Identification of a New Interaction Mode between the Src Homology 2 Domain of C-terminal Src Kinase (Csk) and Csk-binding Protein/Phosphoprotein Associated with Glycosphingolipid Microdomains

Background: Src homology 2 (SH2) domains are noncatalytic regions in tyrosine kinase signaling. Structures of many SH2 domains have been studied, and the regions involved in their interactions with ligands have been elucidated. However, these analyses have been performed using short peptides consisting of phosphotyrosine followed by a few amino acids, which are described as the canonical recognition sites. Here, we report the solution structure of the SH2 domain of C-terminal Src kinase (Csk) in complex with a longer phosphopeptide from the Csk-binding protein (Cbp). This structure, together with biochemical experiments, revealed the existence of a novel binding region in addition to the canonical phosphotyrosine 314-binding site of Cbp. Mutational analysis of this second region in cells showed that both canonical and novel binding sites are required for tumor suppression through the Cbp-Csk interaction. Furthermore, the data indicate an allosteric connection between Cbp binding and Csk activation that arises from residues in the βB/βC loop of the SH2 domain.

Proteins with Src homology 2 (SH2) domains play major roles in tyrosine kinase signaling. Structures of many SH2 domains have been studied, and the regions involved in their interactions with ligands have been elucidated. However, these analyses have been performed using short peptides consisting of phosphotyrosine followed by a few amino acids, which are described as the canonical recognition sites. Here, we report the solution structure of the SH2 domain of C-terminal Src kinase (Csk) in complex with a longer phosphopeptide from the Csk-binding protein (Cbp). This structure, together with biochemical experiments, revealed the existence of a novel binding region in addition to the canonical phosphotyrosine 314-binding site of Cbp. Mutational analysis of this second region in cells showed that both canonical and novel binding sites are required for tumor suppression through the Cbp-Csk interaction. Furthermore, the data indicate an allosteric connection between Cbp binding and Csk activation that arises from residues in the βB/βC loop of the SH2 domain.

Src homology 2 (SH2) domains are noncatalytic regions commonly observed in various types of signal transduction proteins. They function as modules that mediate the interaction between proteins by recognizing a phosphotyrosine (Tyr(P)) in the target proteins. Structural and quantitative binding analyses of many SH2 domains in complexes with related ligand peptides have shown that SH2 domains generally recognize Tyr(P) with three amino acid residues toward the C terminus (Tyr(P)-Xaa1-Xaa2-Xaa3) of the target ligands using two recognition pockets on the surface of the SH2 domains (1–4). One pocket recognizes Tyr(P) of the target primarily through electrostatic interactions and hydrogen bonds, whereas the other recognizes the remaining three amino acids (Xaa1-Xaa2-Xaa3) specifically through hydrophobic interactions. This specificity is considered to generate versatility in the interaction of SH2 domains (5, 6). However, it is still controversial whether the latter pocket alone is sufficient to determine the specificity of the associated interactions (4, 7).

The protein C-terminal Src kinase (Csk) includes SH3, SH2, and kinase domains. This kinase specifically phosphorylates a regulatory Tyr in the C-terminal tail of Src Tyr kinases (SFKs) (8, 9). This event leads to an intramolecular interaction between the Tyr(P)-containing tail and the SH2 domain of the phosphorylated SFK, shifting it to an inactive closed conformation. Thus, Csk negatively regulates the kinase activity of SFKs and plays an important role in physiological functions via signaling pathways for cell proliferation, differentiation, adhesion, and migration (10).

Although SFKs are anchored to membranes via their fattyacylated N termini, Csk, which lacks such a fatty acylation site, exists in the cytoplasm (9). Thus, for Csk to efficiently access SFKs, Csk interacts with the Csk-binding protein (Cbp/PAG), which is localized to membrane microdomains enriched in cholesterol, glycosphingolipids, and lipid rafts, and it is subsequently recruited to the reaction space (11, 12). This interaction is known to occur between the SH2 domain of Csk and the SFK-phosphorylated Tyr-314 of Cbp, but it remains unknown...
whether any other region in Cbp is involved in the interaction (13). In addition, unlike SFKs, which conserve specific Tyr in their C-terminal tails for regulating activity, Csk lacks such a functional tail (9). Nevertheless, the crystal structure of Csk by itself exhibited the existence of two conformers corresponding to active and inactive forms (14). The mechanism of Csk activation is described in detail in a previous review (15). The SH2 domain appears to be required for stabilizing the active form of Csk through connection between the ββ/βC loop (SH2) and the β3/αC loop (kinase). It was also reported that Csk activity is increased through interactions with phosphorylated Cbp or its peptides (13, 16–19). Therefore, it can be speculated that Cbp binding shifts the dynamic equilibrium of at least these two conformers to the more active form. However, the mechanism through which Cbp binding increases Csk activity remains unclear.

To elucidate these mechanisms, we first analyzed the interaction of Csk with various lengths of Tyr-314-containing regions of Cbp using gel filtration chromatography. On the basis of this biochemical result, we determined the tertiary solution structure of the complex of Csk-SH2 with a region of Cbp that contained both Tyr-296 and Tyr(P)-314 using liquid-state nuclear magnetic resonance (NMR) spectroscopy. We found that Csk recognizes not only the four canonical amino acids beginning with Tyr(P)-314 but also a region on the N-terminal side of Tyr(P)-314 that contains Tyr-296.

EXPERIMENTAL PROCEDURES

Preparation of Cbp Mutants—The DNA fragment for rat Cbp peptide (from 289 to 321; Cbp5) was amplified using the pGEX-6P-1 plasmid (GE Healthcare) containing the DNA fragment encoding the region from 195 to 328 of Cbp (Cbp3) by PCR using 5'-CGG GGA TCC AAG AGA TTT AGT TCC TTG TCA-3' and 5'-GCG GAA TTC CGA TTA CTT ACC TGA AGA-3' as primers (Fig. 1A). The amplified DNA fragment was cleaved with BamHI and EcoRI, and the resulting gene was inserted into the BamHI-EcoRI site of pGEX-6P-1. The pGEX-6P-1 plasmid containing the DNA fragment encoding the region from 302 to 321 of Cbp (Cbp6) or a mutant of Cbp5 (Y296F) was produced by the same method as described above using the following primers: 5'-CGC GGA TCC ACA GAA GAG GAG-3' and 5'-CGC GGA TCC ACC TGA AGA-3' as primers (Fig. 1A). The amplified DNA fragment was cleaved with BamHI and EcoRI, and the resulting gene was inserted into the BamHI-EcoRI site of pGEX-6P-1.

Expression and Purification of Various Cbp Peptides—Escherichia coli BL21(DE3) cells were transformed with pGEX-6P-1 containing DNA encoding each of the Cbp peptides and grown in LB media. The cells were incubated at 25 °C with shaking at 240 rpm for 12 h. The expression of GST-fused Cbp peptides was induced by addition of isopropyl β-d-thiogalactopyranoside to a final concentration of 0.1 mM when the absorbance at 600 nm was between 0.3 and 0.6. The cells were further incubated overnight, harvested by centrifugation, and stored at −80 °C. For purification, the cells were dissolved at 4 °C and disrupted by sonication in 100 mM Tris-HCl buffer (pH 7.4) containing 150 mM NaCl, 1 mM EDTA, 5 mM β-mercaptoethanol, and 1% Nonidet P-40 (lysis buffer). After centrifugation, the supernatant was applied to the GSTrap FF affinity column (GE Healthcare), and adsorbed proteins were eluted using lysis buffer (pH 9.0) containing 20 mM reduced glutathione. All purified GST-Cbp fusion proteins were phosphorylated using recombinant Fyn (Millipore) in 50 mM Tris-HCl buffer (pH 7.4) containing 3 mM MgCl₂, 1 mM β-mercaptoethanol, and 4 mM ATP.

Expression and Purification of Csk—Full-length (amino acids 1–450) rat Csk was expressed using a baculovirus vector in Sp9 insect cells, as described previously (13). Cells were lysed using the above-mentioned lysis buffer containing EDTA-free protease inhibitor mixture (Nacalai Tesque) and disrupted using a Dounce homogenizer. The supernatant was collected by centrifugation and applied to the HiTrap Q HP anion exchange column (GE Healthcare) equilibrated with 50 mM Tris-HCl buffer (pH 8.0) containing 1 mM EDTA, 5% glycerol, 5 mM β-mercaptoethanol, and 0.02% octyl-D-glucoside (buffer A). The protein was eluted with a linear gradient of 75–300 mM NaCl. Protein-containing fractions were applied to the HiTrap SP HP cation exchange column (GE Healthcare) equilibrated with buffer A. The protein, eluted with a linear gradient of 50–300 mM NaCl, was applied to the Superdex 200 gel filtration column (GE Healthcare).

Expression and Purification of 13C- and 15N-Labeled Csk-SH2—To obtain 13C- and 15N-labeled proteins, E. coli Origami B(DE3) cells were transformed with pGEX-6P-1(GE Healthcare) containing the gene of Csk-SH2 and were grown in M9 minimal medium containing 1.5 g/liter 15NH₄Cl and 2.0 g/liter creatine care) containing the gene of Csk-SH2 and were grown in M9 minimal medium containing 1.5 g/liter 15NH₄Cl and 2.0 g/liter D-[U-13C₆]glucose, as nitrogen and carbon sources, respectively. For the expression of uniformly 15N-labeled proteins, D-[U-13C₆]glucose was replaced with 4.0 g/liter n-glucose, and 0.1% glycerol was added to the minimal medium. Cells were incubated at 37 °C with shaking. The expression of proteins was induced by the addition of isopropyl β-d-thiogalactopyranoside to a final concentration of 0.5 mM when the absorbance of cells reached an A600 of 0.6. Bacteria were grown for an additional 3 h at 37 °C. Cells were collected by centrifugation and disrupted by sonication in 50 mM Tris-HCl buffer (pH 7.0) containing 400 mM NaCl, 0.1% Tween 20, 1 mM EDTA, and 1 ml of protease inhibitor mixture (Sigma). After centrifugation, the supernatant solution was passed through the DEAE-Sepharose anion exchange column (GE Healthcare) to remove contaminating nucleic acids. The eluate was applied to the GSTrap FF column (GE Healthcare), and the proteins were eluted with 50 mM Tris-HCl buffer (pH 8.0) containing 10 mM reduced glutathione. The eluate was dialyzed for a few hours at 4 °C against 2 liters of the sonication buffer. After addition of 20 μl of PreScission protease (2 units/μl; GE Healthcare) to the solution, it was further dialyzed over 17 h for cleavage of the fusion proteins. In some cases, GST was cut off by the same protease before elution.
New SH2 Domain Binding Mode of Csk to Cbp/PAG

of the fusion protein from the GSTrap FF column during overnight incubation at 4 °C. The proteins were concentrated using an Amicon ultra-4-centrifugal filter unit (Millipore; molecular cutoff, 3000) and were applied to the Superdex 75 gel filtration column (GE Healthcare) with a 20 mm sodium phosphate buffer solution (pH 6.0) containing 50 mM NaCl. The obtained protein (Csk-SH2) ranged from Met-80 to Met-173 with a tag derived from the expression vector (H₂N-Gly-Pro-Leu-Gly-Ser-) attached to the N terminus. Protein concentration was estimated using absorbance at 280 nm (A₂₈₀) with the calculated molar absorption coefficient of 16,000.

Expression and Purification of ¹³C- and ¹⁵N-Labeled Cbp Peptide—The Cbp5 peptide was expressed in E. coli BL21 (DE3) cells (Takara) as a fusion protein with GST, which was labeled with ¹³C- and ¹⁵N-stable isotopes. The procedures for expression and purification were almost the same as those for Csk-SH2. After elution from the GSTrap FF column, the eluate was diluted with the same amount of 20 mM Tris-Cl buffer (pH 8.0). The solution was applied to the HiTrap DEAE FF anion exchange column (GE Healthcare), and the Cbp peptide was phosphorylated by addition of Fyn and ATP prior to removal of GST. A solution containing Csk-SH2 and Cbp5 was applied to the gel filtration column, and the fractions containing the complex were collected.

Protein Binding Assays Using Gel Filtration Chromatography—Each peptide was mixed with equimolar intact Csk or Csk-SH2 in 100 mM Tris-HCl buffer (pH 8.5) containing 150 mM NaCl or 1 mM (NH₄)₂SO₄, 5% glycerol, 5 mM β-mercaptoethanol, and 0.02% octyl-D-glucoside with or without ATP. The solution was applied to the Superdex 200 HR 10/30 (GE Healthcare) gel filtration column; A₂₈₀ was monitored, and each fraction was analyzed by SDS-PAGE. The synthesized peptide and Csk were mixed in a molar ratio of 2:1, and the mixture was assayed as described above.

NMR Spectroscopy—All NMR spectra were acquired at 298 K, except the three-dimensional aromatic ¹³C-edited nuclear Overhauser effect spectroscopy (NOESY), which was performed at 288 K (20), using Bruker DRX-500 and DRX-600 instruments equipped with shielded triple-axis gradient triple-resonance probes, and DRX-800, AvanceII-800, and AvanceIII-950 instruments equipped with z axis gradient triple resonance cryogenic probes. For assignments of ¹H, ¹³C, and ¹⁵N resonances, a series of two- and three-dimensional experiments were performed (21). Two-dimensional ¹H-¹³C heteronuclear single-quantum correlation (HSQC), three-dimensional HNCA, CBCA(CO)NH, HN(CA)CO, and HBCA(CBCA)NH spectra were acquired for assignment of backbone signals. For assignment of the aliphatic side-chain signals, two-dimensional ¹H-¹³C constant time HSQC, three-dimensional ¹³C-edited total correlation spectroscopy (TOCSY) with a mixing time of 79.4 ms, HCCCTOCSY with a mixing time of 22.6 ms, and C(CO)NH and H(CCO)NH with a mixing time of 22.6 ms spectra were used. For assignment of the aromatic side-chain signals, two-dimensional ¹H-¹H double-quantum filtered correlation spectroscopy (22–24) and three-dimensional aromatic ¹³C-edited NOESY with a mixing time of 100 ms were used. The chemical shifts of the ¹H and ¹³C spins in the aromatic residues were assigned by means of two-dimensional (Hβ)Cβ(CγCδ)Hδ and (Hβ)Cβ(CγCδ)He experiments (25). For detection of intermolecular NOEs, a series of filter-related experiments were conducted, namely a ¹³C-filtered/¹³C-edited NOESY using ¹³C,¹⁵N-labeled Csk-SH2 complexed with nonlabeled Cbp5 peptide and a ¹³C-filtered/¹⁵N-edited NOESY and ¹³C,¹⁵N-filtered/¹⁵N-edited NOESY using ¹³C,¹⁵N-labeled Cbp5 peptide complexed with nonlabeled Csk-SH2 (26).

Structure Calculations—The NOE peaks were manually assigned using Sparky. Distance restraints were generated according to the assignment of the NOE cross-peaks, and pseudo-atom corrections were applied to the upper bound restraints involving methyl, methylene, and aromatic ring protons as described previously (27). Torsion angle restraints were derived using TALOS+ (28) with the assigned chemical shifts of ¹H, ¹³Cα, ¹³Cβ, ¹³CO, and ¹⁵N, in reference to the x-ray structure of intact Csk (14). Hydrogen bond restraints, 2.5–3.3 Å for N-O pairs and 1.8–2.3 Å for H-O pairs, were added only to secondary structural regions as confirmed through the corresponding NOE patterns. One disulfide bond between Cys-122 and Cys-164 was confirmed through characteristic ¹³Cα and ¹³Cβ chemical shifts (29) and used as a restraint for structure calculations. Structure calculations with torsion angle dynamics were performed using CYANA-2.1 (30). A total of 100 structures were calculated with 40,000 steps, and after a root-mean-square deviation (r.m.s.d.) for the backbone atoms had reached 1.0 Å, the r⁻⁶ sum averaging method, originally implemented in the CYANA calculations, was applied instead of the pseudo-atom corrections. Finally, the 20 structures with the lowest target functions were selected. Molecular models were prepared using MOLMOL (31) and UCSF Chimera (32). Hydrogen bonds in the complex structure were defined using UCSF Chimera. An accessible surface area for each residue was calculated using MOLMOL. The electrostatic potential of the complex structure was calculated using Delphi (33). All the amino acid sequence alignments were performed using ClustalW, version 2.0 (34).

Relaxation Analysis—The ¹⁵N-spin longitudinal (R₁) and transverse (R₂) relaxation rates and [¹H]–¹⁵N steady-state NOE values were acquired on Bruker DRX-500 and AvanceII-800 spectrometers (26). R₁ relaxation delays were set at 5, 100, 200, 300, 400, 550, 700, 850, 1000, 1150, and 1300 ms, and R₂ relaxation delays were set at 7.2, 36, 72, 108, 144, 180, 216, 2528, 324, and 360 ms in corresponding experiments. Single exponential curve fitting was performed using Sparky (35). The model-free analysis developed by Lipari and Szabo (36), with the isotropic rotational diffusion model assumed, was performed using Tensor2 (37). All NMR data were processed and analyzed using NMRPipe (38) and Sparky, respectively.

zz-Exchange Spectroscopy—Chemical exchange associated with the interaction was monitored using ¹⁵N-labeled Csk-SH2 mixed with each of the nonlabeled phosphopeptides, Cbp5, Cbp6, and Cbp7, at a molar ratio of 2:1 by two-dimensional ¹⁵N zz-exchange spectroscopy (39–41) on a Bruker Avance-III 950 spectrometer with a cryogenic TCI probe. The spectra were acquired with mixing times, τmix, of 0, 20, 50, 100, 200, 350, 550, and 800 ms. For each time point, four peaks observed with
intensities denoted by $I_{AA}(\tau_m)$, $I_{BB}(\tau_m)$, $I_{BA}(\tau_m)$, and $I_{AB}(\tau_m)$ are, respectively, governed by Equation 1,

$$I_{AA}(\tau_m) = I_{AA}(0)p_a + p_b \exp(-k_{ex}\tau_m) \exp(-R_1\tau_m) \quad \text{Eq. 1}$$

$$I_{BB}(\tau_m) = I_{BB}(0)p_b + p_a \exp(-k_{ex}\tau_m) \exp(-R_1\tau_m)$$

$$I_{BA}(\tau_m) = I_{BA}(0)p_b(1 - \exp(-k_{ex}\tau_m)) \exp(-R_1\tau_m)$$

$$I_{AB}(\tau_m) = I_{AB}(0)p_a(1 - \exp(-k_{ex}\tau_m)) \exp(-R_1\tau_m)$$

where $p_i$ represents the relative population for the site $i$; $k_{ex}$ represents the sum of the forward, $k_{ex}$, and reverse, $k_{rev}$, kinetic rate constants for interconversion between the sites; and $R_1$ represents the longitudinal relaxation rate of the $^{15}$N nucleus in the observed spin system. Note that $k_1$ is an apparent pseudo-first-order rate constant that corresponds to the product of the second-order association rate constant, $k_{on}$, and the concentration of free Cbp, i.e. $k_1 = k_{on}[\text{Cbp}]_{\text{free}}$. The total cross-peak intensity ($I_{BA}(\tau_m) + I_{AB}(\tau_m)$) for each time point, $\tau_m$, was fitted to the sum of the third and forth equalities given in Equation 1 to obtain $k_a$ and $R_1$, $p_A$ and $p_B$ were estimated from the intensity ratio of the monomer and complex peaks in a two-dimensional $^1$H-$^{15}$N HSQC spectrum. All NMR data were processed and analyzed using NMRPipe and Sparky, respectively.

**Cells and Gene Transfer—**A549 cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS). Genes were transfected into the A549 cells using a previously described method (42). In brief, cDNAs encoding wild-type human Cbp and its mutant, in which Tyr-296 was substituted with Phe (Y296F) or both Tyr-306 and Tyr-314 were fused with GST and expressed in E. coli. Moreover, we used two types of running buffers in the gel filtration assay, one containing 150 mM NaCl and the other containing 1 M (NH$_4$)$_2$SO$_4$. The results showed that phosphorylated Cbp3 and Cbp5 eluted from the column as one peak and formed a complex with Csk at both salt concentrations, whereas the nonphosphorylated peptides eluted separately from Csk or later than the corresponding phosphorylated peptides without dominantly forming any complex with Csk (Fig. 1B). Moreover, the peak corresponding to Csk in the experiment using the phosphorylated peptides appeared earlier than that using the nonphosphorylated peptides, indicating that the former eluted contained Csk of high molecular weight, because of complex formation with the phosphorylated peptides (Fig. 1B). When Tyr-314 was not phosphorylated, none of the examined peptides dominantly formed a complex in either salt concentration. This indicates that the phosphorylation of Tyr-314 is at least necessary for interaction with Csk. Interestingly, shorter peptides lacking N-terminal regions, namely phosphorylated Cbp6 and Cbp7, formed a complex with Csk during gel filtration in the running buffer containing 150 mM NaCl but not in the buffer containing 1 M (NH$_4$)$_2$SO$_4$ (Fig. 1B). Because the interaction involving the phosphate group of Tyr(P)-314 was weakened by a high salt concentration, it is expected that this interaction is electrostatic in nature. The differences observed in elution patterns of Cbp5 and Cbp6 at high salt concentrations indicate that a region within Cbp5 but outside Cbp6 (i.e., 289–302) generates an additional interaction to Tyr(P)-314 that enables high affinity binding to Csk even at high salt concentrations.

**RESULTS**

**Gel Filtration Assay Using Cbp Peptides of Different Lengths—**To identify the Cbp region responsible for the interaction with Csk, we assayed the interaction between each of the Cbp peptides of different lengths containing Tyr(P)-314 or Tyr-314 and intact Csk using gel filtration chromatography and SDS-PAGE. Deletion mutants of Cbp used for the assay, Cbp3 (195–328), Cbp5 (289–321), Cbp6 (302–321), and Cbp7 (312–321; Fig. 1A), were fused with GST and expressed in E. coli, and intact Csk was expressed in a baculovirus/insect SF9 cell system. The GST-fused Cbp peptides were phosphorylated at Tyr-314 by addition of recombinant Fyn and ATP. We confirmed that this reaction did not phosphorylate Tyr-296 by a peptide mass fingerprinting method. Nonphosphorylated Cbp peptides were prepared as controls by mixing each peptide with Fyn in the absence of ATP.

Each of the GST-fused peptides, containing Tyr-314 or Tyr(P)-314, was mixed with Csk, and the solutions were applied to a gel filtration column. The eluted fractions were applied to SDS-PAGE to detect the contents. To evaluate the affinity qualitatively, we used two types of running buffers in the gel filtration assay, one containing 150 mM NaCl and the other containing 1 M (NH$_4$)$_2$SO$_4$. The results showed that phosphorylated Cbp3 and Cbp5 eluted from the column as one peak and formed a complex with Csk at both salt concentrations, whereas the nonphosphorylated peptides eluted separately from Csk or later than the corresponding phosphorylated peptides without dominantly forming any complex with Csk (Fig. 1B). Moreover, the peak corresponding to Csk in the experiment using the phosphorylated peptides appeared earlier than that using the nonphosphorylated peptides, indicating that the former elute contained Csk of high molecular weight, because of complex formation with the phosphorylated peptides (Fig. 1B). When Tyr-314 was not phosphorylated, none of the examined peptides dominantly formed a complex in either salt concentration. This indicates that the phosphorylation of Tyr-314 is at least necessary for interaction with Csk. Interestingly, shorter peptides lacking N-terminal regions, namely phosphorylated Cbp6 and Cbp7, formed a complex with Csk during gel filtration in the running buffer containing 150 mM NaCl but not in the buffer containing 1 M (NH$_4$)$_2$SO$_4$ (Fig. 1B). Because the interaction involving the phosphate group of Tyr(P)-314 was weakened by a high salt concentration, it is expected that this interaction is electrostatic in nature. The differences observed in elution patterns of Cbp5 and Cbp6 at high salt concentrations indicate that a region within Cbp5 but outside Cbp6 (i.e., 289–302) generates an additional interaction to Tyr(P)-314 that enables high affinity binding to Csk even at high salt concentrations.
FIGURE 1. Gel filtration assay using Cbp peptides of different lengths. A, schematic representation of intact Cbp and its various peptide fragments that appear in the text. The N- and C-terminal residue numbers are indicated on the left and right of each peptide. Tyr(P) represents the 296th or 314th phosphorylated Tyr.

B, SDS-PAGE of gel-filtered fractions, including GST-fused Cbp3, Cbp5, Cbp6, or Cbp7, and Csk showing a secondary binding region between Lys-289 (N-terminal of Cbp5) and Ser-302 (N-terminal of Cbp6) on the N-terminal side of the Tyr(P) of Cbp. NP and P indicate the corresponding nonphosphorylated and phosphorylated Cbp peptides, respectively. Fractions simultaneously containing both the Cbp peptide and Csk indicate that the complex was formed in the corresponding condition. Asterisks indicate nonphosphorylated samples.

C, SDS-PAGE of gel-filtered fractions, including GST-fused Cbp5-Y296F and Csk or GST-fused Csk-SH2, shows that Tyr-296 of Cbp is at least involved in the interaction with Csk or Csk-SH2. NP and P indicate the corresponding nonphosphorylated and phosphorylated Cbp peptides, respectively. All the gel filtration assays were performed in a buffer containing 150 mM NaCl (left) or 1 M (NH₄)₂SO₄ (right).
Gel Filtration Assay Using Y296F Mutant and Phospho-Tyr-296—Here a question emerged as to which residues between 289 and 302 mediate the secondary interaction. We focused on Tyr-296 because it is a unique Tyr in this region, and we expected its phosphorylation to strengthen the interaction with Csk in combination with phosphorylation of the canonical Tyr-314. To study the possible secondary interaction region, we prepared a mutant of GST-fused Cbp5 peptide in which Tyr-296 was replaced with Phe, and we assayed its interaction with Csk or Csk-SH2 using gel filtration chromatography as described above. Although both interactions were maintained in 150 mM NaCl, they were broken in the presence of 1 M (NH₄)₂SO₄ (Fig. 1C). As revealed by NMR spectroscopy (see below), replacement of Tyr-296 with Phe in Cbp5 weakened its interaction with Csk and Csk-SH2, probably because of the absence of a hydrogen bond between the hydroxyl group of Cbp Tyr-296 and the guanidinium group of Csk-SH2 Arg-107.

Overall Structure of the Complex of Csk-SH2 with Cbp5—The biochemical data described above, including gel filtration assays, indicate that Tyr(P)-314 interacts with Csk and that at least Tyr-296 contributes to binding affinity. To obtain detailed structural information about this interaction, we have determined the tertiary structure of the complex of Csk-SH2 with Cbp5 using multidimensional and multiresonance NMR spectroscopy. We confirmed that Csk-SH2 and phosphorylated Cbp5 (without GST) maintained a stable complex during gel filtration, whereas Csk-SH2 and nonphosphorylated Cbp5 did not by MALDI-TOF mass spectrometry. In analysis, we used not only isotope-labeled Csk-SH2 but also isotope-labeled Cbp5, which was overexpressed as a GST fusion protein in

FIGURE 2. Comparison of the amino acid sequence of Cbp5 with those of various peptide ligands associated with SH2 domains registered in PDB. The sequences were aligned using ClustalW2. The sequence of Cbp5 is shown in red, A, comparison of Cbp5 with other peptide ligands associated with different SH2 domains. B, comparison of Cbp5 with other peptide ligands that can interact with Csk-SH2. The secondary binding sites with Csk-SH2 are shown in orange. C, comparison of the N-terminal sequence from Tyr(P) of Cbp5 with the corresponding sequences of other peptide ligands associated with representative SH2 domains. The secondary binding sites with Csk-SH2 are shown in orange.
E. coli. Most of the phosphopeptides that have been used to date for structural analyses were not labeled because they have often been synthesized chemically as phosphorylated forms. However, with the isotopically labeled phosphopeptide of Cbp, we successfully obtained a high quality solution structure of the complex. The Cbp peptide was phosphorylated by Fyn and ATP during purification, and its purity was estimated to be >99% by mass spectrometry. The sequence of Cbp5 includes 33 amino acid residues, which was much longer in the N-terminal direction compared with the sequences of various other peptides that have been used for structural analyses of complexes with SH2 domains (Fig. 2A). This N-terminal sequence is not conserved in other ligands that interact with Csk-SH2 (Fig. 2B).

Fig. 3A shows an overlay of the final 20 structures, which exhibited the best target functions among 100 calculated structures (PDB code 2rsy). In Fig. 3B, a ribbon diagram of the representative structure with the minimum target function is presented. The coordinates of the backbone atoms of Cbp5 were well defined in the wide range from Met-80 to Lys-319 with respect to the coordinates of Csk-SH2. These gave an average r.m.s. value to the mean structure of 0.26 ± 0.04 Å with respect to the backbone atoms in the range from Met-80 to Lys-171 of Csk-SH2 (green) and from Tyr-296 to Lys-319 of Cbp5 (khaki). B, ribbon diagram of the representative structure shows (coral color) an α-helix from Glu-307 to Met-313 and a turn from Arg-299 to Asp-302.

Conventional Interaction Mode—As shown in Fig. 4A, the region of Cbp5 that was expected to be specifically recognized by Csk-SH2, pYSSV, adopted an extended conformation in the complex. In the conformation, the aromatic and phosphate residues of associated ligands (45). In addition, the phosphate group of Tyr(P)-314 formed hydrogen bonds with the hydroxyl group of Ser-109 in βB, the amino (–NH₂) group of Asn-111 in βC, and the amide group of Tyr-116 through the hydroxyl group of Thr-117 in βC (Fig. 4A). Another intermolecular hydrogen bond was found between the amide group of the next

**TABLE 1**

| Statistics of the final 20 best structures of Csk-SH2 and Cbp (289–321) |
|---------------------------------------------------------------|
| **NOE upper distance limits**                                 |
| Total                                                         | 2935 |
| Intrasubunit ([i − j] = 0)                                     | 604  |
| Sequential ([i − j] = 1)                                      | 779  |
| Medium range ([i − j] > 5)                                    | 552  |
| Long range ([i − j] > 5)                                      | 1000 |
| Intermolecular                                               | 202  |
| Hydrogen bonds                                               | 33   |
| Disulfide bond                                               | 1    |
| Dihedral angle restraints violations                          |
| Number >5°                                                   | 0    |
| Mean r.m.s. violations from distance restraints               | 0.0043 ± 0.0005 Å |
| r.m.s deviation to the mean coordinates*                     |
| Backbone atoms                                               | 0.26 ± 0.04 Å |
| Heavy atoms                                                  | 0.68 ± 0.05 Å |

* Pairwise r.m.s.d. was calculated for residues 80–172 and 296–318 among 20 refined structures.
Another interaction region found in the structure of the complex of Csk-SH2 with Cbp5. A, interaction region between Csk-SH2 and Cbp5 is circled in black in the ribbon diagram of the complex of Cbp5 with Csk-SH2. Amino acid residues involved in the interaction are represented using a stick model. B, Cbp5 exhibits an amphipatic character, with hydrophilic residues exposed to the solvent and hydrophobic residues and aliphatic chains oriented inward. Positively and negatively charged residues are labeled in cyan and red, respectively, and hydrophobic residues are labeled in black. C, Tyr-296 and Tyr(P)-314 of Cbp5 are accommodated in the binding pocket of Csk-SH2. The two residues are represented using a stick model. An α-helix and a turn of Cbp formed on Csk-SH2 bring the two Tyr residues close to each other. D, residues of Csk-SH2 interacting electrostatically with the hydroxyl group of Tyr-296. Hydrogen bonds are shown by cyan lines. Amino acid residues of Cbp5 and Csk-SH2 are labeled in orange and black, respectively. E, residues additionally found as contributors to the second interaction site. Amino acid residues of Cbp5 and Csk-SH2 are labeled in orange and black, respectively. Hydrogen bonds are shown as cyan lines. Csk-SH2 is shown in green and Cbp5 in khaki.

Another Novel Interaction Region between Csk-SH2 and Cbp5—Consistent with the results of our gel filtration experiments, structure determination of the complex revealed an additional interaction region in the N-terminal direction from Tyr(P)-314 of Cbp5 (Fig. 5). This region contained a unique α-helix between Glu-307 and Met-313 and a hydrophobic core comprising aliphatic Lys-297 and Arg-299 and side chains of Leu-305, Ile-310, and Met-313. The hydrophilic side chains of Lys-297, Arg-299, Glu-301, Asp-302, Glu-307, Glu-308, and Glu-309 were exposed to the solvent (Fig. 5B). In addition, the residues ranging from Arg-299 to Asp-302 took a turn conformation that exposed the hydrophilic side chains of Glu-300, Glu-301, and Asp-302 to the solvent. The structure of Cbp5, at least in its complex with Csk-SH2, Tyr-296 and Tyr(P)-314 were positioned such that both side chains were inserted together into the pocket of Csk-SH2 with their aromatic rings arranged side by side (Fig. 5C). Most regions (99 ± 1%) of Tyr-296 were buried in the interior of the complex structure, and the accessible surface area of its main chain and side chain was 1 ± 1 (%). This is in contrast to the accessible surface area of Tyr-296 in the free state, which is 30 ± 3 (%). Fig. 5D shows the residues of Csk-SH2 that were involved in the interaction with Tyr-296. The position of the Tyr-296 hydroxyl group allowed electrostatic interactions with Csk-SH2 via the phosphate group of Tyr(P)-314, the main-chain amide group of Thr-110, and the main-chain amide and side-chain amino groups of Asn-111. It also formed a hydrogen bond with the guanidinium group of Arg-107. These data explain the decreased affinity observed in gel filtration experiments when Tyr-296 was replaced with Phe lacking this hydroxyl group. Phosphorylation of Tyr-296 caused decreased affinity between Cbp and Csk-SH2, probably because of steric hindrance and electrostatic repulsion of bulky negatively charged phosphate groups on Tyr-296 and Tyr-314. In addition, the complex structure revealed van der Waals contacts between Arg-299/Glu-300 and Thr-110/Asn-111, and the filtered NOESY experiments exhibited strong intermolecular NOE peaks between these residues. These data indicate that Arg-299 and Glu-300 are also involved in the interaction (Figs. 5E and 6).

We compared the two-dimensional 1H–15N HSQC spectra of Csk-SH2 before and after complex formation (Fig. 7A) and plotted chemical shift differences against the amino acid sequence (Fig. 7B). No significant difference was observed in the overall peak distribution in the spectra. However, Thr-110 and Asn-111 exhibited the weighted average chemical shift deviations over 1.0 ppm (Fig. 7B). These data correspond to interactions of Thr-110 with Tyr-296 and Arg-299 as well as Glu-300 of Cbp and Asn-111 with Tyr-296, Arg-299, and Tyr(P)-314 of Cbp (Figs. 4 and 5). Because we labeled Cbp5 with isotopes, we compared the two-dimensional 1H–15N HSQC spectra of Cbp5 before and after complex formation (Fig. 7C) and plotted chemical shift differences against the amino acid sequence (Fig. 7D). The complex formation led to a wider distribution of Cbp5 amide 1H–15N cross-peaks (Fig. 7C). In particular, Lys-297 exhibited a chemical shift change as large as 1.0 ppm (Fig. 7D). The structure of the complex indicated a hydro-
gen bond between the carbonyl group in the main chain of Lys-297 and the amino group of the side chain of Asn-111, indicating that Lys-297 also contributes to the interaction of Cbp5 with Csk-SH2 (Fig. 5E). Among known ligands of SH2 domains, no residues that correspond to Tyr-296, Lys-297, Arg-299, and Glu-300 of Cbp and Thr-110 or Asn-111 of Csk-SH2 are definitely conserved (Fig. 2C). Likewise, in other SH2 domains, the residues that correspond to Thr-110 and Asn-111 are various and not conserved as expected (45).
FIGURE 8. Tyr-296 is required for a stable Cbp-Csk interaction and tumor-suppressive function of Cbp. A, whole cell lysates (WCL) from A549 cells and Cbp-expressing wild-type cells, Y296F mutant, and Y296F/Y314F double mutants were immunoprecipitated with the indicated antibodies. Cbp was then immunoprecipitated by S. aureus protein A-agarose beads and then immunoblotted with the indicated antibody. Asterisks indicate the location of the immunoglobulin. B, effects of the expression of Cbp and its YF mutants on the tumor growth of A549 cells were examined using colony formation assays in soft agar. Colonies were stained with 3-(4,5-dimethylthiazol-2–2-yl)-2,5-diphenyltetrazolium bromide 10 days after plating. The colony numbers per cm² are presented as mean ± S.E. from three independent experiments. C, effects of the expression of Cbp and another mutant, AASAA, on the tumor growth of A549 cells were examined as described in B. The mutated amino acids are shown in bold letters in the sequence above the graph.

Involvement of Tyr-296 in the Cbp-Csk Interaction in Cells—To evaluate the physiological role of Tyr-296 in stabilizing the Cbp-Csk interaction, we examined the function of a Cbp mutant (Y296F) with a substitution of Tyr-296 by Phe in human lung cancer A549 cells with marked up-regulation of c-Src. We have recently shown that the expression of Cbp is extensively down-regulated in these cells and that the re-expression of Cbp inactivates c-Src and suppresses tumor growth by facilitating the recruitment of Csk to the membrane (44). In Cbp-expressing cells, the Cbp-Csk interaction was readily detected by immunoprecipitation (Fig. 8A). However, in mutant cells, Y296F bound very weakly to Csk. Furthermore, a double mutant, in which Tyr-296 and Tyr-314 were replaced with Phe (Y296/314F), completely failed to interact with Csk. Inversely, the Tyr phosphorylation levels of Cbp, which reflect the activation status of c-Src, were significantly augmented when Cbp mutants were expressed. These observations demonstrate that Tyr-296 contributes to stabilizing the Cbp-Csk interaction even in cells. We next assessed the role of Cbp in cellular function by examining the effects of the expression of Cbp on the colony-forming ability of tumor cells. As described previously, the expression of Cbp significantly suppressed colony formation in A549 cells (Fig. 8B). However, this growth suppression was significantly attenuated when Y296F was expressed. The expression of Y296/314F did not induce significant suppression of the colony-forming ability of these cells, revealing that Tyr-296 supports the tumor-suppressive function of Cbp by stabilizing the Cbp-Csk interaction.

Interestingly, another mutant, in which Tyr-296, Lys-297, Arg-299, and Glu-300 were replaced with Ala (AASAA), attenuated growth suppression more than Y296F did. This effect was comparable with that of mock and Y314F despite retention of Tyr-314, which is critical for the Cbp-Csk interaction (Fig. 8C). As described above, these residues in the wild-type specifically interact with Thr-110 or Asn-111 of Csk-SH2 and form the core of the second interaction region. Hence, it is clear that simultaneous replacement of all of these residues in AASAA led to decreased Csk-binding affinity and that the second recognition region is at least required for tumor suppression via the Cbp-Csk interaction.

Cbp Binding to the SH2 Domain Changes Its Structure to the More Active Form—Previous x-ray analyses of intact Csk demonstrated crystal structures of both active and inactive forms, which probably establish an equilibrium in solution (14). Thus, in order for Csk to be completely active, its conformation must shift to the active form and be maintained at least during the kinase reaction. Interestingly, past experiments showed that Csk activity increased upon interaction with a Cbp phosphopeptide consisting of only six residues (AMPYSSV) or 10 residues (ISAMPYSSVKN) (13, 16–19). However, overlaying the solution structure of the complex on intact crystal structures of the free form showed that the Cbp5 peptide was distant from the catalytic domain (Fig. 9A). Therefore, it is conceivable that the catalytic domain is activated through a possible conformational change in Csk-SH2 triggered by Cbp binding to it.

To investigate how the effect of Cbp binding is structurally transmitted to the SH2 domain, we examined the chemical shift perturbation observed for Csk-SH2 upon binding with Cbp5. Thus, mapping of the perturbed residues on the structure of Csk-SH2 suggested that its βB/βC loop and βD strand underwent conformational or dynamic changes following Cbp5 binding (Fig. 10A). We next compared the structure of Csk-SH2 of the complex with the crystal structure of the free form, which had previously been determined by x-ray analysis (PDB code 3EAC). Consistent with the above-mentioned chemical shift changes, these structures in the βB/βC loops were distinctly different from each other (Fig. 10A). Furthermore, in our complex structure, hydrogen bonds were found in the βB/βC loop around Tyr(P)-314, including those between the amide group of Ser-109 and the carbonyl group of Asp-115, the hydroxyl group of Ser-109 and the amide group of Tyr-112, and the carbonyl group of Tyr-112 and the amide group of Asp-115 (Fig.
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FIGURE 9. Connection between Cbp5 and the kinase domain through Csk-SH2. A, relative configuration between Cbp5 and the kinase domain revealed by overlaying of the solution structure of Csk-SH2 in complex with Cbp5 and the crystal structure of intact Csk (PDB code 1K9A). Csk-SH2 and Cbp5 in the complex are colored in green and khaki, respectively, and intact Csk is colored in gray. The region corresponding to C is shown with a black square. B, intramolecular NOEs observed among Ser-109, Tyr-112, and Asp-115 supporting the presence of the hydrogen bonds between them. The corresponding regions are shown from a 15N-edited NOESY spectrum of 15N-labeled Csk-SH2 in complex with unlabeled Cbp5. C, insertion of Ala-228 into the hydrophobic pocket formed by Tyr-116, Tyr-133, Leu-138, and Leu-149, as observed in the crystal structure of Csk. These residues are shown with a stick model. The pocket is highlighted with a brown circle.

FIGURE 10. Characterization of the structure of Csk-SH2 in complex with Cbp5. A, mapping of the chemical shift deviations in Csk-SH2 upon binding with Cbp5 on the structure of Csk-SH2 and comparison of the solution structure of Csk-SH2 of the complex with the crystal structure of the free form (gray; PDB code 3EAC). The residues with weighted chemical shift deviations larger than 0.8 ppm are colored in red; those between 0.8 and 0.4 ppm are reddish orange; and those between 0.4 and 0.3 ppm are orange. Pro residues are colored in black. Both Tyr-116 and Tyr-133, which are involved in the hydrophobic pocket in contact with Ala-228 in the kinase domain of active Csk (Fig. 9), underwent significant chemical shift changes that were not due to direct contact with Cbp5. B, hydrogen bond network generated between Tyr(P)-314 of Cbp5 and its nearby residues in the βB/βC loop of Csk-SH2. The side chains of these residues are represented by a stick model and hydrogen bonds are shown as blue lines. Amino acid residues of Cbp5 and Csk-SH2 are labeled in orange and black, respectively. C, comparison between the solution structures of the complex of Csk-SH2 with Cbp5 (green) and the crystal structure of intact Csk-SH2 in the active form (gray; PDB code 1K9A). Hydrogen bonds are shown as cyan lines. The r.m.s.d. value of the heavy atoms in the main chains was 0.81 Å. D, βB/βC loops, which have almost the same conformation in both structures, are highlighted with black squares. D, comparison between the solution structure of the complex and the crystal structure of intact Csk-SH2 in the inactive form (PDB code 1K9A). The r.m.s.d. value of the heavy atoms in the main chains was 0.92 Å. βB/βC loops, indicated with a double-headed arrow, take different conformations in the two structures.

NMR Relaxation Analysis—To evaluate changes in the structural dynamics of Csk-SH2 induced by Cbp binding, we measured 15N R1 and R2 relaxation rates, and 1H-15N steady-state NOE values of uniformly 15N-labeled Csk-SH2 with and without uniformly 15N-labeled Cbp5 peptide. Resonances showing severe overlap or low intensity were excluded from the study. The average rotational correlation time τc was estimated as 5.8 and 8.4 ns for Csk-SH2 in the free and complex forms, respectively. Using these relaxation data, we analyzed residue-specific internal motions of Csk-SH2 in the two forms in terms of the order parameters squared (S2) by the model-free approach (Fig. 11, A and B). Fig. 11C shows the difference between the S2 values of Csk-SH2 with and without Cbp5. The two forms of Csk-SH2 exhibited no significant difference in overall dynamics. The average S2 values for Trp-82–Val-172 of Csk-SH2 were 0.85 ± 0.01 (one S.D.) in the free form and 0.83 ± 0.01 in the complex form. However, the S2 values in loop regions exhibited significant differences between the two forms. In particular, Leu-86 in the N-terminal region, Ser-136 in the βD/βE loop (turn), Asp-141 in the βE/βF loop, and Gly-162 in the αB/βG loop had S2 values that were ~0.1 less in the bound form. In contrast, Ser-109 and Asn-111 in the βB/βC loop and Asp-161...
in the αB/βG loop had greater $S^2$ values (about 0.1) in the bound form. These differences are mapped on the structure of Csk-SH2 (Fig. 11D). The $S^2$ values of Cbp5 in the complex exceeded 0.65 (mean; 0.80 ± 0.01) over the range of 22 amino acids from Tyr-297 to Asn-318 (Fig. 11D). These data indicate that a wide range of the region, including a secondary interaction site, stably bound with Csk-SH2.

**zz-Exchange Spectroscopy**—To evaluate the affinity of Csk-SH2 for the secondary binding site of Cbp, we performed zz-exchange spectroscopy of $^{15}$N-labeled Csk-SH2 mixed with each of Cbp5, Cbp6, and Cbp7. Kinetic data were analyzed from well resolved peaks of two residues: Gly-162, which binds directly to Cbp, and Val-172. Although Val-172 is located at the opposite side of the binding region, it exhibited an NMR peak pattern similar to that of Gly-162 because of a possible small conformational change synchronized with Cbp binding. As a result, Cbp5, the longest peptide that includes the secondary binding site, exhibited distinctly lower exchange rates ($k_{off}$ = 0.80 s$^{-1}$ for Gly-162), whereas both Cbp6 and Cbp7 exhibited ∼8–25 times higher $k_{off}$ values for Gly-162 ($k_{off}$ = 6.69 s$^{-1}$ for Cbp6 and $k_{off}$ = 23.1 s$^{-1}$ for Cbp7) (Table 2). These results evidently indicated that longer peptides interacted more strongly and that Cbp5 had the highest affinity because of the additional interaction, which Cbp6 and Cbp7 lack.

**DISCUSSION**

It is generally accepted that SH2 domains bind to ligands by specifically recognizing a Tyr(P) along with the +1 to +3 amino acid residues that follow in the C-terminal direction. Our complex structure of Csk-SH2 with Cbp5 showed that Tyr(P)-314 of Cbp and the three residues that follow toward the C terminus interacted with Csk-SH2 in the same manner as observed in many other phosphopeptides that interact with normal SH2 domains. This observation was supported by our gel filtration experiments and indicates that Csk-SH2 is a normal SH2 domain that mediates typical interaction patterns. However, we have identified a novel binding region of Cbp with Csk-SH2. Analyses of intermolecular NOE and $^{15}$N-spin relaxation rates indicated that the region of Cbp that interacts with Csk-SH2 spans the 23 amino acid residues between Tyr-296 and Asn-318.

Even when nonlabeled Csk-SH2 was added to a solution of nonphosphorylated $[^{15}$N]Cbp5, the two-dimensional $^1$H–$^{15}$N HSQC spectrum of Cbp5 exhibited neither chemical shift changes nor signal broadening in its amide resonance peaks. None of the amide peaks that were assigned to a region close to Tyr-296 in the sequence underwent any chemical shift perturbation either. This indicates that the novel N-terminal interaction region of Cbp alone is not sufficient to enable Cbp5 to bind to Csk-SH2. In fact, addition of a phosphate group to Tyr-314 induced amide chemical shift changes of not only Tyr-314 but also Asp-302 and Lys-289, which are distant from Tyr-314 in the context of the amino acid sequence. In addition, our preliminary molecular dynamic simulation suggested a bent conformation of Cbp5, with Tyr(P)-314 close to Tyr-296 even in its
free state, rather than a randomly fluctuating extended conformation as often seen in short peptides (data not shown). Thus, there may be interference between Tyr-314 and Tyr-296, and when Tyr-314 is not phosphorylated, Tyr-296 may also fail to interact with Csk-SH2. In either case, the result shows that phosphorylation of Tyr-314 dominantly controls association and dissociation with Csk-SH2.

Although the second binding region containing Tyr-296 certainly contributes to the interaction, it appears to enhance the selectivity of Csk-SH2 for this ligand. Intact Cbp possesses six Tyr residues that can undergo phosphorylation (13). Among them, the three residues following Tyr-224 (pYASV) are almost the same as those following Tyr-314 (pYSSV) except for Ala and Ser at the +1 position in the respective sequences. In fact, another ligand for Csk-SH2, SIT, possesses this pYASV sequence as the binding site (50). In addition, the short peptide GDGpYXXXXPLL, in which pYXXX corresponds to any of the sequences pY(T/A/S)(K/R/Q/N)(M/I/V/R), is reportedly recognized by Csk-SH2 (51). Furthermore, sequential alignment of Cbp5 with other ligands containing Tyr(P) residues showed that the N-terminal interaction region around Tyr-296 of Cbp5 is unique and that the corresponding regions are rich in variety in the sequences of other ligands (Fig. 2B). These results indicate that the sequence that includes a few residues following Tyr-296 of Cbp5 with other ligands containing Tyr(P) residues showed that the N-terminal interaction region around Tyr-296 of Cbp5, Cbp6, or Cbp7.

### TABLE 2

| Peptide | SH2   | $P_{i/PA}$ | $k_{on}$ | $k_{off}$ | $R_1$ |
|---------|-------|------------|----------|-----------|-------|
| Cbp5   | Gly-162 | 1.04 ± 0.02 | 0.80 ± 0.05 | 2.00 ± 0.01 | 2.00 ± 0.05 |
|        | Val-172 | 1.22 ± 0.02 | 0.86 ± 0.08 | 2.00 ± 0.01 | 2.00 ± 0.05 |
| Cbp6   | Gly-162 | 1.04 ± 0.02 | 0.80 ± 0.05 | 2.00 ± 0.01 | 2.00 ± 0.05 |
|        | Val-172 | 1.04 ± 0.02 | 0.80 ± 0.05 | 2.00 ± 0.01 | 2.00 ± 0.05 |
| Cbp7   | Gly-162 | 0.90 ± 0.05 | 0.80 ± 0.05 | 2.00 ± 0.01 | 2.00 ± 0.05 |
|        | Val-172 | 0.90 ± 0.05 | 0.80 ± 0.05 | 2.00 ± 0.01 | 2.00 ± 0.05 |

A two-dimensional $^1$H-$^15$N HSQC spectrum of Cbp5 in the free form exhibited most of the main-chain amide signals crowding within a $^1$H resonance range between 8.0 and 8.5 ppm, suggesting that no distinct conformation exists dominantly in the free state of Cbp (Fig. 5C). Although, as described above, particular conformations may be taken, these are unstable. However, Cbp5 in the complex state had a hydrophobic core (Leu-305, Ile-310, and Met-313 and the aliphatic parts of Lys-297 and Arg-299) and hydrophobic residues on the surface (Lys-297, Arg-299, Glu-301, Asp-302, Glu-307, Glu-308, and Glu-309) and was stabilized by adopting a characteristic conformation (Fig. 5B). These characteristics eluded studies of shorter peptide ligands in complex with SH2 domains. Importantly, this structural compaction allowed the Cbp5 peptide to interact with a limited region in the ββ/βc loop of Csk-SH2, in which a characteristic hydrogen bond network was observed.

Our NMR study revealed that Cbp binding to Csk-SH2 formed a hydrogen bond network around the ββ/βc loop, with the phosphate group of Tyr(P)-314 as the center. According to the crystal structure of intact Csk in the active form, Tyr-116 in this hydrogen bond network forms a hydrophobic pocket with Tyr-133, Leu-138, and Leu-149 that interacts with the catalytic domain through the methyl group of Ala-228 (Fig. 9C). However, these residues are geometrically distant from each other in the inactive form. Interestingly, a recent study reported that any mutation in Tyr-116, Leu-138, or Leu-149 in the hydrophobic pocket or in Ala-228 in the kinase domain to Ala or Gly decreases Csk activity to 10–20% of the wild type in the absence and presence of a Cbp peptide (18). These experiments demonstrate that binding of Ala-228 to this hydrophobic pocket is essential for Csk activation. Cumulatively, these observations...
suggest that a hydrogen bond network between Tyr(P)-314, probably involving the novel binding sites, of Cbp and the above-mentioned residues in Csk-SH2 maintains the active form, stabilizes packaging between the hydrophobic pocket, including Tyr-116 of the SH2 domain and Ala-228 in the catalytic domain, and consequently leads to allosteric activation of Csk (Figs. 10B and 9C).

In conclusion, we determined the solution structure of the complex of Csk-SH2 with Cbp. This structure revealed the presence of a novel binding region in Cbp that is separate from the canonical binding region. Furthermore, mutational analysis in cells showed that both canonical and novel binding sites are required for tumor suppression via the Cbp-Csk interaction. These findings indicate that conventionally known interaction sites in target ligands do not explain the biological functions of the associated SH2 domains alone.

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