and 14-3-3y antibodies, respectively. Reciprocal protein detection was performed. As shown in Fig. 1D, both Cables1 and 14-3-3y were detected in the individual immunoprecipitated complexes, 14-3-3y and Cables1, respectively, but not in the control IgG complexes. From these data, we conclude that Cables1 is a natural 14-3-3–binding protein in the native cellular environment.

Most interactions of 14-3-3 with target proteins are mediated by phosphorylation (12, 14, 16, 17, 19). To determine whether the interaction of Cables1 with 14-3-3 is phosphorylation dependent, we performed in vitro phosphatase assays. COS7 lysates expressing both GST–Cables1 and His-14-3-3y were incubated with or without calf intestinal phosphatase (CIP) to induce dephosphorylation or general phosphatase inhibitors to maintain phosphorylation, then GST–Cables1 was probed in the His–14-3-3y complexes (31). Cables1 binding to 14-3-3 was reduced by CIP treatment and the addition of phosphatase inhibitors effectively reversed the CIP effect (Fig. 1E). To test whether the interaction between Cables1 and 14-3-3 could be regulated by endogenous phosphatases, we performed the same phosphatase experiment as described above, but without the addition of the exogenous phosphatase CIP. Lysates were incubated at room temperature, with the intention that these conditions would activate endogenous phosphatases within the cell lysate. Indeed, Cables1 binding with 14-3-3 was reduced by endogenous phosphatases, and this reduction was blocked by the presence of phosphatase inhibitors (Fig. 1E). These results indicate that Cables1 interaction with 14-3-3 is
Cables1 binds 14-3-3 through T44 and T150 sites

The binding of 14-3-3 to target proteins is generally mediated through RSXP/S/TPX and RXXXP/S/TPX motifs where P/S/T represents phosphoserine or phosphothreonine (17). To explore which binding motifs in Cables1 mediate its binding with 14-3-3, we first generated two truncations of Cables1, 1-200 and 201-368, and tested their interaction with 14-3-3. GST-Cables1 truncation 1-200 or 201-368 and His-14-3-3 were cotransfected into COS7 cells and His pulldown and Western blot analysis were performed. As shown in Fig. 2A, truncation 1-200 was able to bind to His-14-3-3γ, whereas truncation 201-368 did not bind to His-14-3-3γ. This result indicates the binding sites on Cables1 that are required for interaction with 14-3-3 are located within residues 1-200 of Cables1. Next, we searched for conserved sequences in Cables1 using ScanSite (www.scansite.com) and identified several potential 14-3-3-binding sites, including T44, S46, S48, T150, and S169. To determine which of the predicted S/T residues are true 14-3-3-binding sites, we mutated all S/T residues to alanine (A) and examined the binding of these mutants with 14-3-3 in His-14-3-3 complexes. F, calyculin A enhances the binding of 14-3-3 with Cables1. Cells overexpressing His-14-3-3γ and GST-Cables1 were treated with the indicated amount of calyculin A for 1 hour, then GST-Cables1 was examined in His-14-3-3γ complexes.

Indeed dynamically regulated by phosphorylation, and the phosphatase responsible for reversing this interaction is endogenously expressed in COS7 cells. Next, we carried out in vivo phosphatase assays using a specific phosphatase inhibitor, calyculin A, to treat COS7 cells expressing both GST-Cables1 and His-14-3-3γ, and then detected the levels of GST-Cables1 in His pulldown complexes. As shown in Fig. 1F, GST-Cables1 binding to His-14-3-3γ was dose dependently enhanced with gradually increasing calyculin A concentrations. These data support the importance of regulated phosphorylation dictating the interaction of Cables1 with 14-3-3.
Akt/14-3-3 Regulates Cables1

Cables1 AA and DD with His-14-3-3γ was clearly weaker than that of Cables1 WT (Fig. 2C). These data suggest that the T44 and T150 sites likely mediate the binding of Cables1 with 14-3-3. As the DD mutant did not interact with 14-3-3, we assume that the DD mutant did not mimic the phosphorylated state of Cables1 required for 14-3-3 binding.

If the T44- and T150-containing regions of Cables1 directly bind 14-3-3, these isolated peptides may be able to compete for the interaction of full-length Cables with 14-3-3. To test this, we performed a competitive binding assay by preincubating the peptides derived from Cables1 with lysates overexpressing GST-Cables1 and His-14-3-3γ for 1 hour. GST-Cables1 was detected in His-14-3-3γ isoform complexes. Direct binding of the pT44 peptide of Cables1 to 14-3-3γ (left) and 14-3-3γ (right), F, direct binding of the pT150 peptide of Cables1 to 14-3-3γ (left) and 14-3-3γ (right). TR-FRET titration assays were carried out in triplicate in a 384-well plate with 5 nMol/L Cables1 peptides and increasing 14-3-3 concentrations for 2 hour. The TR-FRET assay window was calculated as described in Materials and Methods. Both unphosphorylated T44 and T150 peptides were included for comparison.
Taken together, these results indicate that Cables1 may require both pT44 and pT150 sites for effective binding with 14-3-3, possibly in a coordinated fashion (16). Moreover, both T44 and T150 sites are highly conserved among a variety of species, further supporting the potential importance of these two sites through evolution (data not shown).

**Akt phosphorylates Cables1 at 14-3-3–binding sites**

The two 14-3-3–binding sites on Cables1, T44 and T150, reside in sequences that overlap with consensus motifs for potential Akt phosphorylation. To test the hypothesis that Akt phosphorylates Cables1 and then recruits 14-3-3, we carried out the experiment of WT and kinase dead (KD) Akt1 on the binding of Cables1 to 14-3-3. HA-Akt1 WT or KD was cotransfected with GST-Cables1 and His-14-3-3 into COS7 cells, then His pulldown assay and Western blot analysis were carried out. Akt1 WT significantly enhanced the binding of Cables1 and His-14-3-3, whereas Akt1 KD moderately decreased their binding (Fig. 3A). Next, we used a general anti-pAkt substrate antibody that recognizes the motif RXXpS/T to detect phosphorylated levels of Cables1 WT and various single mutants in GST-Cables1 pulled down complexes. As shown in Fig. 3B, both Cables1 T44A and T150A single mutants showed significantly lower levels of pAkt substrate recognition, whereas other Cables1 single mutants showed levels equal to Cables1 WT. To specifically detect the phosphorylated level of Cables1 T44 and T150, we generated corresponding anti-pCables1 T44 and T150 antibodies. The levels of pCables1 T44 and pCables1 T150 were equal for all Cables1 variants except the T44A and T150A mutants, respectively, which showed significantly reduced levels (Fig. 3B). We also used the same methods to examine the phosphorylated levels of the Cables1 AA and DD mutants when coexpressed with Akt1 WT or KD. Phosphorylated levels of GST-Cables1 WT were clearly increased when Akt1 WT was overexpressed and were decreased when Akt1 KD was overexpressed, but phosphorylated levels of the Cables1 AA and DD mutants were significantly reduced and even undetectable under

**Figure 3.**
Akt phosphorylation of Cables1 recruits 14-3-3 binding. A, Akt enhances the binding of Cables1 with 14-3-3. Hist14-3-3γ complexes were pulled down from cell lysates overexpressing His-14-3-3γ, GST-Cables1, and HA-Akt1 WT or KD, followed by SDS-PAGE, and Western blot analysis with the indicated antibodies. B, mutations in T44 and T150 of Cables1 abolish its phosphorylation by Akt. Each GST-Cables1 variant was isolated from transfected cells and detected by the indicated antibodies. C, exogenous Akt phosphorylates Cables1. Phosphorylation of GST-Cables1 WT, AA, and DD was examined in cells overexpressing HA-Akt1 WT or KD. D, Akt interacts with Cables1. COS7 cells were cotransfected with HA-Akt1 WT or KD and GST or GST-Cables1 WT. After 48 hours, cells were lysed, GST pulldown was performed, and proteins were detected by Western blot analysis. E, endogenous Akt phosphorylates Cables1. Cells were transfected with GST-Cables1, and after 24 hours, cells were serum starved for 24 hours. Cells were treated with or without 10 μmol/L Akt1/2 inhibitor or 10 μmol/L LY299002 for 1 hour followed by 100 ng/mL IGF for 15 minutes. Cells were lysed and phosphorylation of GST-Cables1 was measured in the isolated GST-Cables1 complex. F, Akt phosphorylates Cables1 in vitro. The indicated recombinant proteins were incubated with or without recombinant Akt1 in kinase buffer containing γ-[32P]ATP at 30°C for 0.5 hour. Proteins were separated by SDS-PAGE, followed by autoradiography, Coomassie stain, or Western blot analysis.
certain conditions (Fig. 3C). Next, we assessed the interaction between Akt1 and Cable1 by detecting HA-Akt1 WT and KD levels in GST, GST–Cable1 WT, or GST–Cable1 AA complexes, which were pulled down from their overexpressing lysates. HA-Akt1 WT and KD were detectable in GST–Cable1 WT or GST–Cable1 AA complexes but not in GST complexes, and HA-Akt1 WT and KD showed equal interactions with GST–Cable1 WT and the AA mutant (Fig. 3D). To test whether endogenous Akt can also phosphorylate Cable1, we activated endogenous Akt by treating serum-starved GST–Cable1 overexpressing cells with IGF1 and detecting phosphorylated levels of pulled down GST–Cable1. As shown in Fig. 3E, activating endogenous Akt with IGF1 markedly enhanced the phosphorylated levels of GST–Cable1. This enhancement was totally blocked by pretreating cells with the PI3K inhibitor, LY294002, or Akt1/2 inhibitor. To further examine whether Akt is able to phosphorylate Cable1 directly, we performed an in vitro radio labeling kinase assay using recombinant Akt1 and GST–Cable1 WT, T44A, T150A, and AA mutants. The autoradiography results demonstrated that Cable1 WT was effectively phosphorylated by Akt, showing significant labeling with $^{32}$P. Although mutations in Cable1, T44A, and T150A decreased the labeling of $^{32}$P signals of GST–Cable1, the GST–Cable1 AA double mutant exhibited the greatest reduction in Cable1 phosphorylation (Fig. 3F). In addition, Western blot analysis detected pCables1 T44 only with GST–Cable1 WT and T150 mutants, and pCables1 T150 only with GST–Cable1 WT and T44 mutants (Fig. 3F). Together, these data suggest that Akt is an upstream kinase that phosphorylates Cable1 at T44 and T150 sites.

Cable1 overexpression induces apoptosis

Cable1 has been reported to enhance p53-induced cell death in U2OS cells (3). Overexpressing Cable1 alone could also induce apoptosis in several ovarian cancer cells (32). To determine the role of the 14-3-3–binding sites in Cable1–induced apoptosis, we overexpressed control Venus, Venus–Cable1 WT, and AA in HEK293T cell. Apoptosis of Venus-positive cells was analyzed by detecting Annexin V-positive cells as well as cleaved PARP levels by Western blot analysis. As shown in Fig. 4A and B, overexpressing Venus–Cables1 WT induced apoptosis and PARP

Figure 4.

Cable1 overexpression induces cell apoptosis. A, Cable1 overexpression dose dependently induces apoptosis. HEK293T cells in 12-well plates were transfected with increasing amounts of Venus, Venus–Cables1 WT, or AA. After 72 hours, cells were lysed and proteins were detected by Western blot analysis. Cells were stained with Annexin V-PE and induction of apoptosis in Venus-positive cells was analyzed by flow cytometry. B, Cable1 overexpression time dependently induces apoptosis. HEK293T cells in 12-well plates were transfected with 1 μg Venus, Venus–Cables1 WT, or AA. After the indicated times, protein detection and apoptosis analysis were performed as in A. C, Cable1 overexpression induces increased intracellular cleaved PARP. Venus, Venus–Cables1 WT, and AA–overexpressing cells were stained with rabbit anti-C-PARP antibody and goat anti-rabbit IgG with conjugated Texas Red and Hoechst 33342. Images were taken with an Image Xpress 5000. D, Cable1 overexpression inhibits Cdk2 activity. Cdk2 was immunoprecipitated from the lysates of Venus, Venus–Cables1 WT, or AA–overexpressing cells, then used in a kinase assay. Proteins were detected by Western blot analysis. * and ** represent $P < 0.05$ and $P < 0.01$, respectively, for values versus those in the control Venus group. E, Cable1 overexpression increases p21 and decreases pRb levels. Lysates of Venus, Venus–Cables1 WT, and AA–overexpressing cells were used to examine the indicated proteins levels by Western blot analysis.

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HEK293T cells were enhanced withincreasing serum-starvation cleavage in Venus, Venus-Cables1 WT, and AA-overexpressing cells. As shown in Fig. 5B, endogenous pAkt in Venus, Venus-Cables1 WT, and AA-overexpressing HEK293T cells, and AA-overexpressing HEK293T cells, and found that Cables1 AA induced higher p21 and lower phosphorylated Rb protein levels than Cables1 WT. No changes were observed in the protein levels of Bax, p53, p27, p57, cyclin A, cyclin D1, or cyclin E (Fig. 4E).

Activated Akt prevents apoptosis induced by Cables1

The above data suggest that the Akt phosphorylation sites of Cables1 may modulate its inhibition of CdK2, the stability of p21, and the apoptosis induction activity of Cables1. Next, we determined the effects of Akt on apoptosis induced by Cables1. We coexpressed Venus, Venus-Cables1 WT, and AA as well as HA-Akt1 WT and KD in HEK293T cells and analyzed induction of apoptosis as above. As shown in Fig. 5A, overexpressing HA-Akt1 WT significantly inhibited apoptosis and PARP cleavage induced by Venus-Cables1 WT, but moderately inhibited apoptosis and PARP cleavage induced by Venus-Cables1 AA. In contrast, overexpressing HA-Akt1 KD increased apoptosis and PARP cleavage in Venus-Cables1 WT, and AA-overexpressing cells. We also inactivated endogenous Akt by withdrawing serum from the culture medium of Venus, Venus-Cables1 WT, and AA-overexpressing HEK293T cells, and analyzed cell apoptosis. As shown in Fig. 5B, endogenous pAkt S473 levels, which indicate endogenous Akt activity, decreased with increasing serum-starvation time. Apoptosis and PARP cleavage in Venus, Venus-Cables1 WT, and AA-overexpressing HEK293T cells were enhanced with increasing serum-starvation time. These results suggest that activated Akt is able to prevent apoptosis induced by Cables1.

The level of pCables1 is correlated with that of pAkt in human lung cancer patient and A549 xenograft mouse model tissues

The above results demonstrate that Cables1 is phosphorylated by Akt in cell culture. To determine whether this is also the case in tumor tissues, we compared the levels of pCables1 T44, T150, and pAkt S473 in 37 human lung cancer samples by immunostaining with the corresponding antibodies. Information about sex, age, histology, and IHC results of the samples is summarized in Supplementary Table S2, and the IHC images of three representative samples are shown in Fig. 6A. Although sample 1 showed negative staining of pCables1 T44, T150 and pAkt S473, sample 2 showed positive pAkt S473 staining with negative staining of pCables1 T44 and T150, and sample 3 showed positive staining of pCables1 T44, T150, and pAkt S473. The results from the IHC analysis are summarized in Fig. 6B. Positive pAkt S473 staining was present in 13 out of 37 patient tumor tissue samples. Importantly, positive pCables1 T44 and T150 staining was only present in nine out of 37 samples. Importantly, all nine samples also showed positive pAkt S473 staining, suggesting that the levels of pCables1 T44 and T150 in human lung cancer tissues might be controlled by the same mechanism as the activated Akt level. Together, these results in human lung cancer specimens confirm our observations in cell-culture experiments, and indicate that the level of pCables1 is correlated with that of pAkt, supporting a potentially significant role in lung cancer tumorigenesis.
These studies led to our working model (Fig. 7) and suggest that Cables1 growth inhibition activity is antagonized by oncogenic kinases, such as Akt, through phosphorylation of Cables1 at T44 and T150. To test this model, we examined whether Akt status was correlated with Cables1 phosphorylation at these two sites in vivo using a lung cancer A549 xenograft mouse model (33, 34). As shown in Supplementary Fig. S1, tumors treated with vehicle showed relatively high Akt phosphorylation at T473 along with phosphorylated Cables1 at T44 and T150. Conversely, tumors treated with an mTOR kinase inhibitor, INK128, exhibited reduced Akt pT473, and showed decreased phosphorylation of Cables1 at T44 and T150. When tumors were treated with INK128 and a GSK3β inhibitor, SB216763, both the Akt phosphorylation level and the Cables1 phosphorylation level were reversed. Band intensity information was captured by normalizing pAkt and Cables1 at pT44 and pT150 against pan-Akt and Cables1. The statistical analysis (MatLab, corrcoef) of these data led to $P = 0.009$ for pAKT/pT44 of Cables1 with a correlation coefficient ($R$) of 0.717 and $P = 0.001$ for pAKT/pT150 of Cables1 ($R = 0.832$), suggesting highly significant correlation between phosphorylation level of Akt and Cables1 at these sites further supporting the proposed working model in Fig. 7.

**Discussion**

In the present study, we identified a critical mechanism that regulates Cables1 function by which the cell growth inhibition activity, and thus the tumor suppression activity, of Cables1 is suppressed by activated Akt and Akt phosphorylation-induced 14-3-3 binding. We have identified Cables1 as a new 14-3-3 interacting protein and demonstrated that their interaction is phosphorylation dependent and mediated by the T44 and T150 sites of Cables1. Although motif scanning shows that T44 (not T150) is a classical 14-3-3–binding motif, our mutational results suggest that both of these sites mediate 14-3-3 binding, although the binding of synthesized peptides with 14-3-3 in vitro indicates that the Cables1 pT44 peptide binds 14-3-3 more potently than the Cables1 pT150 peptide. Structural analysis of 14-3-3 dimers has revealed that each monomer contains an independent target-protein binding region; therefore, the dimer can interact with two motifs simultaneously, belonging to either a single protein or separate binding partners. Such binding through two sites allows intricate signal transmission and network coordination (16). The binding of the T44 and T150 sites of Cables1 with 14-3-3 most likely occurs in such a coordinated fashion.

We have identified Akt as one kinase that can directly bind to and phosphorylate Cables1, and recruit 14-3-3 binding. Akt, also known as protein kinase B, is a central node in cell signaling downstream of growth factors, cytokines, and other cellular stimuli. Activated Akt phosphorylates many protein substrates and thus has diverse roles in several cellular processes, including cell survival, growth, proliferation, angiogenesis, metabolism, and migration (35). In addition to Cables1,
Akt phosphorylates several Cables1-related proteins and induces their interaction with 14-3-3. Akt is able to phosphorylate Wee1 and promote its cytoplasmic localization by binding to 14-3-3. Relocalized Wee1 cannot phosphorylate Cdk1 and Cdk2 at Y15 sites, which relieves their kinase activity and promotes cell-cycle progression (36). Akt also phosphorylates Cdk2 and causes its cytoplasmic localization through interaction with 14-3-3. This Cdk2 cytoplasmic redistribution is required for cell progression from S to G2–M phase (37). Several groups have reported that Akt also phosphorylates the Cdk inhibitor p27, resulting in its cytosolic sequestration via 14-3-3 binding. Inhibiting p27 nuclear localization enhances its degradation and attenuates its cell-cycle inhibitory effects (38–40). Similarly, Akt phosphorylates another Cdk inhibitor, p21, which, like p27, leads to p21 cytosolic localization by interaction with 14-3-3 (41). Recently, one component of the SCF^Skp2 ubiquitin ligase complex Skp2, which mediates ubiquitination and degradation of several cell-cycle–related proteins, including p21 and p27, was shown to be phosphorylated by Akt. Skp2 phosphorylation by Akt enhances its stability through disrupting the interaction between Cdh1 and Skp2, then triggers SCF^Skp2 complex formation and E3 ligase activity, also leading to 14-3-3–dependent Skp2 relocalization to the cytosol (42, 43). In contrast with these Akt substrates, we did not observe any changes in the localization and stability of Cables1 by Akt-mediated phosphorylation and 14-3-3 binding. Our results showed that Akt phosphorylation and 14-3-3 binding prevented the function of Cables1 in the induction of apoptosis. Although Cables1 has been reported to enhance p53-induced cell death in U2OS cells and to induce apoptosis in several ovarian cancer cells (3, 32), the exact molecular mechanism by which Cables1 induces apoptosis is still unclear. In this study, we found that Cables1 inhibits the kinase activity of Cdk2 by increasing the pCdk2 Y15 level, which is consistent with a previous report (1). Interestingly, our study also showed that Cables1 increases the level of p21 and decreases the level of pRb, but does not affect the other cell-cycle–related proteins we studied. Cdk2 and p21 play critical roles in the control of apoptosis by regulating the function of several apoptosis-related proteins, such as Foxo1, ASK1, and c-Myc (44, 45). Therefore, the inhibition of Cdk2 and upregulation of p21 by Cables1 may contribute to its induction of apoptosis. Moreover, Cables1 AA had stronger effect than WT on decreasing Cdk2 activity and pRb level, increasing p21 level and inducing apoptosis, indicating that these functions of Cables1 are controlled by the phosphorylation of T44 and T150 residues, which are phosphorylated by Akt. Indeed, expressing exogenous Akt prevents Cables1-induced apoptosis, whereas inactivated endogenous Akt potentiates Cables1-induced apoptosis. Thus, in tumor cells with activated Akt, it is possible that the tumor suppressor function of Cables1 is neutralized through phosphorylation of T44 and T150. In support of this, we observed a correlation between the expression of pCables1 and pAkt in cultured cells, in human lung cancer patient samples, and in tumor tissues of an A549 xenograft mouse model. Our working model proposes that under growth conditions, survival signals activate Akt that in turn phosphorylates Cables1 and recruits 14-3-3 binding (Fig. 7) to prevent the induction of apoptosis by Cables1, which occurs partially through inhibiting Cdk2 activity and upregulating p21.

**Figure 7.** Akt phosphorylation and 14-3-3 binding regulate Cables1-mediated induction of apoptosis. Under growth conditions, survival signals activate Akt, which in turn phosphorylates Cables1 and recruits 14-3-3 binding. Induction of apoptosis by Cables1, which occurs partially through inhibiting Cdk2 activity and upregulating p21, is prevented by Akt phosphorylation and 14-3-3 binding.


20. Tsujii K, Mizumoto K, Yamauchi T, Nishimoto I, Matsuoka M. Differential effect of 14-3-3 on the phosphorylation and interaction of Bad with protein kinase B.


Cables1 Complex Couples Survival Signaling to the Cell Death Machinery


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