Solution Structure of the Phosphotyrosine Binding (PTB) Domain of Human Tensin2 Protein in Complex with Deleted in Liver Cancer 1 (DLC1) Peptide Reveals a Novel Peptide Binding Mode

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Background: DLC1 interacts with tensin2 PTB, playing a role as a tumor suppressor in many human cancers.

Results: We solved the tensin2 PTB-DLC1 complex structure and showed its importance for co-localization of DLC1 and tensin2 in cells.

Conclusion: The novel PTB-peptide binding mode provides a molecular basis for understanding the tumor suppression of DLC1 and tensin2.

Significance: A novel PTB-peptide binding mode was observed.

The protein deleted in liver cancer 1 (DLC1) interacts with the tensin family of focal adhesion proteins to play a role as a tumor suppressor in a wide spectrum of human cancers. This interaction has been proven to be crucial to the oncogenic inhibitory capacity and focal adhesion localization of DLC1. The phosphotyrosine binding (PTB) domain of tensin2 predominantly interacts with a novel site on DLC1, not the canonical NPXY motif. In this study, we characterized this interaction biochemically and determined the complex structure of tensin2 PTB domain with DLC1 peptide by NMR spectroscopy. Our HADDOCK-derived complex structure model elucidates the molecular mechanism by which tensin2 PTB domain recognizes DLC1 peptide and reveals a PTB-peptide binding mode that is unique in that peptide occupies the binding site opposite to the canonical NPXY motif interaction site with the peptide utilizing a non-canonical binding motif to bind in an extended conformation and that the N-terminal helix, which is unique to some Shc- and Dab-like PTB domains, is required for binding. Mutations of crucial residues defined for the PTB-DLC1 interaction affected the co-localization of DLC1 and tensin2 in cells and abolished DLC1-mediated growth suppression of hepatocellular carcinoma cells. This tensin2 PTB-DLC1 peptide complex with a novel binding mode extends the versatile binding repertoire of the PTB domains in mediating diverse cellular signaling pathways as well as provides a molecular and structural basis for better understanding the tumor-suppressive activity of DLC1 and tensin2.

Deleted in liver cancer 1 (DLC1) is a tumor suppressor gene located on chromosome 8p21.3–22 first identified in primary human hepatocellular carcinoma (HCC) (1). The DLC1 protein contains an N-terminal sterile α motif domain, a GTPase activation protein domain for Rho family GTPase (RhoGAP), a C-terminal steroidogenic acute regulatory protein-related lipid transfer domain, and a serine-rich unstructured middle region. Loss or down-regulation of DLC1 expression mediated by genetic and epigenetic mechanisms has been associated with the development of many human cancers, including HCC; breast, lung, colon, and prostate cancers; and nasopharyngeal, esophageal, and cervical carcinomas (2–4). Restoration of DLC1 expression was found to suppress cell migration and invasion, inhibit growth, and induce apoptosis of various cancer cell line models in vivo and in vitro (5–8). Functional data regarding the loss of DLC1 in development of hepatocellular carcinoma cells have been shown in a mouse model (9). Accumulating evidence has well demonstrated that DLC1 is a bona fide tumor suppressor in diverse human malignancies. Different mechanisms regulate the tumor-suppressive function of DLC1 (10–12).

The tumor-suppressive function of DLC1 has been shown to be regulated by its association with tensins (13). The tensins are
a family of proteins localized at integrin-mediated focal adhesions bridging the actin cytoskeleton and integrins (14). To date, four members, namely tensin1, tensin2/C1-TEN, tensin3, and tensin4/C-terminal tensin-like (cten), with structural similarity and high sequence homology have been identified (15, 16). All tensins contain C-terminal Src homology 2 and phosphotyrosine binding (PTB) domains, which confer interacting potentials with diverse scaffolding proteins and signaling molecules at focal adhesions (17–20). Identification of tensin2 as the novel interacting partner of DLC1 by our group has provided evidence about the focal adhesion localization of DLC1 (21, 22). Subsequent studies have further demonstrated the functional implication of the interactions between DLC and other tensin proteins in tumor suppression by DLC1 (13, 23). To date, association with tensin proteins is the best characterized regulatory event in the subcellular localization of DLC1 at the protein level. Different binding regions have been identified in DLC1 that interact with Src homology 2 (24) and/or PTB domains of tensins, and they localize to the focal adhesion-targeting region within amino acids 201–500 of DLC1 (25). More detailed biochemical analysis showed that DLC1 utilizes residues 375–385 to predominantly interact with the tensin2 PTB domain, and its growth inhibitory activity depends on this PTB domain interaction (26).

Overall, as a popular protein binding module, a large number of proteins contain the PTB domain, which exhibits a conserved core structure of a PH domain superfold containing a β-sandwich made of two antiparallel β-sheets capped by a C-terminal helix. Proteins containing PTB domains function as adaptors or scaffolds to organize signaling complexes involved in a wide range of physiological processes. The structure of PTB domains confers specificity for binding peptides having a NPXY motif via a structurally conserved mechanism with different requirements for the phosphorylation of tyrosine within this recognition sequence. Structurally, the C-terminal α helix and β5 strand of the PTB domain are involved in this interaction (27). Intriguingly, DLC1 does not contain an NPXY-like motif, indicating it may interact with the PTB domain of tensin2 through a different mechanism.

Despite the differential roles of tensins in different cellular contexts, association with DLC1 plays an unequivocal role in modulating the tumor-suppressive activity of DLC1. As mentioned, DLC1 predominantly interacts with the PTB domain of tensin2. To gain insight into the molecular mechanism of DLC1-tensin2-mediated tumor suppression, it is important to understand the binding mode between DLC1 and the PTB domain of tensin2. In this study, we characterized and verified the binding surface and key residues on the tensin2 PTB domain that are involved in the interaction with DLC1 peptide. We also determined the solution structure of tensin2 PTB in complex with the DLC1 peptide by NMR. The PTB domain of tensin2 binds to a non-phosphorylated peptide of DLC1 in a manner not reported before. The HADDOCK-derived complex structural model reported here reveals a novel peptide binding mode and expands the recognition modes of PTB domains beyond the canonical NPX(p)Y-like motif (where pY is phosphotyrosine) in mediating diverse cellular signaling pathways.

**EXPERIMENTAL PROCEDURES**

**Plasmid Construction**—The PCR-amplified cDNA fragment encoding residues 1262–1409 of PTB domain of human tensin2 (GI 38787956) was cloned into the pET vector (Novagen) by standard molecular cloning techniques with a His6 tag-thrombin cleavage sequence at the N terminus. The coding sequence of DLC1 peptide, corresponding to residues 374–388 of human DLC1 (GI 33188437) with an additional tryptophan residue at the C terminus for protein quantification, was PCR-amplified, purified using a QIAEX II gel extraction kit, and cloned into a PET-derived expression vector. The resulting construct was tagged with a 56-residue B1 immunoglobulin binding domain of streptococcal protein (GB1) and a protein 3C protease cleavage site at the N terminus. For *in vitro* biochemical analysis, the DLC1 peptide was constructed as a GST fusion protein using pGEX-4T-1 vector (GE Healthcare).

**Site-directed Mutagenesis**—Alanine point mutations and deletions in the PTB domain were performed by QuikChange site-directed mutagenesis using *Pfu* DNA polymerase (Sangon Biotech Shanghai) with suitable primer sets. The presence of appropriate mutations was confirmed by DNA sequencing. For immunofluorescence imaging and the colony formation assay, GFP-tagged full-length tensin2 and its mutants were constructed on pEGFP1. The number in the name of full-length tensin2 alanine mutants is the residue’s sequential number in our PTB complex structure. The “Δα” mutant refers to the full-length tensin2 mutant deleted of the N-terminal α1 helix (residues 15–25 of the PTB domain).

**Protein Expression and Purification**—The recombinant proteins were expressed in *Escherichia coli* strain BL21(DE3) at 18 °C overnight. For PTB domain purification, the proteins were purified from the bacterial lysates by sequential chromatography on nickel-nitrioltriacetic acid and Superdex 75 columns (GE Healthcare). For DLC1 peptide purification, proteins were expressed and purified with nickel-nitrioltriacetic acid resin as described above. The N-terminal GB1 tag was cleaved by protein 3C protease and separated by gel filtration. The peak fraction containing DLC1 peptide was desalted in H2O using a HiTrap Desalting column (GE Healthcare) and lyophilized for further study. The molecular weight of purified DLC1 peptide was confirmed by mass spectrometry. For the NMR studies, uniformly isotope-labeled proteins were expressed and purified as described above except that the bacteria were grown in M9 minimal medium using 15NH4Cl and/or [13C]glucose (Cambridge Isotope Laboratories Inc.) as the sole nitrogen and/or carbon sources. The NMR samples were concentrated to ~0.2 (for HSQC-based titration experiments) or ~0.8 mM (for structural determinations) in optimized NMR sample buffer (50 mM sodium phosphate, 100 mM NaCl, 1 mM EDTA, pH 7.0).

**GST Pulldown Assay**—For coupling, prewashed glutathione-Sepharose 4B slurry beads were incubated with bacterial lysates of GST or GST-DLC1(374–392) in an assay buffer (50 mM sodium phosphate, 150 mM NaCl, 1 mM EDTA, 0.1% Nonidet P-40, pH 7.0) at 4 °C for 1 h and washed three times. Then the beads were incubated with bacterial lysates of wild-type or mutated PTB domains for another 2 h at 4 °C and washed three times. The captured proteins were eluted in 50 mM
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Tris, pH 7.9 containing 10 mM glutathione. The samples were then resolved by 15% SDS-PAGE and subjected to Western blot analysis with an anti-His tag antibody (HisProbe, H-3, Santa Cruz Biotechnology).

NMR Binding Studies—To investigate the ligand binding, the two-dimensional 1H-15N HSQC spectra were recorded on uniformly 15N-labeled PTB domain (~0.2 mM) in the presence of different concentrations of DLC1 peptide ranging from 0 to 1.5 mM. Both the PTB sample and the stock solutions of DLC1 were prepared in the NMR buffer (50 mM sodium phosphate, 100 mM NaCl, 1 mM EDTA, pH 7.0). The chemical shift perturbation between the free form and DLC1-bound PTB domain was normalized by the following formula and expressed in ppm.

\[
\text{Chemical shift perturbation} = \sqrt{(\Delta \delta H)^2 + (\Delta \delta N \times \alpha N)^2}
\]

(Eq. 1)

In the equation, \(\Delta \delta H\) and \(\Delta \delta N\) are the differences in chemical shifts of amide protons and nitrogen between the initial and final data points of the titration, respectively. The nitrogen frequency is normalized for proton with the scale factor \(\alpha N = 0.17\), established from estimates of atom-specific chemical shift ranges in a protein environment, in the equation (28).

Fluorescence Polarization Assay—The fluorescence polarization assay was performed on a PerkinElmer Life Sciences LS-55 fluorometer equipped with an automated polarizer at 20 °C. Fluorescence titration was performed by adding increasing amounts of purified PTB domains to a fixed amount of fluorescein 5-isothiocyanate (FITC; Molecular Probes)-labeled DLC1 peptide in 50 mM sodium phosphate, 100 mM NaCl, 1 mM EDTA, pH 7.0. The polarization value of the FITC-labeled DLC1 peptide was measured at each titration point. The \(K_d\) values were obtained by fitting the titration curve with the classical one-site binding model.

NMR Spectroscopy—NMR spectra were acquired at 37 °C on 750- and 500-MHz Varian NMR spectrometers with self-shielded z axis gradients. All spectra were processed using NMRPipe (29, 30) and analyzed using SPARKY 3.6 The 1H, 15N, and 13C resonances of backbone and side chain atoms were assigned by using a standard set of triple resonance experiments on either uniformly 15N, 13C-labeled PTB domain with/without unlabeled DLC1 or uniformly 15N, 13C-labeled DLC1 with/without unlabeled PTB domain at protein concentrations of ~0.8 mM (31–33). The PTB-DLC1 complex was prepared at a ratio of 1:2 between 15N, 13C-labeled and unlabeled components. The initial backbone assignment was generated by Mars (34) and manually checked in combination with 15N-edited three-dimensional NOESY. NOE-derived distance restraints were obtained from 15N- or 13C-edited three-dimensional NOESY spectra, each with a mixing time of 120 ms, complemented by 13C-edited, 15N/13C-filtered three-dimensional NOESY spectra for the intermolecular contact (mixing time, 150 ms) (35). Steady-state heteronuclear [1H]-15N NOE values were determined from spectra recorded with 5-s relaxation and the presence and absence of a proton presaturation period of 3 s in the 500-MHz spectrometer.

NOE Analysis and Structure Calculations—NOE assignment and structure calculations were performed using the CANDID (36) module of the program CYANA2.1 (37). We first calculated the structures of the PTB domain in its free and DLC1-bound forms and vice versa for DLC1 peptide. Next, the initial 200 complex structures of the PTB domain and DLC1 were generated with manually assigned unambiguous intermolecular NOE distance restraints by CNS 1.1 (38). Then the 120 top scoring complex structures from CNS were further refined with additional chemical shift perturbation data using a HADDOCK-type protocol without rigid body docking (39). The calculation details are described in the supplemental Materials and Methods. A cutoff value of 1.4 Å was defined to select the well resolved ensembles of structures. The ten lowest scored structures from each cluster were analyzed further. A family of 10 lowest energy structures in which none showed any restraint violation over 0.5 Å was used for statistical analysis. The dihedral restraints were generated from TALOS (40). The quality of the HADDOCK-derived structural models was assessed using PROCHECK (41) and analyzed by MOLMOL (42). All of the figures representing the structures were generated by PyMOL.

Paramagnetic Labeling—To produce spin-labeled DLC1 peptides, a single cysteine was added to the N- or C-terminal ends, respectively. After purification, the protein was exchanged with phosphate buffer with no reducing agent. A 5-fold molar excess of spin labeling reagent MTSL was added, and the reaction proceeded at 25 °C for 12 h (Toronto Research Chemicals). Excess reagent was removed with a desalting column. A series of 1H-15N HSQC spectra were obtained to monitor the disappearance of a subset of resonances of 15N-labeled tensin2 PTB in the presence of increasing molar ratios of unlabeled, MTSL-coupled GB1-DLC1. A control spectrum was collected at the end of the titration after the addition of 10 mM ascorbic acid to remove the spin label MTSL.

Immunofluorescence Microscopy—To reveal the co-localization of tensin2 and DLC1 in cells, HEK293T cells seeded on coverslips were transfected with Myc-tagged DLC1 full length (FL) and GFP-tagged tensin2 FL or mutants. Transfected cells were fixed with 4% paraformaldehyde, PBS and permeabilized with 0.5% Triton X-100, PBS. Cells were then blocked with 5% bovine serum albumin. Myc-DLC1 was stained by anti-Myc antibody followed by rhodamine-conjugated secondary antibody. Cells were mounted in Vectashield antifade mountant (Vector Laboratories, Burlingame, CA). Images were captured by a Leica Q550CW fluorescence microscope (Leica, Wetzler, Germany).

Colony Formation Assay—The DLC1-deficient human HCC cell line BEL7402 was obtained from Shanghai Institute of Cell Biology, Chinese Academy of Sciences. BEL7402 has been used for the functional characterization of DLC1 and tensin2 elsewhere (5, 21). Cells at 2.5 × 10^4/well were seeded into 12-well tissue culture plates 2 days before transfection. Each of 0.25 μg of GFP-tagged tensin2 and its mutant expression vector (wild-type tensin2, Δα, L139A, D140A, F151A, and ΔPTB) was co-transfected with 1 μg of DLC1-pCS2+MT into cells. One day after transfection, cells were trypsinized and replated at a 1:20 dilution delay and the presence of a proton presaturation period of 3 s in the 500-MHz spectrometer.

6 T. D. Goddard and D. G. Kneller, University of California, San Francisco, unpublished software.
dilution in triplicates into 6-well tissue culture plates. Cells were selected in 700 μg/ml G418 (Merck) for 3 weeks. Colonies formed were fixed with 3.7% formaldehyde and stained with crystal violet solution.

RESULTS

Binding Studies of Tensin2 PTB Domain and the DLC1 Peptide—In a previous study, we identified that amino acids 375–385 of DLC1 were responsible for binding the PTB domain of tensin2 (26) (Fig. 1A). Before the structural study, we first performed an in vitro pulldown assay to verify the binding. The GST-tagged peptide (EDHKPGTFPKALTNG) comprising DLC1 amino acids 374–388 was able to pull down tensin2 PTB domain from bacterial lysate (Fig. 1B), confirming their direct physical interaction. Subsequently, the DLC1 peptide was labeled with FITC to examine its affinity for tensin2 PTB domain in a fluorescence polarization study. As shown in Fig. 1C, the FITC-labeled DLC1 peptide was titrated with increasing amounts of unlabeled PTB, and the polarization value of the FITC-labeled DLC1 peptide was measured at each titration point. A, a section of the overlaid 1H-15N HSQC spectra of 15N-labeled PTB in free form (black) and titrated with non-labeled DLC1 peptide at different molar ratios, 1:1 (blue), 1:2 (magenta), 1:3 (green), 1:4 (yellow), 1:5 (maroon), and 1:6 (red). Residues that undergo significant changes in chemical shifts upon formation of the complex with DLC1 are highlighted by arrows and labeled with peak assignments. E, the weighted chemical shift changes between the free form and DLC1 peptide-saturated PTB obtained from NMR titration experiments are plotted as a function of residue number.

The DLC1 peptide binding properties were further investigated by NMR titration using chemical shift perturbation analyses. A 15N-labeled tensin2 PTB domain was titrated with non-labeled GB1-tagged DLC1 peptide (amino acids 374–388). The residues with chemical shift changes were monitored by two-dimensional 1H-15N HSQC spectra. As shown in Fig. 1D and supplemental Fig. 4, the addition of DLC1 peptide induced significant chemical shift perturbations and resulted in a step-by-step peak shift pattern at many positions in the spectra, indicating a fast exchange regime on the NMR time scale. The 1H-15N HSQC spectra of tensin2 PTB domain showed no change when titrated with GB1 protein alone (data not shown). As shown in Fig. 1E, the weighted chemical shift changes of the amide 1H and 15N obtained from NMR titration experiments were plotted as a function of the residue number. The signals having large chemical shift changes (Δδ > 0.038 ppm) corresponded to the region formed by the helices α1 and α3; β strands β1, β2, and β3; and the loop connecting β7 to the helix α3. The titration results entail the discovery of a new binding region different from the well known PTB substrate binding site reviewed by Uhlik et al. (27).
Overall Structure of the Complex—As our binding studies indicated a new binding mode in which DLC1 peptide occupies the non-canonical binding site of PTB domain, we went further to fully characterize this interaction at the atomic level by multidimensional heteronuclear NMR spectroscopy as described under “Experimental Procedures.” Combining the NMR and mutagenesis data, we determined a HADDOCK-derived structural model for the tensin2 PTB and DLC1 peptide complex. A superimposed ensemble of 10 structures and the ribbon plot of the lowest energy structure are shown in Fig. 2A. The structure ensemble had good backbone geometry, no significant restraint violation, and low pairwise root mean square deviation values (Table 1). The structure of tensin2 PTB domain in the complex presents a typical PH domain superfold, a seven-stranded β-sandwich composed of two antiparallel β-sheets capped by a C-terminal α-helix, α3. Moreover, it contains two additional α-helices, N-terminal α1 and α2, connected to strands β1 and β2, respectively. In the complex, DLC1 peptide is unstructured and interacts with the surface of PTB domain formed by N-terminal helix α1; β-strands β1, β2, and β3; and the loop connecting β7 to C-terminal helix α3, consistent with the titration data (Fig. 2A). Intermolecular NOEs defined the orientation of the DLC1 peptide in the complex (supplemental Fig. 1), which was later independently confirmed by paramagnetic relaxation enhancement experiments as shown in Fig. 3. The backbone amide resonances of Asn-80 and Gln-143 disappear in the presence of the N-terminal spin-labeled DLC1 peptide (Fig. 3A, red), whereas they remain when titrated by C-terminal spin-labeled DLC1 peptide (Fig. 3B, red). This result indicates that residues Asn-80 and Gln-143 are closer to the N terminus of DLC1 peptide than to the C terminus (Fig. 3C). Ascorbic acid can reduce MTSL spin label. The cross-peaks of Asn-80 and Gln-143 recovered after addition of ascorbic acid, indicating that the intensity loss of the two peaks was caused specifically by the paramagnetic spin label (Fig. 3A, black). The paramagnetic relaxation enhancement experiments independently confirmed the orientation of the DLC1 peptide.

Interactions at the Tensin2 PTB Domain and DLC1 Peptide Interface—The binding interface of the tensin2 PTB-DLC1 complex comprises the strands β1–β3, helices α1 and α3, and the loop between β7 and helix α3 (Fig. 2B). The binding site accommodates nearly all the residues of DLC1 peptide in an extended conformation. The DLC1 binding site on PTB domain is characterized by a hydrophobic surface formed by Leu-21, Leu-22, Gln-24, Ala-27, Ser-71, Ala-72, Gln-73, and Thr-76 that helps to occupy the Pro-Gly-Thr-Phe-Pro segment of DLC1. Additionally, the aromatic side chain of DLC1 Phe-381 fills a shallow pocket formed by Lys-69, Val-70, Ser-71, Gln-73, Gly-74, and Thr-76 of PTB domain and is nearly perpendicular to the β2 and β3 strands. Another key residue of DLC1 at the binding interface is His-376. The imidazole ring of His-376 is stabilized by stacking against the side chains of Cys-28, Ser-29, Leu-139, and Pro-141. The hydrophobic residues Ala-384 and Leu-385 of DLC1 are involved in the complex for-
mation, making van der Waals contacts with a narrow cleft formed by the side chains of Arg-23 and Gln-73 of PTB domain. 

**Structure-based Mutational Studies of the Interaction between Tensin2 PTB and DLC1 Peptide**—Based on the complex structure and the chemical shift perturbations observed, a series of point mutants in PTB domain were constructed to evaluate the contributions of these residues to the tensin2 PTB-DLC1 interaction. These point mutants (D20A, L21A, R23A, Q24A, C28A, S29A, V30A, H67A, F68A, K69A, V70A, S71A, R82A, S99A, L139A, D140A, F151A, K154A, and V155A) were expressed and purified to homogeneity. The results of a GST pulldown assay (Fig. 4, B and C) indicate that mutations in the DLC1 binding region of tensin2 affected with tensin2 PTB mutants (Fig. 5, A and B) showed little change from the whole PTB domain, indicating that the core structure was not perturbed. No chemical shift changes could be seen in PTB Δα when titrated with DLC1 peptide. These results indicate that the novel binding mode would only apply to Shc- or Dab-like PTB domains that contain an N-terminal helix.

**Effects of PTB Mutants on the Co-localization of Tensin2 and DLC1 in Cells**—To further confirm the importance of the defined residues in the tensin2-DLC1 interaction, we examined their effect on the co-localization of tensin2 and DLC1 in cells. GFP-tagged wild-type or alanine substitution mutants of full-length tensin2 (GFP-tensin2 FL) were co-transfected with Myc-tagged full-length DLC1 (Myc-DLC1 FL) into HEK293T cells, and the proteins were visualized by immunofluorescence microscopy. As shown in Fig. 5A, GFP-tagged full-length tensin2 co-localized well with Myc-DLC1 FL, whereas the GFP-tensin2 FL mutant (PTB L139A) did so less frequently. In addition, GFP-tensin2 FL mutants (PTB D140A and PTB F151A) failed to co-localize with Myc-DLC1 FL. Similar results were obtained in the DLC1-deficient human BEL7402 HCC cells (data not shown), indicating the role of Leu-139, Asp-140, and Phe-151 in the PTB domain in formation of the tensin2-DLC1 complex.

In a previous colony formation assay, the DLC1 mutant of which the PTB binding region was deleted partially lost the growth-suppressive activity (26). To further address the requirement of tensin2 PTB/DLC1 binding for the tumor-suppressive function of DLC1, we carried out the same colony formation assay using tensin2 mutants derived from our structural and mutational studies in DLC1-deficient BEL7402 HCC cells. The co-transfection of tensin2 with DLC1 significantly suppressed colony formation (p = 0.009) when compared with the transfection of tensin2 only (data not shown). In contrast, growth suppression was abolished when DLC1 was co-transfected with tensin2 PTB mutants (Fig. 5, B and C). These results indicate that mutations in the DLC1 binding region of tensin2 PTB domain reduced the growth suppression activity of tensin2 and DLC1 probably due to the loss of efficient tensin2-DLC1 interaction and that our PTB-peptide complex structure provides a molecular and structural basis for the tumor-suppressive activity of DLC1 and tensin2.

**DISCUSSION**

The tumor suppressor DLC1 utilized a novel binding site for tensin2 PTB domain interaction, and this binding is required for the tumor-suppressive function of DLC1 (26). To obtain structural insight into the molecular mechanism of the above mentioned binding, we performed binding assays with tensin2 PTB and the DLC1 peptide using GST pulldown, fluorescence polarization, and NMR chemical shift perturbation. We also Tested the role of N-terminal helix α1 (residues 15–25 of PTB domain) in the binding. Deletion of the helix α1, which is only seen in Shc- and Dab-like PTB domains, abolished binding of the DLC1 peptide (Fig. 4B). Examination of a two-dimensional 1H-15N HSQC spectrum of PTB Δα showed little change from the whole PTB domain, indicating that the core structure was not perturbed. No chemical shift changes could be seen in PTB Δα when titrated with DLC1 peptide. These results indicate that the novel binding mode would only apply to Shc- or Dab-like PTB domains that contain an N-terminal helix.

**TABLE 1**

| Statistics for structures | r.m.s. value (Å) |
|---------------------------|------------------|
| Final energies (kcal/mol) | 621.89 ± 34.79   |
| van der Waals (kcal/mol)  | 0.9765 ± 0.2475  |

| Violations                  | r.m.s. deviation (Å) |
|----------------------------|----------------------|
| Number of NOE violations   | 0 ± 0                |
| Total NOE distance restraints | 0.0229 ± 0.0028      |
| Number of dihedral angle constraint violations | 0 ± 0 |
| Total NOE violations       | 0.5547 ± 0.0607      |

| Deviations from idealized geometry | r.m.s. deviation (°) |
|-----------------------------------|----------------------|
| Bonds (%)                         | 0.0037 ± 0.0001      |
| Angles (%)                        | 0.4928 ± 0.0294      |
| Improper (%)                      | 0.4766 ± 0.0383      |

| Structural r.m.s.d. to the mean coordinate, bb/heavy (Å) | r.m.s.d. (Å) |
|----------------------------------------------------------|-------------|
| Regions (residue numbers) 15–161, 367–380               | 1.15/1.73   |
| Regions (residue numbers) 20–23, 27–37, 43–56, 64–71, 74–78, 90–91, 95–99, 106–108, 114–124, 127–137, 145–155, 370–377 | 0.80/1.23 |

| Ramachandran plot (°) | r.m.s. deviation (°) |
|----------------------|----------------------|
| Residues in most favored regions | 80.5 |
| Residues in additional allowed regions | 17.1 |
| Residues in generously allowed regions | 1.4 |
| Residues in disallowed regions | 1 |

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determined the solution structure of tensin2 PTB domain and provided a HADDOCK-derived structural model for the ten-
sin2 PTB-DLC1 peptide complex by magnetic resonance spec-
troscopy (NMR), which was the first complex structure model for tensin2 PTB domain. In the complex structure, the binding
features revealed a novel peptide binding mode that has not been observed in PTB-peptide complexes. Our complex structural model is consistent with the binding study, mutagenesis, and cell assay data. However, a future co-crystal structure would be more precise than the present HADDOCK-derived structural model.

Comparison with Representative PTB-Peptide Complexes—As adapter or scaffold proteins, PTB domain-containing proteins play a critical role in regulating the spatial and temporal organization of signaling networks in a wide range of physiological and pathological processes. The structure and function of PTB-peptide complexes have always been a research focus. Most of the complexes reported share a general mode of peptide binding, a conserved location of the peptide binding pocket and a consensus binding motif. In this canonical binding mode, the PTB peptide ligands are bound as an antiparallel pseudo-
H-sheet, forming contacts with the C-terminal helix (Fig. 6, pink region), and the consensus NPXY motifs form a type I β-turn toward their C termini. Although this canonical peptide binding mode does not depend on the phos-
phorylation state of the tyrosine in the peptide, the phosphorylated tyrosine may greatly increase the binding affinity. In light of the structure/function relationships, PTB domains are divided into Shc-like, IRS-like, and Dab-like subgroups (27). The canonical peptide binding mode is seen in the representative complexes structures Shc and TrkA complex (Fig. 6B), IRS-1 PTB and IL-4 peptide complex (Fig. 6C), and Dab1 PTB

FIGURE 3. DLC1 peptide orientation determined by paramagnetic relaxation enhancement experiments. A cysteine residue was added to the N and C termini of the DLC1 peptide by site-directed mutagenesis. The paramagnetic spin label MTSL was introduced to the cysteine residue through the formation of a thioester bond. Sections of 1H-15N HSQC spectra of PTB domain are shown here. A, a section of the overlaid 1H-15N HSQC spectra of 15N-labeled PTB in the presence of the N-terminal spin-labeled GB1-tagged DLC1 peptide (red) and after addition of 10 mM ascorbic acid (black). B, a section of the overlaid 1H-15N HSQC spectra of 15N-labeled PTB in the presence of the C-terminal spin-labeled DLC1 peptide (red) and after addition of 10 mM ascorbic acid (black). Note that the backbone amide resonances of Asn-80 and Gln-143 disappear in the presence of the N-terminal spin-labeled GB1-tagged DLC1 peptide. The paramagnetic relaxation enhancement experiments independently confirmed the orientation of the DLC1 peptide. C, ribbon diagrams of the complex showing the position of the residues Asn-80 and Gln-143, which are in closer proximity to the N terminus of DLC1 peptide than to the C terminus. The backbone of PTB is colored gray. DLC1 is indicated as line mode and colored red. The residues Asn-80 and Gln-143 are shown in stick mode and colored blue and pink, respectively. The middle and right panels are the same as the left panel after a 90° and 180° rotation, respectively, about the z axis.

FIGURE 4. Mutational studies of the interaction between tensin2 PTB and DLC1 peptide in vitro. A, bacterial lysates of wild type or a series of PTB mutants were incubated with immobilized GST or GST-DLC1 peptide. The associated proteins were analyzed by immunoblotting (IB) with anti-His, tag antibody (upper panel). The middle and lower panels indicate a relatively equal amount of GST/GST-DLC1 peptide (DLCpep) and bacterial lysates of PTB mutants used in the GST pulldown (PD) assay. Gluta, glutathione. B, the effect of N-terminal helix of PTB on DLC1 peptide interaction. The wild type (wt) or PTB Δα were pulled down by GST/GST-DLC1 peptide from bacterial lysates and visualized by anti-His antibody (upper panel). The lower panel indicates a relatively equal amount of GST/GST-DLC1 peptide used in the GST pulldown assay.
and ApoER2 complex (43) (Fig. 6E). On the other hand, some peptides bind to the canonical site on the PTB with small differences. As shown in Fig. 6D, which shows the SNT-1 PTB domain in complex with peptide derived from fibroblast growth factor receptor 1, the C-terminal region of the peptide forms antiparallel β-sheets with a unique C-terminal β8 strand and the β5 strand, whereas the N-terminal region of FGFR1 peptide wraps around the β-sandwich structure of the PTB domain (44). Another example is the Numb PTB domain in complex with GPpY-containing peptide (Fig. 6F). The peptide is in a helical turn conformation that does not hydrogen bond to the PTB domain β-sheet (45).

**Novel Peptide Binding Mode of PTB Domain**—Our NMR solution structure provides the first view of the tensin2 PTB domain in complex with its peptide ligand in a non-canonical fashion that has not been observed before. First, the DLC1 peptide occupies a novel binding site on tensin2 PTB domain. The complete solution structure of the tensin2 PTB domain with the binding peptide from DLC1 revealed that the binding cleft is formed by the N-terminal helix α1; β-strands β1, β2, and β3; and the loop connecting β7 to C-terminal helix α3. This region is on the opposite side of the PTB domain from the conserved peptide binding location formed by the β5 strand and the C-terminal α-helix in the canonical binding mode. Second, DLC1 peptide utilizes a non-canonical binding motif for the tensin2 PTB domain interaction. DLC1 peptide does not have the consensus NPXY motif. The binding site on PTB accommodates nearly all the residues of DLC1 peptide in an extended conformation.

In addition, the N-terminal helix α1 is required for this binding, whereas it is either misfolded or not present in many PTB structures obtained. Deletion of helix α1 (amino acids 15–24) prevented pulldown of the PTB domain by the DLC1 peptide, and no chemical shift perturbations indicating binding could be observed in an NMR titration experiment although this deletion did not make noticeable structural changes to the PTB.
domain as judged by the two-dimensional $^1$H-$^{15}$N HSQC spectrum of PTB $\Delta \alpha$. Comparison of the tensin2 PTB domain in complex with the DLC1 peptide and other tensin PTB domain structures (Protein Data Bank codes 2GJY (46), 1WVH (47), 2DKQ, and 3HQC) showed that binding of DLC1 does not substantially alter the structure (see supplemental Fig. 3). However, there is a difference in the orientation of helix $\alpha_1$ in the tensin2 PTB complex structures. In the x-ray crystal structure of tensin2 PTB (Protein Data Bank code 3HQC), this helix protrudes out from the core of the molecule, whereas in the NMR structure (Protein Data Bank code 2DKQ), there is a difference of about 60° in the relative orientation of helices $\alpha_1$ and $\alpha_3$ from that in the bound form. In the complex structure, helix $\alpha_1$ makes close contacts with the C-terminal helix $\alpha_3$ as a network of NOEs between the residues in these helices (Leu-21 to Phe-151 and Leu-22 to Val-155) was observed in extensive analysis of NOESY spectra. This is consistent with the result of the mutational study that the F151A mutant lost binding with no change to the overall structure fold. The loss of binding may be due to the release of helix $\alpha_1$ from $\alpha_3$ caused by Phe-151 mutation. The requirement of the N-terminal helix $\alpha_1$ is a new feature of PTB-peptide binding found in the tensin2 PTB-DLC1 interaction. This indicates that the binding mode shown by DLC1 may only apply to Shc- or Dab-like PTB domains, which have this N-terminal helix.

**Biological Implications**—Although the role of DLC1 as a bona fide tumor suppressor has become clearer recently, its molecular mechanism of biological regulation remains unclear (48). Tensin proteins are the first identified interacting partners of the DLC family. Interaction with tensin proteins not only determines the subcellular localization of DLC1 but more importantly influences the biological activities of both DLC1 and the tensin family. These reported studies have provided compelling evidence about the impact of the regulatory mechanism of DLC1 at the protein level with its interaction with tensin family proteins (13, 21, 23, 49). We have previously documented the interactions between DLC1 and the tensin2 PTB domain and identified the binding site on DLC1. In this study, we further identified the important binding residues in tensin2 PTB domain by structural analysis. Residues Leu-139, Asp-140, and Phe-151 are critical in mediating the association with DLC1 peptide comprising amino acids 374–388. Mutations of these residues in FL tensin2 affected the formation of the DLC1-tensin2 binding complex and further reduced the growth-suppressive activity of the DLC1-tensin2 complex. Our findings have provided evidence that proper co-localization of tensin2-DLC1 through PTB-peptide interaction and their biological activities are tightly associated. These findings substantiate the biological regulation of the tumor suppressor DLC1 by the tensin family and provide a better understanding of DLC1-tensin2 PTB domain interaction at the structural and molecular levels.

In conclusion, we determined the NMR solution structure of tensin2 PTB in complex with the DLC1 peptide. Our HADDOCK-derived complex structural model and binding studies have elucidated the molecular mechanism by which tensin2 PTB domain recognizes DLC1 peptide and revealed a novel PTB-peptide binding mode not reported before that extends the versatile recognition modes of the PTB domains in mediating diverse cellular signaling pathways. This PTB-peptide bind-
ing mode is unique in that peptide occupies a novel binding site separate from the canonical NPXY motif interaction site and that the N-terminal helix of PTB is required for binding. Meanwhile, this PTB-peptide complex structure defines the residues essential for the formation of the tensin2 and DLC1 complex and provides a molecular and structural basis for better understanding the tumor-suppressive activity of DLC1 and tensin2.

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