Phosphorylation of a PDZ Domain Extension Modulates Binding Affinity and Interdomain Interactions in Postsynaptic Density-95 (PSD-95) Protein, a Membrane-associated Guanylate Kinase (MAGUK)*§

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Postsynaptic density-95 is a multidomain scaffolding protein that recruits glutamate receptors to postsynaptic sites and facilitates signal processing and connection to the cytoskeleton. It is the leading member of the membrane-associated guanylate kinase family of proteins, which are defined by the PSD-95/Discs large/ZO-1 (PDZ)-Src homology 3 (SH3)-guanylate kinase domain sequence. We used NMR to show that phosphorylation of conserved tyrosine 397, which occurs in vivo and is located in an atypical helical extension (α3), initiates a rapid equilibrium of docked and undocked conformations. Undocking reduced ligand binding affinity allosterically and weakened the interaction of PDZ3 with SH3 even though these domains are separated by a ~25-residue linker. Additional phosphorylation at two linker sites further disrupted the interaction, implicating α3 and the linker in tuning interdomain communication. These experiments revealed a novel mode of regulation by a detachable PDZ element and offer a first glimpse at the dynamic interaction of PDZ and SH3-guanylate kinase domains in membrane-associated guanylate kinases.

Phosphorylation is one of the most common modifications made to proteins. Depending on the protein substrate, it can regulate the gain or loss of activity through a variety of mechanisms. Most mechanisms studied to date appear to work via a change in steric geometry at the active site or a global conformational change in the protein (1–3), although a few recent examples indicate modulation of protein dynamics that can yield a graded response (4, 5). In some cases, phosphorylation simply provides a protein-protein recognition site (6). Overall, the extent of structural characterization of phosphorylation mechanisms is limited given the ubiquity of this post-translational signaling mechanism and the large diversity of effects observed upon phosphorylation. A greater understanding of how phosphorylation(s) modulates protein activity is essential for a mechanistic understanding of signal transduction.

Many phosphorylated proteins are multidomain, modular proteins (2–4). One class of such proteins that shape signal transduction is the scaffolding proteins, which are frequently phosphorylated (7). Membrane-associated guanylate kinase (MAGUK)§ family proteins are multidomain scaffolding proteins that play roles in ion channel clustering, intracellular trafficking, signal transduction, and cell-cell adhesion (8–10). Malfunction of MAGUKs is associated with central nervous system disorders (11). The defining MAGUK signature is the PDZ-SH3-guanylate kinase "supradomain" architecture. The best known MAGUK protein is postsynaptic density-95 (PSD-95), which is found on the cytoplasmic side of postsynaptic terminals. PSD-95 is a non-catalytic scaffolding protein that clusters glutamate receptors and assembles macromolecular complexes for signal integration in the postsynaptic excitatory neurons (8, 12). It consists of three PDZ domains, an SH3 domain, and a non-catalytic guanylate kinase (GK) domain (see Fig. 1A). PDZ domains are ~90-residue, globular, protein interaction domains that have a conserved fold comprised of two α-helices and six β-strands. The function of PDZs is typically to bind the C-terminal 4–5 residues of target proteins, which bind in a groove between α2 and β2 of the PDZ. PDZ domains are often found in tandem copies and are common to proteins that function as scaffolds for assembling signaling complexes or trafficking components at cell-cell junctions. In PSD-95, the three PDZ domains bind many proteins ranging from NMDA and adrenergic receptors to neuronal nitric-oxide synthase, Pyk2, synaptic Ras GTPase activating protein, and microtubule assemblies (13). Structural work has been carried out on individual PDZs of PSD-95 (14, 15) as well as the SH3-GK module (16, 17). However, little is known about how MAGUK PDZ domains associate/function with the SH3-GK module. A role for PDZs beyond their normal C-terminal ligand binding function has

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§ The abbreviations used are: MAGUK, membrane-associated guanylate kinase; §° order parameter characterizing amplitude of amide bond; PDZ, PSD-95/Discs large/ZO-1; PSD, postsynaptic density; SH3, Src homology 3; GK, guanylate kinase; SEA, solvent-exposed amide; HSQC, heteronuclear single quantum correlation; CSP, chemical shift perturbation; ITC, isothermal titration calorimetry.

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emerged from examples in which intermolecular (or interdomain) contacts occur on novel PDZ interfaces (18, 19) or binding is regulated allosterically (18, 20–23).

Previous work suggests that the third PDZ domain (PDZ3) in PSD-95 family proteins regulates activity at SH3-GK, indicating that this PDZ domain may have an additional regulatory function on top of its C-terminal binding function. MAP1a binds to the GK domain of PSD-95; however, this interaction is weakened by the presence of PDZ3 (and abolished by the presence of all three PDZ domains) and restored by PDZ-binding peptides such as adenomatous polyposis coli, NR2B, and cysteine-rich PDZ-binding protein (24). Interdomain interactions were also characterized in the Drosophila homologue of PSD-95, Dlg-1, in which the domain interaction between PDZ3 and SH3 was regulated by PDZ3-binding peptides (25). The region linking PDZ3 and SH3 is vital to the interdomain communication between these two domains: replacement of the conserved linker region with flexible Gly-Ser repeats disrupts the influence that PDZ3 has on the binding of SH3 to its interaction partner, GukHolder (25). Thus, a major question regarding MAGUKs is what is the role of the PDZ domain preceding SH3-GK and how might it participate in intramolecular, interdomain interactions?

Interestingly, PSD-95 is known to be phosphorylated at up to ~12 sites with several of these sites occurring at linker positions in between structured domains as well as one in a C-terminal helical extension to PDZ3 (see Fig. 1A). To date, phosphorylation has received modest attention as a means of regulating PDZ domain function in general. One exception is the PDZ protein NHERF1 for which phosphorylation has been shown to regulate PDZ-mediated autoinhibition (26). Recently, we showed that folding of a C-terminal extension to PDZ3 into an α-helix (third α-helix (α3)) increases binding affinity to PDZ3 ligand via an allosteric mechanism rooted in changes in picosecond-nanosecond side-chain dynamics (21). The α3 helix is not part of the conserved PDZ fold and hence appeared as an unusual auxiliary element whose function had been unclear. Within the α3 helix, Tyr-397 has been shown to be phosphorylated (27), which in principle could either increase or decrease α3 interactions with the PDZ core. In addition to Tyr-397, the nearby residues Ser-415 and Ser-418 can also be phosphorylated (28, 29). This dense collection of phosphorylation sites suggests that this region (C-terminal extension of PDZ3 and the PDZ3-SH3 linker) is a hot spot for MAGUK regulation. Interestingly, structured “extensions” to PDZ domains are now being discovered in more PDZ domains (30).

To probe the role of PDZ3 in the PSD-95 MAGUK and its transition into the linker and SH3 domain, we chose to study by NMR the effect of phosphorylation at three sites in the C-terminal region of PDZ3 and how such phosphorylations modulate PDZ3 structure, function, and its interaction with SH3-GK. A construct of PDZ3 spanning residues 304–402 was phosphorylated at Tyr-397 (p397-PDZ3) and characterized for structural effects. A combination of chemical shift analysis, hydrogen exchange, $^{15}$N relaxation, and isothermal titration calorimetry showed that phosphorylation induces a fast equilibrium between a docked conformation in which α3 is packed against the PDZ core and a conformational state in which α3 is undocked that has reduced affinity for PDZ3 ligands. The latter state was previously identified via truncation of α3 and shown to alter PDZ3 ligand binding affinity through a novel allosteric mechanism driven by side-chain dynamics (21). The findings here support this mechanism by showing that the dynamic allostery can be triggered by a biologically relevant modification and is likely to be utilized in a cellular context. In a second construct of PSD-95 that includes PDZ3-linker-SH3, phosphorylation at all three sites (397, 415, and 418) was shown to increase the mobility of PDZ3 relative to SH3. These studies provide insight into how phosphorylation affects structure and dynamics of an intrinsically flexible, multidomain system. They also show for the first time that the association of PDZ3 with the SH3 domain in PSD-95 is weakened by multiple phosphorylations.

**EXPERIMENTAL PROCEDURES**

**Protein Expression and Purification—PDZ3** (residues 304–402), Y397E-PDZ3, and PDZ3-SH3 tandem (residues 304–532) from rat PSD-95 were subcloned into the pET21a vector with a tobacco etch virus-cleavable N-terminal histidine tag. The PDZ3-SH3 tandem was stabilized by adding the sequence SGSGSPYIWVPAEREEL to its C terminus (16) where bold residues are from residues 713 to 724, which form the F-strand of the SH3 domain. Y397E-PDZ3, PDZ3-SH3 single mutant (Y397E-PDZ3-SH3), and triple mutant (Y397E/S415E/S418E) were made by multiple site mutagenesis PCR (31).Transformed Escherichia coli BL21(DE3) cells were grown at 37 °C in LB medium or minimal medium containing appropriate isotopes. When cell density reached an $A_{600}$ of 0.6, isopropyl 1-thio-β-D-galactopyranoside was added to a final concentration of 1 mM, and the cells were grown for another 16 h at 22 °C. The cells were harvested by centrifugation and stored at −80 °C until use.

To purify all proteins in this study, the cell pellets were resuspended in 50 mM Tris-HCl, pH 8.0, 200 mM NaCl, 0.1% Triton X-100, 25 mM imidazole, and 1 mM lysozyme. The resuspended cells were frozen and thawed for three cycles, and sonication was applied. The centrifuged cell lysate was loaded to a nickel affinity column and eluted with 500 mM imidazole. The fractions containing target proteins were pooled and subjected to overnight tobacco etch virus cleavage at 4 °C. A 100% cleavage was achieved as suggested by SDS-PAGE. Cleaved samples were further purified by Source Q ion exchange chromatography followed by a G50 size exclusion column equilibrated in a buffer containing 22 mM NaPO$_4$, pH 6.8, 55 mM NaCl, and 1.1 mM EDTA. p397-PDZ3 was separated from PDZ3 using Source Q resin with a 260-ml linear gradient from 0 to 500 mM NaCl in 20 mM Tris-HCl, pH 7.0 (Fig. 2A). The identity and purity of the protein were verified by mass spectrometry and SDS-PAGE. Protein concentrations were determined by UV absorbance with extinction coefficients calculated by the ProtParam tool. Phosphorylation changes the UV absorbance profile of tyrosine with an extinction coefficient of 593 m$^2$ cm$^{-1}$ at 268 nm (32). Taking the other unphosphorylated tyrosine (Tyr-392; $e_{268} = 1114$ m$^2$ cm$^{-1}$) into account, the concentration of p397-PDZ3 was determined using an extinction coefficient of 1707 m$^2$ cm$^{-1}$ at 268 nm.
PDZ3-SH3 single and triple mutants were purified using the aforementioned protocol with modification at the size exclusion chromatography step where elution buffer contained 50 mM Tris-HCl, pH 7.5, 200 mM NaCl, and 1 mM DTT. The plasmid encoding c-Src kinase was a gift from John Kuriyan (University of California, Berkeley) and Sharon Campbell (University of North Carolina at Chapel Hill). The c-Src kinase domain was purified as described (33).

Peptide Synthesis—A 7-mer CRIPPT peptide (Ac-NYKQTSV-COOH) was synthesized and purified as reported (21). The identity and purity of the peptide were verified by mass spectrometry. The peptide concentration was determined using UV absorbance with an extinction coefficient of 1490 M\(^{-1}\)cm\(^{-1}\).

Isothermal Titration Calorimetry Experiments—ITC experiments were carried out on a MicroCal VP-ITC microcalorimeter and repeated twice to estimate error. ITC measurements of PDZ3, p\(^{397}\)-PDZ3, and Y397E-PDZ3 were performed by titrating 1 mM CRIPPT peptide into 0.1 mM protein. In total, 280 ITC measurements were carried out on a MicroCal VP-ITC microcalorimeter using Origin version 5.0 (MicroCal). Data were integrated and analyzed using in-house programs. Briefly, data from “rigid” residues were initially fit to obtain the overall correlation time of the protein, which was determined to be 6.8 ns. As with unphosphorylated PDZ3, anisotropic models of tumbling were not required (21). Models were then selected for individual residues (supplemental Table S1) to fit the Lipari-Szabo dynamics parameters (\(S^2\), \(\tau_a\), etc.).

Solvent-exposed Amide (SEA)-HSQC Experiments—Solvent-exposed amide HSQC spectra were collected at 500 MHz using a \(^1\)H/\(^15\)N/\(^13\)C probe and z axis pulsed field gradients at 298 K. The original SEA-HSQC pulse sequence was developed by Lin et al. (36) using a clean chemical exchange-phase modulated-PM mixing scheme. In this study, a Biopack version of the SEA-HSQC pulse sequence was used. The intertransient delay was set to 1.5 s, and the mixing time was 0.1 s. The spectra of PDZ3 and p\(^{397}\)-PDZ3 were collected at a protein concentration of 1 mM with 16 transients. NMR intensity is a function of several factors such as protein concentration, nuclear relaxation time, solution conductivity, and protein tumbling time. Of these factors, protein tumbling time is significantly influenced by phosphorylation (see “Results”). This necessitates a method to scale the SEA-HSQC intensities of PDZ3 and p\(^{397}\)-PDZ3 to allow direct comparison. The scaling procedure used the residues that are not affected by phosphorylation, i.e. those residues that experience slight or nearly no chemical shift perturbation as shown in Fig. 2A. The SEA-HSQC peak intensities of p\(^{397}\)-PDZ3 for these residues were scaled to match those of PDZ3.

RESULTS

Phosphorylation of Tyr-397 and Purification of Phosphorylated PDZ3—From the crystal structure of PDZ3 (14), the side chain of Tyr-397 is partially buried between a3 and the core domain of PDZ3 (Fig. 1B). This tyrosine residue is phosphorylated in mouse brain (27), although it is unknown which kinase in vivo is responsible for its phosphorylation. Tyr-397, located at the linker between PDZ3 and SH3, is highly conserved throughout MAGUK proteins of various organisms (Fig. 1A). Using the ExPasy Proteomics Server, we analyzed the first 100 homologue sequences of PSD-95, which belong to different organisms, different subfamilies, and different splicing forms. Only one conserved mutation (Tyr to Phe) was found. The high conservation of Tyr-397 implies its significance within the MAGUK family. Members of the Src family are known to interact with PSD-95 (37). Using the Group-based Prediction System (38), Src kinase was predicted to be capable of phosphorylating Tyr-397. Treatment of \(^1\)N-labeled PDZ3 with c-Src resulted in a new set of resonances in the HSQC spectrum of PDZ3. The relative peak intensities suggested that only 10% of PDZ3 was phosphorylated. The reaction went further to completion after removing ADP by dialysis and adding fresh ATP, which was repeated four times. This yielded ~40% putative p\(^{397}\)-PDZ3.

Ion exchange chromatography was used to purify p\(^{397}\)-PDZ3 based on the additional negative charge introduced by phosphorylation. The elution profile displayed five discrete species, three of which tested positive for protein (Fig. 2A). The first peak was confirmed to be PDZ3 from HSQC, and the third peak was confirmed to be pure phospho-PDZ3. An overlay of HSQC spectra of PDZ3 and putative p\(^{397}\)-PDZ3 shows peaks shifting in regions of PDZ3 consistent with modification at Tyr-397 (Fig. 2B). The identity of the third peak was also analyzed by mass spectrometry (supplemental Fig. S2). Although the electrospray ionization ion trap mass spectrum indicated the correct mass for a singly phosphorylated species (supplemental Fig. S2A), MS-MS MALDI-TOF results suggested that the
dominant species of the third peak was p397-PDZ3, although small amounts of p392-PDZ3 (10%) could not be ruled out (supplemental Fig. S2B). A species other than p397-PDZ3 was not found in 1H-15N HSQC, HNCO, HNCACB, or CBCA-(CO)NH spectra. Based on this, the following NMR-based observations were interpreted to arise solely from p397-PDZ3.

The above procedure was used to obtain milligram quantities of p397-PDZ3 for further biophysical characterization.

Phosphorylation-induced Undocking of α3 Lowers Affinity for CRIP1 Ligand—To determine the effect of phosphorylation at Tyr-397 on PDZ3 structure, the backbone chemical shifts of p397-PDZ3 were assigned and compared with unphosphorylated PDZ3 (Fig. 3A). A species other than p397-PDZ3 was not found in 1H-15N HSQC, HNCO, HNCACB, or CBCA-(CO)NH spectra. Based on this, the following NMR-based observations were interpreted to arise solely from p397-PDZ3. The above procedure was used to obtain milligram quantities of p397-PDZ3 for further biophysical characterization.

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In unphosphorylated PDZ3, the side chain of Tyr-397 packs into side chains of the β2 strand, the β3 strand, and the α1-β4 loop of the PDZ core. These hydrophobic interactions help to define how α3 packs against the PDZ core (Fig. 3B). One possible consequence of phosphorylation is therefore destabilization/unfolding of α3 because the phosphate group might occlude proper packing of α3. In this sense, phosphorylation of Tyr-397 would effectively convert PDZ3 into a C-terminally truncated form, much like PDZ(303–395) characterized previously and referred to as Δ7ct (21). To test this idea, we compared 1H, 15N amide chemical shifts of PDZ3, p397-PDZ3, and Δ7ct by superimposing HSQCds of the three proteins. Nearly all residues experiencing significant CSPs in p397-PDZ3 (Fig. 3) display a linear trend of HSQC peaks in the order PDZ3, p397-PDZ3, and Δ7ct (Fig. 4A). To quantitatively demonstrate linearity of these three sets of resonances, correlation values...
Values of linear fitting were calculated and plotted (Fig. 4B). Most R values are around 1, indicating high linearity for many residues. Because chemical shifts are highly nonlinear with respect to positioning of structural elements, such extensive linearity cannot be reasonably explained as accidental. Rather, the simplest explanation is that the peak positions of p397-PDZ3 are averaged due to fast exchange between two conformations: one in which H9251 is packed as in native PDZ3 and one in which H9251 is undocked and makes no contributions to chemical shifts in PDZ core residues. Thus, p397-PDZ3 exists as an equilibrium of two dynamically exchanging states represented by PDZ3 and H7ct, respectively. Preliminary 15N Carr-Purcell-Meiboom-Gill relaxation dispersion experiments (39) on p397-PDZ3 did not detect this exchange; and thus, we conclude that exchange takes place on the microsecond or possibly even nanosecond timescale. Because p397-PDZ3 peaks occurred on average approximately half-way between unphosphorylated PDZ3 and H7ct peaks, the fraction of undocked H9251 was estimated to be ~50%.

It was shown previously that truncation of α3 decreases binding affinity of PDZ3 with CRIPT peptide by 21-fold (21). If p397-PDZ3 is indeed a mixture of PDZ3- and H7ct-like states, an intermediate binding affinity to CRIPT peptide should be observed. To test this, we performed ITC experiments to measure the binding affinity of p397-PDZ3 to CRIPT C-terminal peptide. The binding affinity of p397-PDZ3 was found to be ~4 times weaker than PDZ3, falling in between PDZ3 and H7ct (Table 1). Importantly, this change in $K_d$ due to phosphorylation is mainly attributed to entropy, which is the same signature observed for H7ct (21).

Phosphorylation of Tyr-397 Increases Disorder in H9251—Phosphorylation of Tyr-397 disrupts the packing of H9251 with the core domain of PDZ3 as described above. However, it is unknown how phosphorylation affects the local structure of H9251. Does H9251 still maintain its secondary structure when H9251 is undocked from PDZ3? To answer this question, we analyzed 13C chemical shifts, amide exchange rates, and 15N backbone relaxation in phosphorylated and unphosphorylated PDZ3.

The 13C chemical shift relative to random coil values is a sensitive indicator of secondary structure (40). These relative chemical shift values for unphosphorylated PDZ3 and p397-PDZ3 are shown for the C-terminal residues that include H5385–392 and α3 H394–398 (Fig. 5). Phosphorylation clearly increases disorder in H9251.
Phosphorylation Tunes PDZ3 Interactions in PSD-95

Amide proton exchange experiments on solvent accessibility and the stability of secondary structure. We measured these exchange rates using the SEA-HSQC experiment. This approach was used because the amides in the α3 region exchange rapidly (data not shown). In a SEA-HSQC spectrum, the intensity of an amide resonance is proportional to the extent to which it is solvent-exposed (36). As shown in supplemental Fig. S3A, only surface-exposed residues of PDZ3 are observed in the spectrum. The undocking of α3 is expected to increase solvent exposure, which should be evident from SEA-HSQC intensity changes. However, it is not appropriate to directly compare SEA-HSQC intensities of PDZ3 and phospho-PDZ3 samples as intensities are differentially affected by protein concentration and tumbling times. Therefore, intensities were normalized (see “Experimental Procedures”) to facilitate direct comparison. SEA-HSQC peak intensities of PDZ3 and p397-PDZ3 are plotted against one another in supplemental Fig. S3B. A good correlation of these peak intensities is obtained as indicated by an R value of 0.98, and a slope of 0.74 (for p397-PDZ3 relative to PDZ3) was observed. As expected, α3 and its docking region displayed increased solvent exchange (supplemental Fig. S3, C and D), whereas other regions have few significant changes in solvent accessibility.

To directly determine the degree of flexibility in p397-PDZ3, we used 15N backbone relaxation coupled with model-free analysis (41) to yield an order parameter, S2, for each residue. The most significant change in S2 upon phosphorylation was in the α3 region, which showed decreases of 0.1–0.3 (Fig. 6). For residues in α3 with significant helicity (394–400), the average decrease in S2, however, is only 0.10. Given that the undocked fraction was ~50% and that the docked state showed average S2 values of ~0.85, this suggests that the undocked fraction yields on average S2 values of ~0.65 for residues 394–400; this is much higher than expected for a fully unfolded C-terminal tail. Thus, it is more likely that upon undocking these “α3” residues redistribute some of their helical character into a mixture of extended and random coil-like states with the latter contributing most significantly to a decrease in S2. This would allow for relatively high S2 values and the observed 13Cα chemical shifts because extended backbone conformations would help to reduce the downfield shifts from random coil values (Fig. 5). In addition to S2 effects at the C terminus, a slight overall decrease in S2 was observed throughout the PDZ core that is suggestive of an overall small increase in backbone dynamics in p397-PDZ3. These results are similar to that found in Δ7ct, which showed large increases in side-chain dynamics and subtle increases in backbone dynamics (21). From the 15N relaxation

### TABLE 1

| Binding parameters | PDZ3 | p397-PDZ3 |
|--------------------|------|-----------|
| K_s (μM)           | 3.60 ± 0.28 | 14.00 ± 0.47 |
| ΔH (kcal/mol)      | -8.57 ± 0.31 | -8.80 ± 0.29 |
| -TΔS (kcal/mol)    | -1.16 ± 0.26 | -2.18 ± 0.31 |

in the context of full-length PSD-95 (Fig. 1A). Although it may not be feasible to probe this 80-kDa protein for phosphorylation effects by NMR, we hypothesized that some full-length behavior might be captured in shorter multidomain constructs. Thus, we set out to characterize the behavior of PDZ3 in a construct that spans residues 304–532, which includes the SH3 domain and the linker preceding it. In addition, the F-strand from the GK domain (residue numbers 713–724) was added to the C terminus to enhance stability (16). This PDZ3-SH3 construct yielded reasonable quality HSQC spectra (supplemental Fig. S4) from which the effects of phosphorylation could be monitored.

In PDZ3 alone, phosphorylation at Tyr-397 initiated α3 undocking and a redistribution of its sampled conformations. In the intact MAGUK, because α3 precedes a ~25-residue linker that connects to the SH3, phosphorylation may affect putative domain packing between PDZ3 and SH3. To test this, 15N backbone transverse relaxation times (T2) were measured in PDZ3-SH3. T2 times are highly sensitive to the rate of tumbling experienced by individual sites. An overall increase in T2 is expected for a reduced degree of domain-domain packing because loss of packing would lead to more time spent tumbling as an isolated domain. For these experiments, phosphorylation at 397 in PDZ3-SH3 was mimicked by the Y397E mutation, which yielded a ~7-fold reduction in CRIPT binding affinity in the context of the PDZ3 construct (supplemental Fig. S5). The validity of the mimic was also confirmed by 13Cα chemical shifts. These shifts indicate an almost identical loss of helicity (relative to unphosphorylated PDZ3) in Y397E-PDZ3 compared with p397-PDZ3 (Fig. 5).

T2 values for the mutant and wild-type constructs were measured, and their differences are shown in Fig. 7A. In general, Y397E-PDZ3-SH3 showed T2 values that are ~3–25 ms longer relative to wild type. The increase in T2 at virtually all residues suggests that phosphorylation at Tyr-397 reduced the interaction between PDZ3 and SH3, which appeared to be stabilized by the presence of α3 helical secondary structure.

Two additional serine phosphorylation sites (Ser-415 and Ser-418) within the PDZ3-SH3 linker region have been reported (42). The fragment where Ser-415 and Ser-418 are located corresponds to a fragment that was shown to contribute to communication between PDZ3 and GK in the PSD-95 homologue Dlg-1 (25). Specifically, deletion or replacement of this “A2” fragment with random coil segments (from Gly-Ser repeats) abolishes the PDZ3-dependent interaction of the GK domain with GukHolder. Because of the proximity of these res-
idues to α3 and their conserved nature (Fig. 1B), we created a triple phosphorylation mutant to test the effect of phosphorylation at these two additional sites. 15N T2 measurements were made on a Y397E/S415E/S418E triple mutant of PDZ3-SH3. Compared with the Y397E single mutant, the triple mutant exhibited even higher T2 values (Fig. 7B), suggesting stronger dissociation of PDZ3 and SH3 domains.

DISCUSSION

Regulation in PDZ Domains—When PDZ domains are excised from their parent proteins, they typically behave as simple binders of C termini, which is documented by the many determined structures of individual PDZs and their cognate ligands (43). Current evidence indicates, however, that PDZ function can be influenced by flanking sequences and/or domains as well as intermolecular interactions (30). For example, the PDZ domain of Par6 alters its binding affinity for ligand upon binding of Cdc42 to the semi-Crib motif, which is immediately N-terminal to the PDZ domain, and augments its β-sheet (18). In a reverse sense, “PDZ proteases” are regulated by ligand-induced changes in PDZ structure (20). PDZ function has also been shown to be regulated by redox events within the PDZ domain (44), and this redox potential can be allosterically regulated by binding to yet another PDZ domain (45). The idea that PDZ domains are involved in higher order regulatory events is consistent with their locations (often in tandem) in proteins involved in organization of signaling complexes, subcellular transport, and cell-cell adhesion (9, 43).

Here, we show how PDZ3 from PSD-95 is regulated by phosphorylation of Tyr-397, which lies outside the strict PDZ boundary in a C-terminal extension that augments the PDZ core structure (14, 21). Although phosphorylation was shown to regulate function of the PDZ protein NHERF1, the phosphorylation sites are far from the PDZ domains, and their precise mechanism of release of autoinhibition remains to be elucidated (26). In contrast to phosphorylation of PDZ domains themselves, there are numerous examples of phosphorylation of PDZ substrates (46–54). Phosphorylation typically occurs at positions −2 to −4, and although this usually results in weakening of the PDZ-ligand interaction, there are examples of phosphorylation strengthening the interaction (49, 53, 54). Thus, phosphorylation appears to be a recurring regulatory mechanism in PDZ-mediated interactions. Although there is currently little direct characterization of how phosphorylation influences PDZ domains, proteomics work suggests that phos-
Phosphorylation in PDZ core or flanking sequences is relatively common. As collected in the UniProt database, numerous phosphorylation sites have been experimentally confirmed (mostly in murine proteins) for the PDZ proteins ZO-1, GRIP1, MAGI2, MAGI3, and Par3, for example. Many of these phosphorylation sites exist in short linkers between domains as observed here in PSD-95 but also within the PDZ domains.

Phosphorylation at Tyr-397 Allosterically Regulates Binding of C Terminus—We successfully phosphorylated (at Tyr-397) a construct of PDZ3 that includes an α-helical C-terminal extension that ends at position 402. A combination of chemical shift analysis, amide hydrogen exchange, and 15N relaxation showed that phosphorylation partially undocks the α3 helix from the PDZ3 core. The linear change in chemical shifts (Fig. 4) indicates that p397-PDZ3 exists as a rapid equilibrium between a docked state in which α3 is natively packed and an undocked state in which “α3 residues” have lost some helical character. This conclusion was only possible because of previous characterization of Δ7ct, which eliminates the native α3 by truncation of residues 396–402 (21). In that study, α3 was shown to confer a 21-fold increase in binding affinity to CRIPT peptide. Without α3, the PDZ3 core has enhanced side-chain dynamics that are quenched upon CRIPT binding. This confers an entropic penalty to binding, which was detected through isothermal titration calorimetry. Interestingly, because α3 does not contact CRIPT peptide, this was considered a novel dynamic allosteric mechanism. Here, we show that this dynamic mechanism has biological significance and was induced by phosphorylation at Tyr-397. The 4-fold reduction in CRIPT binding affinity (as opposed to 21-fold) is consistent with the ~50% population of undocked α3. The behavior of p397-PDZ3 also confirmed that Δ7ct serves as an excellent proxy for phosphorylated PDZ3 constructs.

Phosphorylation Tunes Interdomain Interactions in PDZ3-SH3-GK MAGUK Core—A caveat of the results on p397-PDZ3 (residues 304–402) here is that the construct remains artificial as an excised domain. (One reason in support of using this construct is that phosphorylation at Tyr-397 was less efficient upon further extension of the C terminus and hence yields of such phosphorylated constructs were unacceptably low.) Nevertheless, the role of the linker may be important. Although there is no crystal structure of full-length PSD-95 or the PDZ-SH3-GK supradomain from any MAGUK, there is evidence for interaction between PDZ3 and SH3-GK. From electron microscopy, intact PSD-95 appears as a “C-shaped” arrangement of domains (55). In other work, binding assays on various constructs of PSD-95 family MAGUKs suggest interdomain allostery between PDZ3 and the SH3-GK module. In the case of PSD-95, PDZ3 (as well as PDZs 1 and 2) was shown to influence binding of MAP1a, which binds to the GK domain (24). A similar result was found for SAP97 (a closely related PSD-95 family MAGUK) binding to guanylate-kinase-associated protein (56). In the case of the PSD-95 homologue Dlg-1 from Drosophila, PDZ3 influences binding of GukHolder, which binds to the SH3 domain (25). Structural insight into how PDZ3 and its following linker interact with SH3-GK is needed to unravel the basis for these interactions and how they are regulated.

To gain insight into how PDZ3 interacts with SH3-GK and whether phosphorylation plays a role, we studied a PDZ3-linker-SH3 construct. 15N T2 measurements within the PDZ3 domain for wild type as well as for a Y397E phosphomimic and a triple phosphomimic construct, Y397E/S415E/S418E, were made. Ser-415 and Ser-418 have also been confirmed to be phosphorylated (28, 29), and therefore a combination of these sites may be phosphorylated under different signaling and cell contexts. Phosphorylation at Tyr-397 increased the mobility of PDZ3, and this effect was increased further for the triple mutant (Fig. 7). This is consistent with a model in which PDZ3 links to SH3-GK.
interacts with SH3-GK via the SH3 domain and phosphorylation at 397 in α3 and 415/418 in the linker disrupts this interaction (Fig. 8). To our knowledge, these are the first reported physical data that relate PDZ3 to SH3 for any MAGUK. To date, there are no reported structures for any construct containing PDZ3 and SH3 domains, although structural models of PDZ3-SH3-GK have been proposed (56, 57). The $T_2$ results suggest that additive phosphorylation interferes with the putative PDZ3-SH3 interaction. Because positions 415 and 418 are well beyond the PDZ3 domain (and extension), the $T_2$ data also suggest that the linker plays a role in the association of PDZ3 and SH3. It is currently unknown whether phosphorylation at these sites occur in an ordered pattern, but it is possible that phosphorylation of Ser-415/Ser-418, which are seated in a putatively flexible fragment (SSLGSG), precedes phosphorylation of Tyr-397, which is mostly buried. Phosphorylation of Ser-415/Ser-418 may then facilitate Tyr-397 phosphorylation by loosening the domain packing between PDZ3 and SH3 or providing binding sites for the kinase that phosphorylates Tyr-397. Overall, our NMR relaxation measurements suggest that phosphorylations of Tyr-397, Ser-415, and Ser-418 cooperate together to regulate interdomain communication of PSD-95.

It is currently unclear how conserved these phosphorylation events are. The PSD-95 family of proteins comprises PSD-95, PSD-93, SAP97, and SAP102. Although there is high sequence conservation in the α3 and linker among these proteins, phosphorylation at Tyr-397 has only been confirmed for PSD-95 and PSD-93, and phosphorylation at 415/418 has only been observed so far in PSD-95. Although it may be possible that differences in phosphorylation patterns in this region contribute to functional differences between these MAGUKs (which also have functional redundancies), this cannot be resolved until more in-depth phosphorylation mapping is carried out.

Concluding Remarks—PDZ domains are primarily appreciated for their simple function of binding C-terminal sequences. However, PDZs are typically embedded in lengthy protein sequences characterized by multiple other structured domains and significant stretches of unknown structure/flexibility. Understanding native PDZ function therefore often requires consideration of these additional elements. There are now a number of examples of PDZ domains whose core structures are augmented by N- or C-terminal extensions (14, 22, 58–60), and a recent review predicts that one-third of PDZ domains contain such structural augmentations (30). We show here that the effects of such additional segments can be dynamically modulated by phosphorylation, building upon previous work that showed that the external segments can regulate PDZ function. In many multidomain proteins, the role of interdomain linkers is not well understood. By mimicking phosphorylation at Tyr-397 and in the linker of the PDZ3-linker-SH3 construct from PSD-95, progressive increases in mobility in PDZ3 due to linker phosphorylation were detected. This suggests that the linker plays a key role in supradomain assembly. We propose that linkers in MAGUKs and in other PDZ proteins can adopt varying degrees of structure/flexibility that serve to tune interdomain communication.

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