Origins of Allostery and Evolvability in Proteins: A Case Study

Graphical Abstract

Highlights

- Adaptation in a PDZ domain involves intermediate mutations that bridge ligand classes
- A simple model shows that class-bridging mutations are evolutionarily preferred
- Class-bridging mutations act allosterically
- The origin of allostery in proteins could be in the capacity to adapt

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In Brief
Mutations that are neutral but that potentiate functional protein adaptation by a subsequent mutation result in enhanced conformational plasticity at the distal “business end” of the protein. The data suggest a model in which the origin of allostery lies in the capacity for adaptation.

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Origins of Allostery and Evolvability in Proteins: A Case Study

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SUMMARY
Proteins display the capacity for adaptation to new functions, a property critical for evolvability. But what structural principles underlie the capacity for adaptation? Here, we show that adaptation to a physiologically distinct class of ligand specificity in a PSD95, DLG1, ZO-1 (PDZ) domain preferentially occurs through class-bridging intermediate mutations located distant from the ligand-binding site. These mutations provide a functional link between ligand classes and demonstrate the principle of “conditional neutrality” in mediating evolutionary adaptation. Structures show that class-bridging mutations work allosterically to open up conformational plasticity at the active site, permitting novel functions while retaining existing function. More generally, the class-bridging phenotype arises from mutations in an evolutionarily conserved network of coevolving amino acids in the PDZ family (the sector) that connects the active site to distant surface sites. These findings introduce the concept that allostery in proteins could have its origins not in protein function but in the capacity to adapt.

INTRODUCTION
Proteins display the capacity to fold, often into well packed three-dimensional structures, and to carry out biologically essential activities such as catalysis, signal transmission, and allosteric regulation. The amino acid sequence reflects the constraints arising from these properties, and considerable prior work has focused on understanding how the sequence encodes folding and biochemical function (Anfinsen, 1973; Bowie et al., 1990; Halabi et al., 2009; Morcos et al., 2011; Suel et al., 2003). However, it has been appreciated for decades that there may be non-trivial pressures on proteins that come not just from the physics of folding and function but also from the process of evolution itself (Anfinsen, 1973; Bershtein et al., 2006; Smith, 1970; Tokuriki and Tawfik, 2009; Wagner, 2005). For instance, proteins must arise through a random, iterative, stepwise process of mutation and selection, and they must be capable of adaptive variation as conditions of fitness vary in the environment. These considerations may place unique and yet unknown “design” constraints on evolved proteins, a missing aspect of our current understanding. An example of such a constraint might be functional connectivity in adaptive paths—the requirement that adaptation between fitness peaks proceeds through intermediates that maintain function above the threshold of selection (Smith, 1970).

Recent work has begun to elucidate the general properties of adaptation that are likely critical for the evolution of biological systems (Aakre et al., 2015; Bershtein et al., 2006; Bloom et al., 2005, 2006; Draghi et al., 2010; Draghi and Plotkin, 2011; Hayden et al., 2011; Tokuriki and Tawfik, 2009; Wagner, 2005). One such property is conditional neutrality, a special case in which mutations have no significant effect in the existing genetic or environmental background but have a significant effect upon subsequent changes in either the genome or environment (Draghi et al., 2010; Draghi and Plotkin, 2011; Hayden et al., 2011; Wagner, 2005). Such variations are “cryptic” in the sense that they hide their effects on fitness until exposed in the right setting and can therefore accumulate and pre-exist in populations as standing genetic variations (Luria and Delbrück, 1943). Because conditionally neutral mutations arise without selection and only express their fitness advantages upon subsequent events, they are said to be pre-adaptive (or “exaptive”); Gould and Vrba (1982) and represent a pool of variants that can facilitate the emergence of novel adaptive phenotypes. Indeed, conditional neutrality has been convincingly demonstrated to facilitate adaptation both theoretically (Draghi et al., 2010) and experimentally (Hayden et al., 2011) and, conceptually, represents the key link between the two major driving forces for genetic variation in populations: neutral drift and selection (Draghi and Plotkin, 2011). Understanding the prevalence and structural principles of conditional neutrality in protein molecules represents a key step in linking biophysical variation at the molecular level to evolutionary viability.

In this work, we carry out a series of experiments and simulations in the PSD95, DLG1, ZO-1 (PDZ) family of protein interaction modules to explore the prevalence, structural origin, importance, and mechanism of conditional neutrality. We show that conditional neutrality emerges from a distributed network of physically contiguous amino acids that permits a specific set of distantly positioned residues to unlock new structural and functional states at the ligand binding site. These adaptive positions are contained within the PDZ protein sector—a conserved...
and coevolving network of amino acid positions in the entire protein family (Halabi et al., 2009; McLaughlin et al., 2012). These findings allow us to propose basic structural principles of adaptation, and they open the hypothesis that the capacity to adapt may be the origin of allosteric mechanisms in proteins.

RESULTS

A Two-Mutation Path to New Specificity

A case study of protein adaptation is evident in PSD95

\[ \text{pdz3} \], a member of the PDZ protein family (Halabi et al., 2009; McLaughlin et al., 2012). PDZ domains are roughly 100-amino acid mixed \( \alpha/\beta \) folds that typically recognize the C-terminal region of target proteins at a groove formed between the \( \beta2 \) strand and \( \alpha2 \) helix (Figure 1A); for simplicity, we refer to this groove as the “active site.” Ligand specificity is largely defined by the amino acid sequence in the last four positions (numbered in reverse order from the carboxyl terminus [position 0]), with the major determinant of specificity being the identity of the residue at the antepenultimate (−2) position (Songyang et al., 1997; Stöffler et al., 2007; Tonikian et al., 2008). For example, class I domains bind ligands with a consensus of \( \text{S/T-X-}X\phi\text{-COOH} \) (\( \phi \) is any amino acid, \( \phi \) is hydrophobic) and class II domains with a consensus of \( \text{S/T-X-}X\phi\text{-COOH} \). PSD95

\[ \text{pdz3} \] is an archetypical class I domain, binding its cognate ligand \( \cdot\text{TKNYKQ5SV-COOH} \), derived from the cysteine-rich interactor of PDZ (CRIPT) (Niethammer et al., 1998) with an affinity (indicated by the equilibrium dissociation constant, \( K_d = 0.8 \mu \text{M} \)) that is 45-fold higher than to a class-switching mutant peptide \( \cdot\text{T2F, TKNYKQ5FSV-COOH} \), specifying a Thr-to-Phe substitution at the −2 position (Figure 1B). Because PDZ domains operate with dissociation constants in the 1−15\( \mu \text{M} \) range (Stöffler et al., 2007), this dynamic range represents a physiologically significant degree of specificity. Previous work shows that a double mutant of PSD95

\[ \text{pdz3} \] (G330T,H372A) displays a near-complete reversal of specificity for these two peptides (McLaughlin et al., 2012). It shows roughly a 45-fold preference for the T-2F ligand and an affinity (\( K_d = 0.5 \mu \text{M} \)) that is similar to that of the wild-type protein for the CRIPT ligand (Figure 1B).

These data frame a minimal instance of the problem of adaptive mutational paths. H372A and G330T represent two distinct strategies for achieving the class-switching specificity change. Position 372 is directly located at the ligand binding site and interacts with the residue at the −2 position of the target peptide. For example, in PSD95

\[ \text{pdz3} \], the histidine residue at this position makes a hydrogen bond with the hydroxyl group on the −2 threonine side chain of the ligand (Figure 1C), an interaction that is thought to represent the basis for the Ser/Thr specificity of class I PDZ domains (Doyle et al., 1996). Not surprisingly, the H372A mutation shows a substantial (34-fold) loss of affinity for the CRIPT ligand (Figure 1B). It also shows gain of function for the T-2F ligand, and, thus, this single active site mutation on its own provides a partially class-switching phenotype (13-fold T-2F preference) that is then enhanced by the G330T mutation to quantitatively complete the specificity switch.

In contrast, the G330T-first path is strikingly different. Position 330 is positioned on a surface loop (\( \beta2-\beta3 \)) that is packed against the region of position 372.

Figure 1. A Two-Mutation Path to New Functional Specificity in a PDZ Domain

(A) The structure of the PDZ domain (PSD95

\[ \text{pdz3} \], PDB: 1BE9) bound to the CRIPT C-terminal peptide (yellow stick bonds). Positions G330 and H372 in the protein and T-2 in the ligand peptide are shown as spheres with an overlay of the van der Waals surface.

(B) A thermodynamic cube showing the effects of the G330T and H372A mutations in the context of the wild-type CRIPT ligand (top face) and the T-2F ligand (bottom face). Wild-type PSD95

\[ \text{pdz3} \] shows a 45-fold preference for the CRIPT ligand, whereas the G330T,H372A double mutant shows a 45-fold preference for the T-2F ligand.

(C) Stereochemical details around ligand position −2. H372 makes a hydrogen bond with the class I-defining threonine side chain of ligand position −2, and G330 occurs on a surface loop (\( \beta2-\beta3 \)) that is packed against the region of position 372.
A Comprehensive View of Ligand Specificity in PDZ Domains

To test the notion that G330T acts as a functional bridge between class specificities, we measured the binding of wild-type, G330T, H372A, and the double-mutant protein to a library of all possible peptide ligands defined by randomizing the C-terminal four amino acid positions (2^{16} = 160,000 total ligands). This analysis is made possible by a quantitative bacterial two-hybrid (BTH) assay for PDZ function in which transcription of a reporter gene is tuned to be linearly proportional to the binding free energy of PSD95 to target ligands (Figure S1; McLaughlin et al., 2012). The reporter gene is chloramphenicol acetyl transferase (CAT), and thus the binding profile of any PDZ variant over the full space of all ligands can be assessed simply by selecting bacterial cells carrying the BTH on chloramphenicol and deep sequencing of the library before and after selection. The binding between PDZ and each ligand is given by ΔE_r, the normalized log ratio of observing ligand x in the selected and unselected libraries (Experimental Procedures; Figure S1). We obtained excellent counting statistics for 154,521 of the 160,000 possible ligands for all four PSD95 variants (Table S1)—a near-complete global profile of ligand specificity over the adaptive path defined by the G330T, H372A double mutant.

Hierarchical clustering of a few top binding sequences provides an intuitive preview of the full dataset (Figure 2A). Consistent with the data in Figure 1B, PSD95 binds ligands with T/S preference at the -2 position (class I), H372A and the double mutant bind ligands with hydrophobic amino acids at -2 (class II), and G330T binds ligands in both classes. The full dataset is very high-dimensional and impossible to directly visualize. Accordingly, we used principal components analysis (PCA) to project all peptides bound with better than 15 μM affinity by any of the four PSD95 variants (2,359 total) onto a two-dimensional space based on their profile of binding (Figure 2C). This low-dimensional projection is well justified. The first two principal components (V1 and V2) capture nearly 97% of the total variance representing relationships between the 2,359 ligands with physiologically relevant affinity (Figure 2B). The principal components also have clear biochemical meaning. V1 separates ligands by amino acid preference at the class-defining position —2 to left of the origin, bulky hydrophobic [class II] to the right, and V2 separates ligands by a combination of preference at both —2 and —3 (Figure 2C). Thus, the PCA provides a statistically accurate and intuitive representation of ligand specificity over the adaptive path.

Figure 3 shows the binding profiles of wild-type, G330T, H372A, and the double mutant variants of PSD95 projected onto the two-dimensional space. With the 15 μM cutoff for physiological relevance, the wild-type exclusively binds peptides in the class I space (Figure 3A), and both H372A and the double mutant bind ligands exclusively in the class II space (Figures 3C and 3D). This confirms that both the H372A single mutant and the double mutant are able to switch class specificity. In contrast, G330T shows a distinct binding profile that provides a link between the space of class I and class II ligands (Figure 3B). Thus, G330T is indeed a functional bridge—a single mutation that makes PSD95 capable of recognizing both classes of ligands. Amino acid profiles of ligands bound by the four PSD95 variants over a range of threshold binding affinities reinforce these findings (Figure S2). These data also provide a data-rich illustration of the concept of conditional neutrality—a mutation that preserves...
existing function while opening up new functions; in this case, binding to physiologically distinct ligand variants.

Note that G330T is not a "generalist" or "promiscuous" protein in any typical sense. It maintains high-affinity binding and only recognizes 846 ligands of 154,521 total, a number that is in the same range as the other, more class-specific PSD95\textsuperscript{pdz3} variants (Table S1). It also recognizes ligands only in the lower half of the ligand space spanned by the top principal components (Figure 3B), showing that the effect is mainly in the shape, rather than the breadth, of the binding profile. Thus, G330T is a targeted class-bridging mutation capable of providing a near-neutral path from the native class I ligand (CRIPT) to a specific region of class II specificity. Subsequent acquisition of H372A to make the double mutant then localizes and reconfigures the binding specificity completely in the class II space (Figure 3D).

**Preferential Adaptive Path**

Which path—G330T first or H372A first—is evolutionarily more likely to achieve the specificity switch defined by the double mutant in PSD95\textsuperscript{pdz3}? Given that H372A provides a partial class switch in just one mutation (Figure 1B), it is important to understand what advantage (if any) there is in the class-bridging phenotype of G330T as an intermediate in the adaptive path. To study this, we carried out computational simulations of evolutionary dynamics between the wild-type and G330T, H372A double mutant states over a range of mutation rates and ligand-switching rates. Each trial of simulation is initiated with a population of wild-type PSD95\textsuperscript{pdz3} genotypes (N = 1000), which is large given the small number of genotypes considered (four; Figure 4A). At each generation, single mutations between genotypes are allowed with a probability $m$ and double mutations with probability $m^2$, the target ligand switches between the class I CRIPT peptide and the class II T-2F peptide every $t$ generations, and fitness at every generation is defined as a fraction-bound of ligand determined from the experimentally defined equilibrium dissociation constants (Figure 1B). The total ligand concentration is set to 100μM, a value in the middle of the specificity range of wild-type PSD95\textsuperscript{pdz3}—the relevant regime for non-trivial dynamics. In essence, this simulation gives an opportunity to study how the flux between the two paths to the double mutant state depends on both internal parameters (mutation rate and population size) and external parameters (environmental switching between the two class-distinct ligands).

A representative simulation trajectory at one particular mutation and ligand-switching rate illustrates properties of the adaptive process (Figure 4B). In this case, $t = 500$ and $N\mu = 1$, meaning that the ligand switches every 500 generations, and one single mutation is made, on average, at every generation. As expected, the wild-type genotype (black trace) is the most fit in the presence of the class I CRIPT ligand, with a small fraction of other genotypes stochastically occurring in the population according to the mutation rate and on their fitness relative to the wild-type. Switching to the T-2F ligand causes the population to ultimately switch to the double mutant state (blue), the genotype that is most fit for the boundary is defined by affinity ≤ 15μM. The data show that the wild-type binds exclusively to class I ligands, that H372A and the double mutant bind exclusively to class II ligands, and that G330T is a bridge between class specificities, binding a subset of ligands in both class I and class II regions.
T-2F ligand under these simulation conditions. However, the path of switching can show considerable trial-by-trial variability with regard to intermediates. For example, in this trajectory, G330T (green) is more prevalent in trial 1 and H372A (red) more prevalent in trial 2. Averaged over many trials of switching (~500 events) from CRIPT to T-2F, we find that G330T is by far the preferred path of adaptation to the double-mutant state given the selected mutation and ligand-switching rates (Figure 4D).

How can we understand this result mechanistically? Because both G330T and H372A can bind the T-2F ligand about equally well (Figure 1B), the path simply depends on the relative availability of these genotypes in the population at the moment of switching (Figure 4B, insets). This property, in turn, depends on the fitness of G330T and H372A while in the CRIPT environment, a factor that heavily favors G330T over H372A (Figure 1B; \( K_{d}^{G330T} = 2.2 \pm 0.33 \mu M \) and \( K_{d}^{H372A} = 26.9 \pm 6.3 \mu M \)). As a consequence, G330T typically comprises the majority of the cryptic genetic variation in the CRIPT environment, more likely to be present and able to support transition to the double-mutant state when the environment switches to T-2F.

How does this result depend on mutation rate and the switching rate of target ligand? Simulations show that conversion to the double-mutant state is sensitive to the rate of mutation and the rate of ligand switching. For example, at a mutation rate of 0.1 per generation, the population is more likely to remain in the wild-type state, whereas at a mutation rate of 10 per generation, the population is more likely to reach the double-mutant state (Figure 4C–E).

Figure 4. The Preferred Path of Adaptation
(A) A population dynamics model for the path of adaptation between the wild-type and G330T, H372A double mutant genotypes. Simulations are initiated with a population of \( N \) wild-type individuals, and, at each generation, single mutations are allowed with rate \( m \) and double mutations with rate \( m^2 \), ligands switch between CRIPT and T-2F every \( r \) generations, and the fitness of each genotype is defined as the fraction bound of ligand. The simulation permits a quantitative analysis of the relative flux through G330T or H372A along the path to the double-mutant state.

(B) One simulation trajectory at a particular mutation and ligand-switching rate (\( N_m = 1 \) and \( r = 500 \)), showing two trials of adaptation in response to switching from the CRIPT ligand (class I) to the T-2F ligand (class II). These examples show that different proportions of the two single mutants can act as intermediates, depending on the pre-existing population of G330T and H372A variants at the moment of ligand switching (insets). Circles mark every 10 generations in main panels and every generation in insets.

(C–E) Histograms of the fraction of G330T (\( n_{G330T}/(n_{G330T} + n_{H372A}) \)), \( n \) represents integrated counts over the period of switching to the double mutant state for ~500 trials of switching from the CRIPT to T-2F ligands. The analysis is shown for three regimes of mutation rate: (1) \( N_m = 0.1 \), where mutations are rare (C), (2) \( N_m = 1 \) (D), and (3) \( N_m = 10 \), where mutations are abundant (E). G330T is always the preferred path of adaptation to the double-mutant state upon ligand switching.

(F) The preference of a hypothetical “G330T” variant (thus in quotation marks) over H372A for adaptation to the T-2F ligand as a function of computationally varying the affinity for the CRIPT ligand from 1 \( \mu M \) to 26.9 \( \mu M \), the same affinity as H372A. Thus, given model parameters, affinities as low as 15 \( \mu M \) can still provide a statistical advantage over H372A in facilitating adaptation (Wilcoxon rank-sum test).

472 Cell 166, 468–480, July 14, 2016
Figure 5. The Structural Basis for Ligand Specificity Switching

Shown are high-resolution crystal structures of wild-type (WT), H372A, G330T, and double mutant variants of PSD95<sup>pdz3</sup>, either unliganded (apo) or bound to CRIPT or T-2F ligands as labeled.

(A) The WT-CRIPT structure, recapitulating features of class I ligand recognition. The threonine hydroxyl of –2 is hydrogen-bonded to histidine 372, and G330 is located on a well ordered β2–β3 loop packed against the region of 372 (G330<sub>1</sub>, the subscript indicates conformation 1).

(B and C) The structures of H372A bound to CRIPT (B) or T-2F (C) show truncation of the 372 side chain and little other conformational change, a local perturbation permitting accommodation of the phenylalanine side chain at –2 without steric clash. The loss of both bulk and hydrogen bonding potential at position 372 is consistent with the partially class-switching phenotype of H372A.

(D) Binding of the T-2F ligand to wild-type PSD95<sup>pdz3</sup> causes rotation of H372 to a new, non-native rotamer state (to prevent steric clash), and induction of two partially occupied conformational states of the β2–β3 (G330<sub>1</sub> and G330<sub>2</sub>).

(E–G) The G330T mutation (apo state) stabilizes the β2–β3 loop in the alternate conformation 2 (E), a state that can permit either rotamer state of H372 without steric clash. Thus, in G330T, the H372 side chain occupies the native rotamer in binding CRIPT (F) and the alternative rotamer in binding T-2F (G).

(legend continued on next page)
double-mutant state is only achieved over a certain regime of ligand-switching rate. This makes sense: if ligand switches too rapidly to permit fixation of the double mutant, the population converges to the only genotype that is fit for the average of both ligand environments—G330T (Figure S3). However, in any regime of ligand-switching rate in which the double mutant fixes in the population, G330T is always more preferred than H372A in mediating adaptation (Figures 4C–4E). This is true when mutations are rare (Nμ < 1; Figure 4C) and when mutations are abundant (Nμ >> 1; Figure 4E). Thus, it is the neutral, class-bridging genotype rather than the direct class-switching genotype that represents the likely intermediate in adaptation to new ligand class specificity.

But how “neutral” does a mutation have to be to be statistically preferred over a class-switching mutant such as H372A in mediating adaptation? Indeed, even G330T is not strictly neutral in the CRIPt environment (K^G330T = 2.2 ± 0.33 μM and K^WT = 0.8 ± 0.09 μM); this is the reason why it is considerably less competitive than the wild-type (Figure 4B). To study this, we carried out a series of simulations in which we examined the effect of varying the affinity of G330T for the CRIPt ligand from near wild-type (1 μM) to that of the class-switching mutant (26.9 μM; Figure 1B). The data show that, given the conditions of the simulation, affinities up to the limit of physiological PDZ binding (<15 μM) will be statistically preferred to H372A (Figure 4F). This result relaxes the notion of conditional neutrality, defining a limit of protein function at which a mutant can still contribute to the cryptic genetic variation and be distinguished in adaptive capacity from direct class-switching mutations.

Structural Basis for Class-Bridging Ligand Binding

The dominance of the class-bridging G330T mutation in adaptation to new ligand specificity is interesting because it is not structurally obvious. Position 330 occurs on a surface loop (β2–β3) that lies behind the substrate binding pocket and makes no direct contact with ligand (Figures 1A and 1C). How does mutation at this distant site create a dual-function PSD95<sup>Δ<sub>2–3</sub></sup> binding pocket capable of recognizing both class I and class II ligands? To address this issue, we solved the high-resolution crystal structures of the four PSD95<sup>Δ<sub>2–3</sub></sup> variants in either the apo (unliganded) state or bound to either CRIPt or T-2F ligands—a total of eight structures (Figure 5; Tables S2 and S3). All structures were solved under near-isomorphous conditions—in the same crystal form (P4₁2₁2₁) with unit cell constants within 0.5% of each other—and models were refined to a similarly high resolution (≤2.0 Å) with excellent statistics and geometry (Tables S2 and S3). Thus, we are in a position to make statements about the mechanism of action of the mutations from a comparative study of atomic structures.

The structure of PSD95<sup>Δ<sub>2–3</sub></sup> bound to the class I CRIPt ligand—the wild-type complex—shows H372 in a rotameric state in which it can hydrogen-bond with T-2 and a well ordered β2–β3 loop that makes backbone hydrogen bonds with the H372 region (Figure 5A; Doyle et al., 1996). Not surprisingly, mutation of position 372 to alanine abrogates class I recognition by eliminating the hydrogen-bonding partner for the Thr/Ser residue at the –2 position (Figure 5B) but also creates space for accommodating a bulky hydrophobic side chain at the –2 position without steric clash (Figure 5C). No other conformational changes are evident, indicating that the direct class-switching phenotype of H372A is due to effects that are spatially localized to the site of adaptive challenge.

The origin of the class-bridging phenotype of G330T is qualitatively different. To explain, consider the effect of the T-2F ligand in binding to wild-type PSD95<sup>Δ<sub>2–3</sub></sup>, a low-affinity complex. Binding of the T-2F ligand involves a propagated structural perturbation in which the side chain of H372 (presumably due to steric clash) is forced to adopt a new rotamer state with two split conformations with roughly equal occupancy, and the β2–β3 loop (containing position 330) is, in turn, induced to partially adopt an alternate conformational state (Figures 5D and 5I). The conformational heterogeneity at both H372 and the β2–β3 loop are consistent with the poor affinity of the wild-type protein for the T-2F ligand (Figure 1B). How does G330T provide for high-affinity binding of both CRIPt and T-2F ligands? The G330T mutation stabilizes the β2–β3 loop in the non-native alternate conformation (Figures 5E and 5I), a structural change that permits the H372 side chain to adopt either rotamer state without steric penalty (Figures 5F and 5G). Thus, the G330T variant can recognize both class I and class II ligands with high affinity, switching the rotameric state of H372 in a ligand-dependent manner (compare Figures 5F and 5G). To complete the path of adaptation, it is straightforward to see that addition of H372A in the background of G330T would (just as in the wild-type background) abrogate class I ligand recognition, resulting in the class II specificity observed in the double mutant (Figure 5H).

In summary, H372A works directly and locally at the binding pocket to simultaneously eliminate class I ligand binding and to promote class II ligand binding—the phenotype of direct switching. In contrast, G330T works allosterically to open up conformational plasticity at the binding pocket, which enables both class I and class II recognition—the phenotype of class-bridging binding. It is important to note that this plasticity does not come in the form of general flexibility of the binding pocket reminiscent of low-affinity, broad-specificity interfaces, such as in the immature, germline isoforms of antibodies (Wedemayer et al., 1997). Instead, it opens up just one additional macroscopic conformational state (Figures 5F and 5G), which results in high-affinity, dual-class ligand specificity (Figure 3B).

Spatial Distribution of Conditional Neutrality

The detailed study of G330T and H372A motivates a comprehensive analysis of all adaptive mutations to define the general structural rules. Such a study is made possible by a

(H) The structure of the G330T,H372A double mutant bound to T-2F is similar to that of H372A alone (C), except that the β2–β3 loop is in conformation 2, consistent with G330T.

(I) Conformational states of the β2–β3 loop in all eight PSD95Δ2–3 variants presented in this work. The loop is in one conformation (conf 1) in the wild-type PSD95Δ2–3-CRIPt, H372A-CRIPt, and H372A-T-2F complexes; in two partially occupied conformations (conf 1 and conf 2) in the WT-T-2F complex; and in conf 2 in all variants in a G330T background. The dashed lines are drawn through the C<sub>α</sub> atom of position 330.
dataset comprising a total saturation mutagenesis of PSD95\textsuperscript{pdz3} (McLaughlin et al., 2012), reporting the effect of every possible amino acid substitution at every position in the PDZ domain (1,598 total) on binding to either the class I CRIPT ligand (ordinate) or the class II T-2F variant (abscissa). The shaded regions describe physiologically significant (\(\leq 15\mu M\)) binding to the T-2F ligand, with either associated loss of function (red) or physiological neutrality (green) for the CRIPT ligand.

(B) Three rotations of PSD95\textsuperscript{pdz3}, with all positions containing adaptive mutations for T-2F in sphere representation with colors as shown in (A). Thus, red spheres correspond to positions with direct switching mutational phenotypes, and green spheres indicate positions with class-bridging phenotypes. The number of mutations at each position with that phenotype are shown in parentheses. The blue mesh indicates the protein sector—the network of coevolving positions in the PDZ family. The data show that nearly all adaptive mutations are contained in the protein sector (marked with asterisks), that direct-switching phenotypes localize to the site of adaptive challenge (ligand – 2), and that class-bridging phenotypes arise from a distributed, contiguous network of residues leading away from the binding pocket through the sector.

Figure 6. Spatial Architecture of Adaptive Mutations in Response to T-2F

(A) The effect of all possible single mutations in PSD95\textsuperscript{pdz3} on binding to either the class I CRIPT ligand (ordinate) or the class II T-2F variant (abscissa). The shaded regions describe physiologically significant (\(\leq 15\mu M\)) binding to the T-2F ligand, with either associated loss of function (red) or physiological neutrality (green) for the CRIPT ligand.

(B) Three rotations of PSD95\textsuperscript{pdz3}, with all positions containing adaptive mutations for T-2F in sphere representation with colors as shown in (A). Thus, red spheres correspond to positions with direct switching mutational phenotypes, and green spheres indicate positions with class-bridging phenotypes. The number of mutations at each position with that phenotype are shown in parentheses. The blue mesh indicates the protein sector—the network of coevolving positions in the PDZ family. The data show that nearly all adaptive mutations are contained in the protein sector (marked with asterisks), that direct-switching phenotypes localize to the site of adaptive challenge (ligand – 2), and that class-bridging phenotypes arise from a distributed, contiguous network of residues leading away from the binding pocket through the sector.
residues linking the class-switching active site residues to class-bridging regions distributed throughout the protein structure (Figures 6B and 7). The network is not isotropically organized in space around the T-2F site in a manner consistent with a simple model of spatial proximity to the site of adaptation. Instead, it is an anisotropic network that fractures through the protein structure to include some distantly positioned residues at the expense of some more proximal ones (Figures 6B and 7). For example, position 373 is in the immediate vicinity of T-2F but shows no mutations capable of adaptation. In contrast, position 362 is nearly 15 Å from ligand position –2 but has four mutations that create a binding pocket capable of recognizing both class I and class II ligands. Thus, the data argue that all class-bridging mutations are fundamentally allosteric in nature, forming specific networks of amino acids within the protein structure to influence active site function from a distance.

The Protein Sector as the Origin of Adaptive Mutations
What principle of protein structure underlies the complex spatial organization of conditionally neutral mutations? The special relevance of this class of mutations for evolutionary dynamics and the finding that they originate from wire-like amino acid networks within the protein structure (Figures 6B and 7) provide an important clue. Previous studies have introduced the concept of “protein sectors,” groups of amino acids that are conserved and coevolve over the long-term evolutionary history of a protein family (Halabi et al., 2009; Smock et al., 2010). Sectors comprise sparse, contiguous networks of amino acids that typically link the protein active site to distantly positioned surface sites (Lockless and Ranganathan, 1999; Suel et al., 2003) and have been linked to the constraints on protein folding (Reynolds et al., 2013; Socolich et al., 2005) and functional properties such as catalysis (Halabi et al., 2009; Reynolds et al., 2011), binding (McLaughlin et al., 2012; Russ et al., 2005), signal transmission (Ferguson et al., 2007; Lee et al., 2008, 2009; Suel et al., 2003), and allosteric regulation (Reynolds et al., 2011; Suel et al., 2003). In short, sectors represent a model for the relevant cooperative action of amino acid positions in proteins.

We compared the pattern of adaptive mutations—both direct class-switching and class-bridging—with the pattern of coevolution in the PDZ domain family (McLaughlin et al., 2012). The sector in the PDZ domain family (blue mesh, Figures 6B and 7) comprises a group of 20 amino acid positions (~20% of total residues, default parameters, SCA5.0) that form a network linking the ligand binding pocket to three regions: the β2–β3 loop, the α1 helix, and the end of the β4 strand (positions 362–363) (Figures S4 and S5B). Essentially all of the adaptive mutations, both directly class-switching and class-bridging, are contained within the sector (38 of 44 mutations at 8 of 12 positions; Figures 6B and 7) (p = 0.001, Fisher’s exact test; Figure S5), and the six remaining mutations occur at four surface sites (358, 333, 371, and 380; marked in yellow in Figure 7) that contact the peripheral edges of the sector. Importantly, keeping the number of top-scoring positions the same as for the sector (i.e., 20), neither spatial proximity to the site of adaptive challenge (p = 0.063, Fisher’s exact test) nor position-specific conservation (p = 0.47, Fisher’s exact test) show such significant correlation with adaptive positions (Figures S5D–S5F). Thus, at least for the primary specificity site, the capacity to adapt in the PDZ domain emerges from an evolutionarily ancient coevolving network of residues. The coevolution of amino acids within the sector is consistent with the cooperativity and allosteric effect of class-bridging mutations and generalizes the role of protein sectors as not only functional units of proteins but as adaptive units of proteins.

DISCUSSION

A Structural Model for Protein Adaptation
The motivation for this work is the concept that evolutionary dynamics place non-trivial constraints on the design of natural proteins. An example of such a constraint is the existence of the class of epistatic mutations termed conditionally neutral—mutations that do not influence the existing functional activity but that open up new activities that can be selected under the right environmental conditions (Hayden et al., 2011; Luria and Delbrück, 1943; Wagner, 2005). Such mutations can contribute to the standing genetic variations in populations and can facilitate the acquisition of new phenotypes as selection conditions fluctuate (Draghi et al., 2010; Hayden et al., 2011). Thus, the elucidation of structural principles of conditional neutrality in proteins is a key next step in understanding their mechanisms and origin.

In this work, we demonstrate the existence, evolutionary relevance, and mechanism of conditionally neutral mutations in a member of the PDZ family of protein interaction modules. The main result is that conditional neutrality is generally allosteric in nature, working from a distance through a network of amino acid interactions to open up new conformational states at the ligand binding pocket. In contrast, adaptive mutations located at the active site have the property of direct switching of ligand class-specificity—new ligand binding is gained at the expense of binding to the existing ligand. Simulations of evolutionary dynamics confirm the notion that it is the class-bridging mutations, not the direct switching ones, that are likely to serve as intermediates in adaptation. Thus, we conclude that, in addition to its contributions to functional properties such as signal transmission and regulation, intramolecular allosteric plays a key role in facilitating the evolutionary process.

Recent high-throughput methods for mutagenesis (Fowler and Fields, 2014; McLaughlin et al., 2012; Stiffler et al., 2015) will facilitate testing of the generality of this conclusion. However, the findings here are consistent with data in at least one other protein system—TEM-1 β-lactamase, an enzyme that confers resistance to specific antibiotics in bacteria (Salverda et al., 2010). Deep mutational scanning reveals a class of mutations underlying adaptation that shows conditional neutrality; that is, neutral with regard to the existing substrate but gain of function toward a new substrate (Stiffler et al., 2015). As in PDS95Daz3, these mutations occur at sites that are distant from the active site, connecting through physically contiguous networks within the protein structure. Interestingly, in TEM-1, distance from the active site correlates with the robustness of neutrality to increasing selection pressure, a finding that might help explain why conditionally neutral mutations form wire-like networks that extend far from the active site.

Taken together, these findings suggest an “outside-in” structural principle for protein adaptation. The idea is that the path of
Figure 7. Relationship of the Protein Sector to Adaptive Positions

(A) A space-filling representation of PSD95pdz3, with the protein sector shown as blue mesh and positions capable of adaptation to the T-2F ligand colored red (direct-switching, in sector), green (class-bridging, in sector), or yellow (class-bridging, non-sector). The data show that the four non-sector adaptive mutations (labeled) occur at surface-exposed sites distant from ligand position –2 (shown as red stick bonds).

(B–G) Serial slices through PSD95pdz3 at the planes indicated in (A). The views are from the left as indicated. The data show that adaptive positions are nearly all contained within the protein sector (overlap of blue mesh with red and green spheres), and the four surface-exposed positions with class-neutral mutations are connected to the peripheral regions of the protein sector. Overall, adaptive positions comprise a wire-like system of physically connected residues that connects the site of adaptive challenge (ligand –2) through the protein structure.
adaptation likely starts from the acquisition of mutations at positions distant from the active site but that are wired up through a pre-existing network of cooperative amino acid interactions. Through action at a distance, these mutations have the capacity for opening up active-site conformational states that can introduce new functional phenotypes without abrogating existing function—the property of conditional neutrality. From a point of view of evolutionary dynamics, the key benefit of such mutations is that they can temporally unlink the appearance of adaptive mutations from environmental fluctuations that alter selection pressures. The neutrality protects against purifying selection and enables adaptive mutations to pre-exist in populations as cryptic variations. In contrast, mutations that cause direct phenotypic switching (even when structurally more obvious) can only support adaptation with temporal coupling of mutation and selection pressures. More generally, we propose that the degree of neutrality of adaptive mutations toward existing functionality will set the timescale (relative to the mutation rate and environmental switching rate) over which it can support adaptation. Thus, conditionally neutral mutations would seem to represent a pool of natural variation that is the engine for the evolution of new phenotypes.

**Implications for Protein Engineering**

The outside-in concept for protein adaptation is interesting because it is essentially opposite to the current practice of structure-guided protein engineering. For example, classic work on switching the primary (P1 site) substrate specificity of the serine protease trypsin to that of chymotrypsin (Hedstrom, 2002; Hedstrom et al., 1994) began with mutations at sites directly contacting the P1 side chain (the S1 pocket). The result was initial loss of enzyme function, a phenotype explained by collapse of the S1 pocket upon mutation (Perona et al., 1995). Successful transfer of specificity required the subsequent addition of mutations at positions distributed through the protein structure, which has the effect of restoring stability (and new functional specificity) to the S1 pocket. Similarly, attempts to switch the activities of type II restriction enzymes (Lukacs et al., 2000; Morgan and Luyten, 2009), transcription factors (Poelwijk et al., 2011), and β-lactamases (Stiffler et al., 2015) show that active site mutations tend to display loss of function, whereas combinations with structurally non-obvious peripheral mutations facilitate acquisition of new function. In light of the work presented here, a useful avenue for protein engineering might be to target mutations not by the principle of spatial proximity to the active site but by the spatial pattern of adaptive mutations.

In this regard, it is interesting that the adaptive mutations, both direct-switching and class-bridging, occur within the network of coevolving positions (the sector) in the PDZ and serine protease families (Halabi et al., 2009). This finding strongly argues that the pattern of adaptive mutations is not merely an idiosyncratic feature of each model system but is, instead, a deeply conserved aspect of the entire protein family that can be predicted through sequence analysis alone. It will be interesting to combine sector predictions and the principle of outside-in mutagenesis to explore new general strategies for the evolution and engineering of new protein functions.

**Origins of Allostery**

The finding that the protein sector contains adaptive mutations offers an interesting hypothesis about the origin of this cooperative internal architecture within protein tertiary structures. Sectors are coevolving units of protein structures and have been associated with various functional properties of proteins—catalysis, binding, and allosteric signaling (Halabi et al., 2009; Reynolds et al., 2011; Smock et al., 2010; Suel et al., 2003). A natural inference might be that the wire-like architecture of sectors, connecting active sites to distant surface sites through the protein core (Figure 7), emerged in evolution as a consequence of selection for the corresponding functional property. However, it is not obvious how such a network of cooperative amino acid interactions could be built through a process of stepwise variation and selection given that intermediate genotypes are not guaranteed to be functional.

The data presented here suggest another model. Sectors are primarily a consequence of a history of adaptation to fluctuating conditions of fitness, with the wire-like distributed architecture evolving simply because conditional neutrality is enabled by non-local allosteric mechanisms. That is, we propose that the origins of allostery lie in evolvability, not in function. According to this model, functional properties that make use of allostery (e.g., long-range regulation and signal transmission) are derivatives that emerge easily at multiple surface sites through engagement of the pre-existing allosteric network. Indeed, experiments suggest that it is possible to naively engineer new allosteric control into proteins through engagement of sector-connected surface sites (Lee et al., 2008; Reynolds et al., 2011). The recent development of techniques for fast continuous evolution of proteins (Esvelt et al., 2011) may help with designing experiments to test these ideas.

**Experimental Procedures**

**Global Analysis of PDZ Ligand Specificity**

The comprehensive study of PDZ binding specificity is made possible by a modified version of a BTH system (McLaughlin et al., 2012) in which transcription of the chloramphenicol acetyltransferase (pZER1RM-CAT plasmid) reporter gene is made quantitatively dependent on the binding between each PDZ domain variant (fused to the bacteriophage λ-c1 DNA binding domain, pZS22 plasmid) and a library of ligands (fused to the N-terminal domain of the Escherichia coli RNA polymerase α subunit [pZA31 plasmid, total theoretical library complexity 20^4 = 160,000] (Figure S1; Table S1)). The details of construction of the ligand library and the BTH assay are given in the Supplemental Experimental Procedures. After selection, cultures were subject to plasmid DNA isolation, PCR amplification of the ligand region of pZA31, and standard preparation for Illumina Hi-Seq 2500 sequencing (University of Texas Southwestern [UTSW] genomics core). Sequencing data were analyzed using home-written codes and MATLAB (MathWorks, available upon request) and used to compute ΔE = log(f̂_u/f̂_s) - log(f̂_u/f̂_u), the enrichment of each ligand x in the selected (s) and unselected (u) libraries relative to a reference sequence o with similar affinity for each PDZ variant. The reference sequence was CRIP1 for wild-type and G330T variants and T-2F for H372A and the double mutant variants.

**Expression and Purification of PSD95Δ32 Proteins**

Wild-type or mutant PSD95Δ32 (amino acid range 297–415) were expressed as glutathione S-transferase (GST)-fusions in Escherichia coli BL21(DE3) cells and purified to near homogeneity through sequential affinity chromatography, cleavage of the GST tag, source 15Q anion exchange chromatography, and...
size exclusion chromatography. Complete details are given in the Supplemental Experimental Procedures. Purified proteins were concentrated to 35 mg/ml and subsequently either flash-frozen in liquid N₂ for storage at −80°C or used immediately for crystallization. Substrate peptides for co-crystallization (CRIP - [acyetyl-TKNYKGTAVG], T-2F - [acyetyl-TKNYKQFSV-COOH]) were synthesized using standard fluorenylmethyloxycarbonyl chloride (Fmoc) chemistry (UTSW Proteomics Core Facility), high-performance liquid chromatography (HPLC)-purified, and lyophilized.

Crystallization and Structure Determination of PSD95\(^{pdz3}\) Variants

Crystallization of PSD95\(^{pdz3}\) variants was performed by the vapor diffusion hanging drop method at 16°C. Details are given in the Supplemental Experimental Procedures, and specific crystallization conditions for each mutant are shown in Table S4. Diamond-shaped crystals appeared either spontaneously or with microseeding after 1–5 days and grew to 100–200 µm in length over several weeks. Single crystals were cryoprotected by serial equilibration into crystallization buffer with increasing amounts of glycerol (up to 25%) and flash-frozen in liquid N₂.

Diffraction data were collected at 100 K either at the UT Southwestern Structural Biology Laboratory or at the Advanced Photon Source (Argonne National Laboratory, 19-ID) and indexed and scaled in HKL-2000 (Otwinski and Minor, 1997) (HKL Research). Resolution cutoffs were chosen based on χ and CC 1/2 (Tables S2 and S3). Phasing and refinement were carried out using PHENIX (Python-based hierarchical environment for integrated xtallography) (Adams et al., 2010) with manual modeling in COOT (crystallographic object-oriented toolkit) (Emsley et al., 2010). The data collection and refinement statistics are summarized in Tables S2 and S3. An initial model was obtained from rigid body and temperature factor refinement using published structures of PSD95\(^{pdz3}\) (PDB: 1BFE and 1B8H, with ligand removed) and subjected to 0.5-Å coordinate randomization followed by Cartesian simulated annealing to minimize phase bias. Further computational refinement steps involved iterative rounds of positional and temperature factor minimization, manual model building, solvent placement, and translation/libration/screw (TLS) refinement, decreased by increase in crystallographic R-factors. Figures were prepared with PyMol (DeLano, 2002). The atomic coordinates and structure factors were deposited in the PDB: 5HEB (PSD95\(^{pdz3}\)(WT)-CRIP), 5HED (PSD95\(^{pdz3}\)(WT)-T-2F), 5HET (PSD95\(^{pdz3}\)(G330T)-apo), 5HEY (PSD95\(^{pdz3}\)(G330T)-CRIP), 5HF1 (PSD95\(^{pdz3}\)(G330T)-T-2F), 5HFB (PSD95\(^{pdz3}\)(H372A)-CRIP), 5HFC (PSD95\(^{pdz3}\)(H372A)-T-2F), and 5HFF (PSD95\(^{pdz3}\)(G330T, H372A)-T-2F).

Computational Simulations

The population dynamics model shown in Figure 4A was simulated using custom-written code developed in MATLAB and executed on a LINUX high-performance computing cluster (BioHPC, UT Southwestern Medical Center). The complete annotated codes and details of the simulation are given in the Supplemental Experimental Procedures. The model simulates the dynamics of a constant-sized population (here, N = 1000, large relative to the number of genotypes) comprising the four PDZ variants (wild-type, G330T, H372A, and the double mutant), with mutation and selection under a fluctuating environment of either CRIP or T-2F ligands. For each trial of switching from CRIP to T-2F in which the double mutant ultimately goes to fixation in the population, we computed the fraction of G330T and H372A in the interval from the switch to fixation of the double mutant. Limits for integration were automatically determined by empirical fitting of the probability density of the double mutant in each trial (see codes). The data in Figures 4C–4F were obtained from ∼500 trials of CRIP1 to T-2F ligand switching each.

Statistical Tests

Fisher’s exact tests of the association between adaptive positions and classification of PSD95\(^{pdz3}\) positions by sector, positional conservation, or spatial proximity were carried out using MATLAB. Positional conservation and sector were defined as reported previously (McLaughlin et al., 2012), and spatial proximity was computed using coordinates from a high-resolution crystal structure of PSD95\(^{pdz3}\) (PDB: 1BFE) (Doyle et al., 1996) as the closest distance between any pair of atoms of ligand T-2 and every other amino acid; see Figure S5 for the contingency matrices.

ACCESSION NUMBERS

The accession numbers for the atomic coordinates and structure factors reported in this paper are PDB: 5HEB (PSD95\(^{pdz3}\)(WT)-CRIP), 5HED (PSD95\(^{pdz3}\)(WT)-T-2F), 5HET (PSD95\(^{pdz3}\)(G330T)-apo), 5HEY (PSD95\(^{pdz3}\)(G330T)-CRIP), 5HF1 (PSD95\(^{pdz3}\)(G330T)-T-2F), 5HFB (PSD95\(^{pdz3}\)(H372A)-CRIP), 5HFC (PSD95\(^{pdz3}\)(H372A)-T-2F), and 5HFF (PSD95\(^{pdz3}\)(G330T, H372A)-T-2F).

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, five figures, and four tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2016.05.047.

AUTHOR CONTRIBUTIONS


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The BTH assay is modified from previous work (Dove et al., 1997; McLaughlin et al., 2012) to give a quantitative readout of the affinity between a PDZ domain and its peptide ligand in the expression of a target gene. The PDZ domain is expressed as a C-terminal fusion with the DNA binding domain of the bacteriophage λ-c1 protein (with the E34P mutation to optimize dynamic range (McLaughlin et al., 2012)). The peptide ligand is expressed as a C-terminal fusion to the N-terminal domain of the E.coli RNA polymerase α-subunit, and the target gene is chloramphenicol acetyl transferase (cat).

A library of potential PDZ ligands (randomized in terminal four residues, \(20^4 = 160,000\) total) is transformed into E.coli MC4100-Z1 cells expressing the other components of the BTH assay, and are selected on chloramphenicol for PDZ function (see methods for details). The ligand library is subject to Illumina HiSeq sequencing to count the frequency of each C-terminal peptide in the library in both the unselected and selected cell populations. As shown for a few examples, legitimate ligands are highly enriched in the selected population while non-binding ligands are absent. Quantitatively, we compute the enrichment of each ligand \(x\) relative to reference ligand \(o\) ("relative enrichment," \(\Delta E\)) according to the equation shown. The reference ligand is chosen for same absolute affinity over PDZ variants to put datasets on the same scale for quantitative comparison; thus, CRIPT for wild-type and G330T and T-2F for H372A and G330T,H372A.

A standard curve showing the relationship of relative enrichment to the equilibrium free energy of binding for a set of 83 single point mutations in the PSD95\(^{pdz3}\) domain to the CRIPT ligand. The experiment shows that the sequencing based assay provides a quantitative measure of ligand binding.
Figure S2. Sequence Profiles of Ligand Peptides Bound by PDZ Variants along the Adaptive Path, Related to Figures 2 and 3

Each panel shows the pattern of amino acid preferences at ligand positions (labeled −3 to 0) as a “sequence logo” plot for the PDZ variant and affinity cutoff indicated. In each panel, the bar graph shows a more detailed, quantitative view of the amino acid profile at position −2, the main specificity determinant. The units of amino acid preference are “bits,” where a frequency of unity for a particular residue would give a bit score of \( \log_2 \) (20). The data permit two conclusions. First, that the amino acid profiles are robust to affinity cutoff. Second, that wild-type PSD95\(^{pdz3}\) (WT) displays class I (S/T) preference at position −2, H372A and the G330T,H372A double mutant display class II (bulky hydrophobic) preference, and G330T displays a class-bridging phenotype, capable of binding both classes of ligands.
Figure S3. The Population Dynamics Model as a Function of $t$, the Ligand Switching Rate, Related to Figure 4

(A) A portion of one simulation trajectory, mutation rate $Nmu = 1$ (meaning production of about one single mutation per generation) and the wait-time for ligand switching $t = 20$ (meaning the ligand switches between CRIP and T-2F every 20 generations). At this fast switching rate, the only genotype that dominates in the population is the one capable of binding both ligands well, G330T.

(B) For comparison, a slower ligand switching rate (here, $t = 500$) permits adaptation to the genotypes that are most fit for the CRIP and T-2F environments (wild-type and double mutant, respectively), with varying proportion of G330T and H372A as intermediates.
Figure S4. The Protein Sector in the PDZ Domain Family, Related to Figures 6 and 7
Statistical coupling analysis (SCA (Halabi et al., 2009; Lockless and Ranganathan, 1999)) reveals a network of coevolving amino acids in the PDZ domain family - a "sector" - that links the ligand binding pocket to a few distant surface sites. Residues G330 and H372 are indicated.
Figure S5. The Relationship of Adaptive Positions to the Protein Sector, Positional Conservation, and Spatial Proximity to the Site of Ligand Variation, Related to Figures 6 and 7

(A) A list of positions in PSD95<sup>pdz3</sup> that contain mutations capable of adaptation to the T-2F ligand (shaded regions, Figure 7). The number of adaptive mutations at each position is indicated (total, 44).

(B) As defined in (McLaughlin et al., 2012), the group of statistically co-evolving positions (the “sector”) in the PDZ protein family comprises 20 positions, indicated at left (shown in Figure S4). For comparison, the middle and right columns show a ranked list of the top 20 conserved positions and the top 20 spatially closest positions to ligand position /C0, respectively. Positional conservation is computed by the Kullback-Liebler relative entropy (Cover and Thomas, 2006), and spatial proximity is measured as a closest distance between atoms comprising the T-2 ligand residue and each residue in PSD95<sup>pdz3</sup>. Yellow highlight indicates a match with an adaptive position.

(C) The fraction of total adaptive positions included in each indicated classification of PSD95<sup>pdz3</sup> positions. These data show that the majority of adaptive mutations occur in the protein sector, and that fewer are accounted for by the other classifications.

(D–F) 2 × 2 contingency matrices describing the relationship between classifications of PSD95<sup>pdz3</sup> positions by adaptive capacity (rows) and by sector, positional conservation, or spatial proximity to the T-2F ligand position (columns). These data permit a Fisher Exact Test, with the null hypothesis that the two classifications are statistically independent. p values show that the null hypothesis is rejected at a p < 0.05 level for the PDZ sector, and is not rejected otherwise. Note that the near-significant p value for spatial proximity makes sense; since physical interactions between atoms arise largely from very short-range forces, we expect that adaptive mutations for T-2F will be enriched in spatially local sites. But the data show that the concept of spatial proximity is incomplete, discounting the adaptive effect of a number of positions (e.g., 330, 362, 322, etc.) that work through allosteric mechanisms. The data suggest that the sector provides a better model for the adaptive effect of mutations by taking such mechanisms into account.
Supplemental Information

Origins of Allostery and Evolvability in Proteins:
A Case Study

Arjun S. Raman, K. Ian White, and Rama Ranganathan
Origins of allostery and evolvability in proteins: Supplementary Information

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Contents

I. Supplementary Experimental Procedures 1
   A. Global analysis of PDZ ligand specificity. 1
   B. Construction of the ligand library 1
   C. Expression and purification of PSD95\textsuperscript{pdz3} proteins. 2
   D. Crystallization and structure determination of PSD95\textsuperscript{pdz3} variants. 2
   E. Computational simulations and codes 3

II. Supplementary Tables 7
   A. TableS1: Sequencing statistics of the ligand library selection experiments 7
   B. TableS2: Crystallographic data collection and refinement statistics - Part 1 8
   C. TableS3: Crystallographic data collection and refinement statistics - Part 2 9
   D. TableS4: Crystallization conditions for PSD95\textsuperscript{pdz3} variants 10

References 11

I. SUPPLEMENTARY EXPERIMENTAL PROCEDURES

A. Global analysis of PDZ ligand specificity.

Comprehensive study of PDZ binding specificity is made possible by a modified version of a bacterial two-hybrid system (McLaughlin et al., 2012) in which transcription of the chloramphenicol acetyl transferase (CAT) reporter gene (pZE1RM plasmid, pRM+ promoter, ampicillin resistant) is made quantitatively dependent on the binding between a PDZ domain (fused to the pRM+ promoter-binding bacteriophage \(^\lambda\)-c1 DNA binding domain, pZS22 plasmid, IPTG inducible, trimethoprim (trm) resistant) and its ligand (fused to the N-terminal domain of \textit{E.coli} RNA polymerase \(\alpha\) subunit, pZA31 plasmid, anhydrotetracycline (aTC) inducible, kanamycin resistant) (Fig. S1). Electrocompentent MC4100-Z1 cells containing pZE1RM-CAT and pZS22-PDZ3 variant plasmids were transformed with 1 \(\mu\)l of 20 ng/\(\mu\)l pZA31-RNA\(\alpha\)-ligand library (see below), recovered for one hour in LB media, grown in 20 \(\mu\)g/ml trm, 50 \(\mu\)g/ml kan, 100 \(\mu\)g/ml amp to OD\(_{550}\) of 0.04, and induced using 50 ng/ml doxycycline plus antibiotics for 3 hours to an OD\(_{550}\) of 0.1. 10ml of the induced culture was used to innoculate 100mL LB + antibiotics as above for selection; the remainder was reserved as the pre-selection population for deep sequencing. Selection was carried out with 150 \(\mu\)g/ml chloramphenicol for 6 hours (taking care that OD\(_{550}\) \(\leq\) 0.1), washed in LB medium, and grown overnight at 37\(^\circ\)C. Both pre- and post-selection cultures were subject to plasmid DNA isolation, PCR amplification of the ligand region of pZA31, and standard preparation for Illumina Hi-Seq 2500 sequencing (UT Southwestern genomics core). Sequencing data were analyzed using home-written codes and MATLAB (Mathworks Inc., provided upon request) and used to compute \(\Delta E_x = \log(f_x^s/f_x^u) - \log(f_x^o/f_x^u)\), the enrichment of each ligand \(x\) in the selected \((s)\) and unselected \((u)\) libraries relative to a reference sequence \(o\) with similar affinity for each PDZ variant. The reference sequence was CRIPT for wild-type and G330T variants, and T-2F for H372A and the double mutant variants.

B. Construction of the ligand library

The library of PDZ ligands (randomized in the C-terminal four amino acid positions, total theoretical library complexity \(20^4 = 160,000\)) was generated as C-terminal fusions with the N-terminal domain of \textit{E.coli} RNA polymerase \(\alpha\) subunit. The library was made using NNS oligonucleotide-directed mutagenesis with a pZA31-RNA\(\alpha\) template containing a non-binding PDZ ligand (N-TKNYKQGGG-COOH) to eliminate background binding. Two oligonucleotides (one sense, one antisense) were synthesized (IDT) with each sequence complementary to 15 base-pairs (bp) on either side but with one oligo containing four consecutive NNS codons at the target positions; \(N\) is a mixture of A, T, C, and G, and \(S\) is a mixture of G and C. This results in 32 codons at each position encoding all 20 amino acids. The
oligos encode a type IIIs restriction site (BsaI), designed to optimize cloning efficiency by enabling a unimolecular ligation protocol. We carried out a single round of PCR, amplifying the entire plasmid while encoding the full library of ligand sequences. This product was subsequently restricted with BsaI, subject to a unimolecular ligation reaction (1 ml, incubated overnight at 16°C), and purified into a final volume of 10 µl (Zymo purification kit). Ten individual transformations into MaxDH10B E. coli (Invitrogen) were made, grown overnight after recovery, and plasmid DNA prepped so as to minimize any possible bottlenecking effect. Transformation of the final library into MC4100-Z1 cells for selection yielded greater than 10^8 transformants, and a near complete representation of the theoretical complexity (Table S1).

C. Expression and purification of PSD95<sup>pdz3</sup> proteins.

pGEX-4T-1 plasmids containing Glutathione-S-transferase (GST)-fusions of wild-type or mutant PSD95<sup>pdz3</sup> (amino acid range 297–415) were transformed into Escherichia coli BL21(DE3) cells and grown overnight on LB plus 100 µg/mL ampicillin (amp) plates. Streaks of colonies were used to start overnight cultures (LB + amp), used to inoculate 1L cultures (Terrific Broth or ZYM-5052 auto-inducing medium (Studier, 2005) + 100 µg/mL amp), and grown to an optical density (600 nm) of 0.6-0.8 at 37°C, induced overnight at 18°C (supplemented with 1 mM isopropyl-D-thiogalactopyranoside if manual induction), and then harvested by centrifugation. Pellets were resuspended in lysis buffer composed of phosphate-buffered saline (PBS: 140 mM NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.8 mM KH<sub>2</sub>PO<sub>4</sub>, pH 7.3) supplemented with 1% glycerol, 1 mg/ml hen egg white lysozyme, 1 mM dithiothreitol (DTT), EDTA-free protease inhibitor cocktail (Roche). The cell suspension was subjected to sonication and centrifugation, and clarified lysate was then incubated with glutathione sepharose 4B resin (GE Healthcare). Bound protein was washed with PBS supplemented with 1% glycerol and 1 mM DTT, and the GST tag was cleaved through bovine thrombin (Calbiochem) proteolysis overnight at room temperature in PBS supplemented with 10% glycerol and 1 mM DTT. Thrombin was removed by benzamidine sepharose (GE Healthcare) and the PDZ domain was purified to near-homogeneity using a Source 15Q anion exchange (GE Healthcare) column employing a linear gradient from low salt (20 mM Tris HCl pH 7.5, 1% glycerol, 1.0 mM DTT) to high salt (20 mM Tris HCl pH 7.5, 1 M NaCl, 1% glycerol, 1 mM DTT). The protein was dialyzed into 10 mM HEPES pH 7.2, 10 mM NaCl, concentrated and subject to size-exclusion chromatography (Superdex 75, GE Healthcare). Peak fractions were pooled, concentrated to 35 mg/mL, and subsequently either flash frozen in liquid N<sub>2</sub> for storage at −80°C or used immediately for crystallization. Substrate peptides for co-crystallization (CDRIPT (Acetyl-TKNYKQFSV-COOH)), T-2F (Acetyl-TKNYKQFSV-COOH)) were synthesized using standard FMOC chemistry (UTSW Proteomics Core Facility), HPLC purified, and lyophilized.

D. Crystallization and structure determination of PSD95<sup>pdz3</sup> variants.

Crystallization of PSD95<sup>pdz3</sup> variants was performed by the vapor diffusion hanging drop method. In all cases, purified protein was diluted to a final concentration of ~9 mg/mL in protein buffer (10 mM HEPES pH 7.2, 10 mM NaCl). Where applicable, peptide was included in protein buffer to a final molar ratio of 2:1 relative to protein. Reservoir solutions typically contained 1 M sodium citrate, pH 7.0; specific crystallization conditions for each mutant are shown in Table S4. Equal amounts (1.5 µl) of protein and reservoir solution were mixed and equilibrated against 500 µl of crystallization buffer at 16°C. Diamond-shaped crystals appeared either spontaneously or with microseeding after 1–5 days and grew to 100–200 µm in length over several weeks. To prepare microseeding solutions, wild-type crystals of the appropriate state were crushed and resuspended in crystallization buffer. Single crystals were cryoprotected by serial equilibration into crystallization buffer with increasing amounts of glycerol (up to 25%) and flash frozen in liquid N<sub>2</sub>.

Diffraction data were collected at 100 K at either at the UT Southwestern structural biology laboratory or at the Advanced Photon Source (Argonne National Laboratory, 19-ID) and indexed and scaled in HKL-2000 (Otwinowski and Minor, 1997) (HKL Research). Resolution cutoffs were chosen based on I/σ and CC 1/2 (Tables S2-S3). Phasing and automated refinement was carried out using PHENIX (Adams et al., 2010) with manual modeling in COOT (Emsley et al., 2010); the data collection and refinement statistics are summarized in Tables S2-S3. An initial model was obtained from rigid body and temperature factor refinement using published structures of PSD95<sup>pdz3</sup> (PDB 1BFE and 1BE9, with ligand removed), and subject to 0.5 Å coordinate randomization to help reduce phase bias. Further computational refinements steps involved iterative rounds of positional and temperature factor minimization, manual model building, solvent placement, and TLS refinement, guided by decrease in crystallographic R-factors. Figures were prepared with PyMol (DeLano, 2002). The atomic coordinates and structure factors have been deposited in the Protein Data Bank with the following acces-
E. Computational simulations and codes

The model shown in Fig. 4 and Fig. S3 simulates the dynamics of a constant sized population (here, \(N = 1000\), large relative to the number of genotypes) comprising the four PDZ variants (wild-type, G330T, H372A, and the double mutant) with mutation and selection under a fluctuating condition of fitness. In each generation, single mutations occur with probability \(\mu\), double mutations with probability \(\mu^2\), and selection re-draws the frequency of each genotype according to its ability to bind ligand relative to all other available genotypes. The fractional binding of each genotype is determined from the experimentally measured equilibrium dissociation constants (Fig. 1B), and the ligand (CRIPT or T-2F) switches every \(\tau\) generations. For each trial of switching from CRIPT to T-2F in which the double mutant ultimately goes to fixation in the population, we compute the fraction of G330T and H372A in the interval from the switch to fixation of the double mutant; limits for integration were automatically determined by empirical fitting of the probability density of the double mutant in each trial (see codes). The data in Figs. 4C-F were obtained from \(~500\) trials of CRIPT to T-2F ligand switching each. The codes were executed using a custom shell script on a high-performance LINUX cluster (BioHPC, UT Southwestern Medical Center).

```matlab
function [out] = evSim_RWR(pinit, mu, tau, nswitches,Kds,switch_mode)
% [sim]=evSim_RWR([1000;0;0;0],mu,tau,nswitches, switch_mode,f);

% The population dynamics model in Raman et al, "Origins of allosterery and
evolvability in proteins: a case study". This function models the
% dynamics of population shifts in WT, G330T, H372A, and the double mutant
% given an initial population structure, mutation rate, ligand switching
% time, and other parameters, described below.
% Inputs:
% (1) "pinit" is the initial population vector, in order wild-type,
% G330T, H372A, and the double mutant. For example, pinit=[1000;0;0;0] to
% start with 1000 WT individuals. sum(pinit) gives the total population
% size, \(N\). In paper pinit = [1000;0;0;0].
% (2) "mu" is the mutation rate - the probability of a single mutation at
% each generation. So, for example, \(mu=0.001\) gives \(N*mu=1\). The paper
% describes simulations at \(mu = 0.0001, 0.001,\) and 0.01.
% (3) "tau" is the wait time in generations for ligand switching. So,
% \(tau=100\) means the ligand switches every 100 generations.
% (4) "nswitches" is total number of ligand switches, and so
% nswitches=tau is the total number of generations simulated.
% (5) "Kds" is a 2 X 4 matrix of equilibrium dissociation constants,
% assumed in units of micromolar. The columns correspond to WT, G330T,
% H372A, G330T,H372A in order, and rows to CRIPT or T-2F ligand in
% order. Values are given in Fig. 1B. Kds(:,1)=[0.8 2.2 26.9 22.1] and
% Kds(:,2)=[36 1.8 1.9 0.5].
% (6) "switch_mode" is a flag that determines whether ligand switching is
% regular (every \(tau\) generations, switch mode=0), or is Poisson
% distributed with a mean wait time of \(tau\) generation.
% (switch_mode="Po". Default switch_mode=0.
% Outputs:
% "out" is a structure with two fields...out.output has all the inputs and
% out.output has "F", the population vector at each generation,
% "gen_switch", the generation at which each ligand switch happens
% (always starts with CRIPT).

% ******************************************************
% ******************************************************
% Step 1: Set Inputs
% ******************************************************
if nargin>6
  switch_mode=0;
end
```

The function `evSim_RWR` simulates the dynamics of a constant sized population. The inputs include `pinit` for the initial population structure, `mu` for the mutation rate, `tau` for the wait time between ligand switches, `nswitches` for the total number of ligand switches, `Kds` for the equilibrium dissociation constants, and `switch_mode` for the mode of ligand switching (regular or Poisson distributed). The code is designed to model the dynamics of population shifts in WT, G330T, H372A, and the double mutant given an initial population structure, mutation rate, and ligand switching times. The outputs include the population vector at each generation and the generation at which each ligand switch happens. The code is executed using a custom shell script on a high-performance LINUX cluster (BioHPC, UT Southwestern Medical Center).
```
if size(Kds,1)==2; Kds=Kds'; end  \% in case Kds matrix needs transposition
f=10./(10*Kds);

% *************************************************
% Step 2: Initialization
% *************************************************
len=(nswitches+1)*tau; \% set number of generations

% Defines svect, which stores the generations at which ligands switch.
% Regular switching every tau generations if switch_mode=0 and Poissonian
% switching otherwise.
svect=zeros(1,len);
if switch_mode==0
    svect((1:tau+len))=1; svect(1)=0;
else
    draws=round(exprnd(tau,1,nswitches));
    index=0;
    for i=1:numel(draws)
        svect(draws(i)+index)=1;
        index=index+draws(i);
    end
    svect(1)=0;
end
% set initial ligand environment, 1 is CRIPIT, -2 is T-2F
env=1; sel=1;

% initialize population vector (numbers of each species, a 4 X 1 vector)
% and individual vector (the actual members of the population, labelled
% according to type, a 1000 X 1 vector).
P=zeros(len);
P(:,1)=pinit;
psize=sum(pinit); \% total population size
labels = [0 1 2 3]; \% the genotype (labels) in decimal. 0=WT, 1=G330T, 2=H372A, 3=the double mutant
f_vect=pinit./sum(pinit); \% frequencies
i_vect=(labels==endr miał(1,f_vect,psize))'; \% make initial individual vector from frequencies (in decimal)
m_vect=dec2bin(i_vect(:,1),2); \% binary version of the population vector
m_vect=[str2num(m_vect(:,1)) str2num(m_vect(:,2))]; \%change from char to num. This is the initial population vect

% *************************************************
% Step 3: Simulation
% *************************************************

% The idea is to pick an environment (that defines the selections),
% randomly draw single and double mutations using binary representation,
% convert to decimal population, apply selection to adjust frequencies,
% redraw population, convert to binary representation, and iterate. Very
% likely more efficient ways to do this...

for i=2:len
    if svect(i)==1
        env=env; \% switch ligands if at the generation specified in svect
        sel=sel-env;
    end
    f_vect=(f(:,sel))./(sum((P(:,1-i)./psize).*f(:,sel))); \% convert fraction bound to relative fitness
    m_vect=abs((poissrnd(mu,2,psize)==0)-m_vect); \% apply random mutation given mu
    p_vect=bin2dec(num2str(m_vect)); \% switch to decimal
    P_tmp=[numel(find(p_vect==0));numel(find(p_vect==1));numel(find(p_vect==2));numel(find(p_vect==3))]; \% get
    f_vect=(f_vect.*P_tmp)./(sum(f_vect.*P_tmp)); \%get frequencies with applying selection on species in
    i_vect=(labels==endr miał(1,f_vect,psize))'; \% generate new population, now after mutation and selection
    m_tmp=dec2bin(i_vect(:,2),2); \% back to binary matrix
    m_vect=[str2num(m_vect(:,1)) str2num(m_vect(:,2))]; \% switch from char to num
    P(:,1)=numel(find(i_vect==0));numel(find(i_vect==1));numel(find(i_vect==2));numel(find(i_vect==3)); \%Write
    new population vector after mutation and selection
end

% *************************************************
% Step 4: Make output structure
% *************************************************

% first, we determine the generations at which CRIPIT changes to T-2F (every
% other one)
gen_switch=gen_switch+1;\% gen_switch=
```

out.input.tau = tau;
out.input.nswitches = nswitches;
out.input.f = f;
out.input.switch_mode = switch_mode;
out.output.P = P;
out.output.gen_switch = gen_switch;

% ***********************************
% Step 5: Analysis
% ***********************************

% This is optional, and is best avoided for high-throughput study of many
% conditions. Comment out below if not desired.

% a plot of the population dynamics over the simulation length
h_sim=figure;clf;hold on;grid on;
plot(P(:,1,:),'b','LineWidth',1.6);
plot(P(2,:),'g','LineWidth',1.6);
plot(P(3,:),'r','LineWidth',1.6);
plot(P(4,:),'k','LineWidth',1.6);
plot(svect*Sparse,'--k');
hold off;
% analysis of the simulation. Here, we extract the population dynamics
% following each trial of ligand switching (CRIPIT to T-2F), isolate the
% events in which the double mutant goes to near fixation, determine the
% interval over which the intermediate genotypes should be integrated, and
% compute the fractional flux through the G330T state.

% pre-allocate variables
events_post=zeros(4,tau,nnumel(gen_switch)); % population dynamics in each trial
lim=zeros(1,nnumel(gen_switch)); % limit for integration in each trial
Npost=zeros(2,nnumel(gen_switch)); % integrated counts of G330T and H372A per trial
frac_G330T=zeros(1,nnumel(gen_switch)); % fraction of G330T per trial
count_double=0;

for k=1:nnumel(gen_switch)
    events_post(:,k,i)=(.,gen_switch(k),gen_switch(k):tau-1));
    if max_events_post(4,:)>800 % minimal double mutant level to consider trial (arbitrary)
        count_double=count_double+1;
        ydat=smooth(diff(events_post(4,:)),20);
        xdat=[1:nnumel(ydat)];
        try
            pd=fit(ydat',diff(events_post(4,:))','gauss1'); % Gaussian fit of double mutant population
            lim(k)=pd.bi*(2*pd.cl); % limit for integrating G330T, H372A populations
            if lim(k)<tau
                Npost(1,k)=numel(events_post(2,1:floor(lim(k),k),2)) % Number G330T
                Npost(2,k)=numel(events_post(3,1:floor(lim(k),k),2)) % Number H372A
                frac_G330T(k)=Npost(1,k)/(Npost(1,k)+Npost(2,k)); % the fraction G330T
            else
                frac_G330T(k)=Inf; % if limit is not less than tau
            end
        catch
            frac_G330T(k)=Inf; % if fitting throws an exception
        end
    end
end
end
end
end
end

% clean up events for exceptions
ind=find(~issubf(frac_G330T) & ~issubf(frac_G330T) & frac_G330T==0 & frac_G330T==Inf); % the indices of valid trials
frac_G330T_clean=frac_G330T(ind); % clean frac_G330T for valid switching trials
lim_clean=lim(ind); % clean lim for valid switching trials
mean_frac_G330T=mean(frac_G330T_clean); % mean fraction G330T over simulation
sem_frac_G330T=std(frac_G330T_clean)/sqrt(nnumel(ind)); % standard error of the mean, since we want confidence in
the mean value, and not the scatter over trials.

% plotting
h_analysis=figure;clf;
plot(frac_G330T_clean,'ok', 'LineWidth',1.5);hold on;grid on
plot([0 nnumel(gen_switch)], [mean_frac_G330T mean_frac_G330T], 'or', 'LineWidth',1);
if isempty(frac_G330T_clean)
text(1,.05,mean((frac_G330T_clean)),['fraction G330T = ', num2str(mean_frac_G330T)],'FontWeight','bold','FontSize',12);
end
axis([0 nnumel(gen_switch) 0 1]);
P_double=count_double/nnumel(gen_switch);
sem_frac_G330T mean_frac_G330T P_double
end
% Step 6: Add analysis to output structure
% ************************************************
out.analysis.frac_G330T=frac_G330T_clean;
out.analysis.index_valid=ind;
out.analysis.integration_limits=lin_clean;
end
### II. SUPPLEMENTARY TABLES

#### TABLE S1
**Sequencing statistics of the ligand library selection experiments.** Related to Figures 2-3. Sequencing statistics from Illumina HiSeq2500 runs for the unselected and selected populations of peptide libraries for wild-type, G330T, H372A, and the double mutant experiments. The unselected populations were combined over all experiments. The total number of reads, \(1.07 \times 10^8\) represented approximately 97% coverage of all peptides in the library.

<table>
<thead>
<tr>
<th></th>
<th>Number of reads</th>
<th>Number of Ligands</th>
<th>Number of Ligands (&gt; 50 counts)</th>
<th>Number of Ligands Bound by Protein (&gt; 15 (\mu)M)</th>
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<tr>
<td><strong>Total Input Library</strong></td>
<td><strong>1.07 \times 10^8</strong></td>
<td>154,521</td>
<td>96.7</td>
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<tr>
<td><strong>Unselected Library Statistics</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>Selected Library Statistics</strong></td>
<td></td>
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<tr>
<td>WT</td>
<td>46,598,840</td>
<td>56,640</td>
<td>55,278</td>
<td>185</td>
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<tr>
<td>G330T</td>
<td>51,195,397</td>
<td>90,735</td>
<td>83,056</td>
<td>846</td>
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<tr>
<td>H372A</td>
<td>29,419,042</td>
<td>59,935</td>
<td>43,488</td>
<td>1051</td>
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<tr>
<td>H372A/G330T</td>
<td>48,989,769</td>
<td>123,295</td>
<td>86,255</td>
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<td>Crystallographic data collection and refinement statistics - Part 1</td>
<td>WT-CRIPT</td>
<td>WT-T_F</td>
<td>G230T-Apo</td>
<td>G230T-CRIPT</td>
</tr>
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<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
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<tr>
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</tr>
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<td>5HED</td>
<td>5HE7</td>
<td>5HEY</td>
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<td>Wavelength (Å)</td>
<td>1.54178</td>
<td>1.54178</td>
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<td>0.97937</td>
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<tr>
<td>Resolution range (Å)</td>
<td>40.15−1.65 (1.709−1.65)</td>
<td>31.66−1.7 (1.761−1.7)</td>
<td>36.52−2.001 (2.073−2.001)</td>
<td>27.21−1.5 (1.554−1.5)</td>
</tr>
<tr>
<td>Space group</td>
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<td>P4_32</td>
<td>P4_32</td>
<td>P4_32</td>
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<td>Unit cell (Å, °)</td>
<td>89.771 89.771 89.771 90 90 90</td>
<td>89.551 89.551 89.551 90 90 90</td>
<td>89.445 89.445 89.445 90 90 90</td>
<td>90 90 90 90 90 90</td>
</tr>
<tr>
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<td>87993</td>
<td>45447</td>
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<tr>
<td>Unique reflections</td>
<td>15083 (1156)</td>
<td>13972 (1259)</td>
<td>8697 (805)</td>
<td>25425 (1645)</td>
</tr>
<tr>
<td>Multiplicity</td>
<td>12.9 (2.5)</td>
<td>9.2 (3.0)</td>
<td>10.0 (8.4)</td>
<td>21.9 (14.7)</td>
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<tr>
<td>Completeness</td>
<td>0.98</td>
<td>0.99</td>
<td>0.99</td>
<td>0.98</td>
</tr>
<tr>
<td>%</td>
<td>Mean I/σ</td>
<td>61.571 (2.478)</td>
<td>60.017 (2.627)</td>
<td>35.095 (2.077)</td>
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<td>R-merge</td>
<td>0.034 (0.389)</td>
<td>0.031 (0.302)</td>
<td>0.064 (N/A)</td>
<td>0.119 (N/A)</td>
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<tr>
<td>R-meas</td>
<td>0.034 (0.333)</td>
<td>0.033 (0.361)</td>
<td>0.068 (N/A)</td>
<td>0.112 (N/A)</td>
</tr>
<tr>
<td>R-pim</td>
<td>0.008 (0.199)</td>
<td>0.010 (0.193)</td>
<td>0.022 (0.397)</td>
<td>0.035 (0.665)</td>
</tr>
<tr>
<td>CC1/2</td>
<td>0.87</td>
<td>0.805</td>
<td>0.713</td>
<td>0.555</td>
</tr>
<tr>
<td><strong>Refinement</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reflections used in refinement</td>
<td>15082 (1156)</td>
<td>13971 (1259)</td>
<td>8696 (805)</td>
<td>25425 (1645)</td>
</tr>
<tr>
<td>Reflections used for R-free</td>
<td>1509 (116)</td>
<td>1398 (130)</td>
<td>871 (80)</td>
<td>2124 (145)</td>
</tr>
<tr>
<td>R-work</td>
<td>0.1666 (0.2273)</td>
<td>0.1704 (0.2291)</td>
<td>0.1837 (0.2369)</td>
<td>0.1724 (0.2705)</td>
</tr>
<tr>
<td>R-free</td>
<td>0.1945 (0.2745)</td>
<td>0.1973 (0.2694)</td>
<td>0.2150 (0.2586)</td>
<td>0.3826 (0.2765)</td>
</tr>
<tr>
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<td>1054</td>
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<td>1110</td>
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<td>RMS (bonds)</td>
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<td>99</td>
<td>98</td>
<td>95</td>
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<td>Ramachandran allowed (%)</td>
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<td>0.72</td>
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<td>0</td>
<td>0</td>
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<td>Rotamer outliers (%)</td>
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<td>N/A</td>
<td>N/A</td>
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<td>Solvent</td>
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<td>29.79</td>
<td>38.7</td>
<td>31.27</td>
</tr>
<tr>
<td>Number of TLS groups</td>
<td>3</td>
<td>3</td>
<td>1</td>
<td>16</td>
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</table>

TABLE S2 Crystallographic data collection and refinement statistics - Part 1. Related to Figure 5. All proteins were crystallized in the same space group (P4_32) and showed unit cell constants within 0.5% of each other. Ligand-bound datasets were collected to a resolution higher than 2.0 Å.
<table>
<thead>
<tr>
<th></th>
<th></th>
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<th></th>
<th></th>
</tr>
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<tbody>
<tr>
<td>PDB ID</td>
<td>5HE1</td>
<td>5HEB</td>
<td>5HFC</td>
<td>5HEF</td>
</tr>
<tr>
<td>Source</td>
<td>APS 19-ID</td>
<td>UTSW SBL</td>
<td>UTSW SBL</td>
<td>APS 19-ID</td>
</tr>
<tr>
<td>Wavelength (Å)</td>
<td>0.97918</td>
<td>1.54178</td>
<td>1.54178</td>
<td>0.97918</td>
</tr>
<tr>
<td>Resolution range (Å)</td>
<td>40.13–1.747 (1.81–1.747)</td>
<td>40.67–1.617 (1.675–1.617)</td>
<td>36.52–1.851 (1.918–1.851)</td>
<td>36.57–1.749 (1.812–1.749)</td>
</tr>
<tr>
<td>Space group</td>
<td>P4_32</td>
<td>P4_32</td>
<td>P4_32</td>
<td>P4_32</td>
</tr>
<tr>
<td>Unit cell (Å, °)</td>
<td>89.724 89.724 89.724 90 90 90 89.592 89.592 89.592 90 90 90 89.445 89.445 89.445 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90</td>
<td>89.592 89.592 89.592 90 90 90 89.445 89.445 89.445 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90</td>
<td>89.592 89.592 89.592 90 90 90 89.445 89.445 89.445 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90</td>
<td>89.592 89.592 89.592 90 90 90 89.445 89.445 89.445 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90 89.575 89.575 89.575 90 90 90</td>
</tr>
<tr>
<td>Total reflections</td>
<td>358023</td>
<td>203441</td>
<td>68030</td>
<td>264373</td>
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<tr>
<td>Unique reflections</td>
<td>13007 (1237)</td>
<td>15108 (556)</td>
<td>10759 (114)</td>
<td>12879 (1179)</td>
</tr>
<tr>
<td>Multiplicity</td>
<td>27.3 (26.3)</td>
<td>13.3 (1.6)</td>
<td>6.3 (2.5)</td>
<td>20.3 (20.1)</td>
</tr>
<tr>
<td>Completeness</td>
<td>0.99</td>
<td>0.93</td>
<td>0.98</td>
<td>0.99</td>
</tr>
<tr>
<td>Mean I/σ(I)</td>
<td>67.804 (1.844)</td>
<td>65.515 (0.936)</td>
<td>35.543 (1.000)</td>
<td>50.222 (1.826)</td>
</tr>
<tr>
<td>Wilson B-factor</td>
<td>19.04</td>
<td>15.74</td>
<td>21.65</td>
<td>18.64</td>
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<tr>
<td>Rmerge</td>
<td>0.055 (N/A)</td>
<td>0.035 (0.338)</td>
<td>0.040 (0.448)</td>
<td>0.063 (N/A)</td>
</tr>
<tr>
<td>Rfree</td>
<td>0.056 (N/A)</td>
<td>0.036 (0.470)</td>
<td>0.044 (0.547)</td>
<td>0.065 (N/A)</td>
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<tr>
<td>R-factor</td>
<td>0.013 (0.381)</td>
<td>0.009 (0.325)</td>
<td>0.017 (0.304)</td>
<td>0.016 (0.478)</td>
</tr>
<tr>
<td>CC1/2</td>
<td>0.721</td>
<td>0.695</td>
<td>0.689</td>
<td>0.766</td>
</tr>
<tr>
<td>Refinement</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Reflections used in refinement</td>
<td>13006 (1217)</td>
<td>15109 (556)</td>
<td>10759 (114)</td>
<td>12880 (1179)</td>
</tr>
<tr>
<td>Reflections used for R-free</td>
<td>1301 (123)</td>
<td>1513 (56)</td>
<td>1078 (97)</td>
<td>1285 (114)</td>
</tr>
<tr>
<td>R-work</td>
<td>0.1993 (0.2911)</td>
<td>0.1751 (0.3004)</td>
<td>0.1635 (0.2094)</td>
<td>0.1740 (0.2314)</td>
</tr>
<tr>
<td>R-free</td>
<td>0.2179 (0.3097)</td>
<td>0.2033 (0.3281)</td>
<td>0.2051 (0.3146)</td>
<td>0.2233 (0.2761)</td>
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<tr>
<td>Number of non-hydrogen atoms</td>
<td>1182</td>
<td>1204</td>
<td>1161</td>
<td>1191</td>
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<td>Macromolecules</td>
<td>1080</td>
<td>1055</td>
<td>1037</td>
<td>1080</td>
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<tr>
<td>Ligands</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>12</td>
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<tr>
<td>Protein residues</td>
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<td>123</td>
<td>124</td>
<td>128</td>
</tr>
<tr>
<td>RMS (bonds)</td>
<td>0.006</td>
<td>0.006</td>
<td>0.012</td>
<td>0.011</td>
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<tr>
<td>RMS (angles)</td>
<td>0.94</td>
<td>0.95</td>
<td>1.46</td>
<td>1.27</td>
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<tr>
<td>Ramachandran favored (%)</td>
<td>99</td>
<td>99</td>
<td>95</td>
<td>97</td>
</tr>
<tr>
<td>Ramachandran allowed (%)</td>
<td>0.73</td>
<td>0.73</td>
<td>4.5</td>
<td>0.71</td>
</tr>
<tr>
<td>Ramachandran outliers (%)</td>
<td>0</td>
<td>0</td>
<td>0.76</td>
<td>2.1</td>
</tr>
<tr>
<td>Rotamer outliers (%)</td>
<td>4.4</td>
<td>13.3 (1.6)</td>
<td>2.7</td>
<td>6.2</td>
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<tr>
<td>Clashscore</td>
<td>4.63</td>
<td>1.91</td>
<td>10.18</td>
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<td>Average B-factor</td>
<td>28.71</td>
<td>22.31</td>
<td>30.76</td>
<td>25.02</td>
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<tr>
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<td>20.99</td>
<td>30.12</td>
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<td>Ligands</td>
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<td>N/A</td>
<td>N/A</td>
<td>57.25</td>
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<tr>
<td>Solvent</td>
<td>36.72</td>
<td>31.72</td>
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<td>31.53</td>
</tr>
<tr>
<td>Number of TLS groups</td>
<td>5</td>
<td>5</td>
<td>8</td>
<td>7</td>
</tr>
</tbody>
</table>

**Table S3 Crystallographic data collection and refinement statistics - Part 2.** Related to Figure 5. All proteins were crystallized in the same space group (P4_32) and showed unit cell constants within 0.5% of each other. Ligand-bound datasets were collected to a resolution higher than 2.0 Å.
<table>
<thead>
<tr>
<th>Protein</th>
<th>[sodium citrate] (M)</th>
<th>crystallization buffer pH</th>
<th>[protein] (mg/ml)</th>
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<tr>
<td>WTapo</td>
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<tr>
<td>WT,CRIPW</td>
<td>1.0</td>
<td>7.0</td>
<td>9</td>
</tr>
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<td>WT,T-2F</td>
<td>1.125</td>
<td>7.1</td>
<td>9</td>
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<td>G330Ta apo</td>
<td>1.05</td>
<td>7.0</td>
<td>7</td>
</tr>
<tr>
<td>G330T,CRIPW</td>
<td>1.2</td>
<td>7.0</td>
<td>8</td>
</tr>
<tr>
<td>G330T,T-2F</td>
<td>1.2</td>
<td>6.8</td>
<td>9</td>
</tr>
<tr>
<td>H372Aapo</td>
<td>0.95</td>
<td>7.0</td>
<td>9</td>
</tr>
<tr>
<td>H372A,CRIPW</td>
<td>1.05</td>
<td>7.0</td>
<td>7</td>
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<tr>
<td>H372A,T-2F</td>
<td>1.05</td>
<td>7.0</td>
<td>7</td>
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<tr>
<td>G330T,H372Aapo</td>
<td>1.0</td>
<td>7.0</td>
<td>13</td>
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<tr>
<td>G330T,H372A,CRIPW</td>
<td>1.25</td>
<td>7.0</td>
<td>9</td>
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<tr>
<td>G330T,H372A,T-2F</td>
<td>1.2</td>
<td>6.75</td>
<td>7</td>
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</table>

TABLE S4 Crystallization conditions for PSD95<sup>PDZ3</sup> variants. Related to Figure 5. Details of protein expression, purification, and general aspects of crystallization are given in the methods section.
References