Coupling of Conformational Transitions in the N-terminal Domain of the 51-kDa FK506-binding Protein (FKBP51) Near Its Site of Interaction with the Steroid Receptor Proteins

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Background: FK506-binding protein 51 (FKBP51) inhibits and FKBP52 stimulates transcription by various steroid receptors. Exchange of a single residue largely reverses this pattern of regulation.

Results: Unlike FKBP52, FKBP51 has two distinct coupled conformational transitions surrounding this mutation site.

Conclusion: Structural analysis of FKBP51 transient states can inform inhibitor design by conformational selection.

Significance: The differential plasticity of the FKBP domains may underlie their differential regulation.

The FK506-binding protein of 51-kDa (FKBP51) and FKBP52 proteins, respectively, has been reported to largely reverse the inhibitory (FKBP51) or stimulatory (FKBP52) effects of these co-chaperones on the transcriptional activity of glucocorticoid and androgen receptor-protein complexes. Previous NMR relaxation studies have identified exchange line broadening, indicative of submillisecond conformational motion, throughout the β₁-β₂ loop in the FK1 domain of FKBP51, which are suppressed by the FKBP52-like L119P substitution. This substitution also attenuates exchange line broadening in the underlying β₃ and β₃a strands that is centered near a bifurcated main chain hydrogen bond interaction between these two strands. The present study demonstrates that these exchange line broadening effects arise from two distinct coupled conformational transitions, and the transition within the β₃ and β₃a strands samples a transient conformation that resembles the crystal structures of the selectively inhibited FK1 domain of FKBP51 recently reported. Although the crystal structures for their series of inhibitors were interpreted as evidence for an induced fit mechanism of association, the presence of a similar conformation being significantly populated in the unliganded FKBP51 domain is more consistent with a conformational selection binding process. The contrasting reduced conformational plasticity of the corresponding FK1 domain of FKBP52 is consistent with the current model in which FKBP51 binds to both the apo- and hormone-bound forms of the steroid receptor to modulate its affinity for ligand, whereas FKBP52 binds selectively to the latter state.

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The atomic coordinates and structure factors (code 4R0X) have been deposited in the Protein Data Bank [http://wwpdb.org/].

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2The abbreviations used are: FKBP51, FK506-binding protein of 51-kDa; FKBP52, FK506-binding protein of 52-kDa; FK506, immunosuppressive drug primarily used in organ transplant therapies; FK1, first FKBP-binding domain; PDB, Protein Data Bank.

Interchanging Leu-119 for Pro-119 at the tip of the β₃-β₄ loop in the first FK506 binding domain (FK1) of the FKBP51 and FKBP52 proteins, respectively, has been reported to largely reverse the inhibitory (FKBP51) or stimulatory (FKBP52) effects of these co-chaperones on the transcriptional activity of glucocorticoid and androgen receptor-protein complexes. Previous NMR relaxation studies have identified exchange line broadening, indicative of submillisecond conformational motion, throughout the β₁-β₂ loop in the FK1 domain of FKBP51, which are suppressed by the FKBP52-like L119P substitution. This substitution also attenuates exchange line broadening in the underlying β₃ and β₃a strands that is centered near a bifurcated main chain hydrogen bond interaction between these two strands. The present study demonstrates that these exchange line broadening effects arise from two distinct coupled conformational transitions, and the transition within the β₃ and β₃a strands samples a transient conformation that resembles the crystal structures of the selectively inhibited FK1 domain of FKBP51 recently reported. Although the crystal structures for their series of inhibitors were interpreted as evidence for an induced fit mechanism of association, the presence of a similar conformation being significantly populated in the unliganded FKBP51 domain is more consistent with a conformational selection binding process. The contrasting reduced conformational plasticity of the corresponding FK1 domain of FKBP52 is consistent with the current model in which FKBP51 binds to both the apo- and hormone-bound forms of the steroid receptor to modulate its affinity for ligand, whereas FKBP52 binds selectively to the latter state.

The FK506-binding protein of 51-kDa (FKBP51)² and FKBP52 are two closely homologous proteins, each containing a pair of FK506-binding domains (FK1 and FK2) followed by a tetratricopeptide repeat domain that mediates their interactions with Hsp90. FKBP52 was first characterized as a co-chaperone of Hsp90 in the activated hormone-bound steroid receptor complexes that are formed with the progesterone, androgen, or the glucocorticoid receptor proteins (1). FKBP52 enhances the binding of the glucocorticoid receptor complex to the dynein motor protein, thus facilitating microtubular transport of the receptor from the cytosol to the nucleus (2). FKBP51 is the predominant Hsp90 co-chaperone for the unliganded state of the glucocorticoid receptor (3, 4), which induces a lower binding affinity for glucocorticoids (5) and dynein (6). Steroid hormone binding to the FKBP51-bound glucocorticoid receptor is believed to induce the release of FKBP51 and the subsequent binding of FKBP52 (7).

The hormone-bound glucocorticoid receptor strongly induces the fkbp5 gene encoding FKBP51, yielding a direct negative feedback control loop (8). Presumably acting via the glucocorticoid receptor, single nucleotide polymorphisms in the fkbp5 gene exhibit a strong correlation with recurrence of depressive episodes, the rate of response to antidepressant therapies, and in psychological stress disorders (9, 10). As a necessary chaperone for the Akt-specific phosphatase PHLPP (11, 12), FKBP51 also provides indirect feedback regulation by inhibiting glucocorticoid receptor phosphorylation via the Akt-p38 kinase pathway (13, 14).

The FK1 domain mediates the selectivity of interaction for steroid receptor binding exchange (15, 16), dynein binding (6), and inhibition of the Akt kinase (12). The peptidylprolyl isomerase activity of this domain is not required for receptor binding exchange (16) or Akt kinase inhibition (12). Using both

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yeast heterologous expression and murine FKBP52 knock-out experiments, Smith and colleagues (16) demonstrated that a FKBP52-like L119P mutation near the tip of the β1-β2 loop in the FK1 domain of human FKBP51 increases reporter gene expression by ∼3.5-fold for the human glucocorticoid and androgen receptors. The complementary P119L mutation in FKBP52 yielded a ∼2-fold decrease in reporter gene expression, indicating that the transcriptional activity of these steroid receptors can be substantially reversed by a single point mutation.

The conformational plasticity of the steroid receptor protein offers the opportunity for its various interaction partners to couple to these conformational transitions in the process of regulation. Although the conformational states of unliganded steroid receptor proteins remain poorly characterized, crystal structures of various ligand-bound states demonstrate that the conformational transitions of the ligand binding domain which are induced by steroid antagonists generally differ from those induced by steroid agonists, and these distinct conformations can differentially interact with co-regulators (17). In addition, a concurrent conformational transition in the Hsp90 subunits has been proposed to serve as a component of a larger scale allosteric response (18).

Arguably most relevant to the present study, the binding of Hsp70 to the isolated glucocorticoid receptor-ligand binding domain induces a conformational transition that affects the residues neighboring the ligand binding pocket, and not only markedly reduces the intrinsic hormone binding affinity of apo-glucocorticoid receptor-ligand binding domain but also stimulates the release of hormone from the liganded receptor (19). Based on this Hsp70-mediated modulation of glucocorticoid receptor-ligand binding domain hormone affinity, Agard and colleagues (19) have proposed an ATP-dependent regulatory mechanism for altering receptor activity in response to the intracellular hormone concentration.

The markedly differing conformational dynamics of the FK1 domains of FKBP51 and FKBP52 provide a potential mechanism for differentially coupling to the regulatory transitions of the steroid receptor proteins as well as providing a potential basis for selective drug design (20). Unlike FKBP52, FKBP51 exhibits elevated 15N R2 relaxation rates throughout much of the β3-β2 loop with a magnetic field strength dependence that indicates conformational dynamics in the submillisecond time frame (21). This conformational exchange line broadening is quenched in the L119P variant of FKBP51. The complementary P119L variant of FKBP52 induces a FKBP51-like pattern of exchange line broadening for the residues along the β1-β2 loop, indicative of a similar underlying conformational transition (20).

The FKBP52-like L119P variant of FKBP51 also suppresses exchange line broadening of residues in the β2 and β3a strands near Phe-67, which has its structurally buried phenyl ring packed directly under the tip of the β1-β2 loop in the unliganded crystal structures (22). The bifurcated main chain hydrogen bonding of the amides of Phe-67 and Asp-68 with the carbonyl oxygen of Gly-59 disrupts the standard antiparallel hydrogen bonding pattern between the β2 and β3a strands. We earlier proposed that a transition in this bifurcated hydrogen bonding interaction could provide the structural basis of the conformational exchange line broadening observed for those strands (20). Interestingly, the analogous P119L mutation in FKBP52 has no effect on the line broadening of the β2 and β3a strands, indicating an additional nonequivalence in the conformational plasticity of FKBP51 and FKBP52.

It has remained unresolved whether the exchange line broadening in the β1-β2 loop and in the β2 and β3a strands of FKBP51 arise from a single concerted conformational transition or from two distinct coupled transitions. In the year following our initial characterization of the conformational exchange line broadening dynamics surrounding the bifurcated main chain hydrogen bonding of the amides of Phe-67 and Asp-68 with the carbonyl oxygen of Gly-59, Haush and colleagues (23) reported the crystal structures of two so-called iFit inhibitor-bound forms of the FK1 domain from FKBP51 in which the conformation of the β3a strand surrounding Phe-67 is altered so as to disrupt this bifurcated main chain hydrogen bonding interaction. They interpreted these structures to demonstrate an induced-fit mechanism of binding, despite the fact that the structure of the final ligand-bound state cannot discriminate between an induced-fit or a conformational selection-based mechanism (24). Particularly germane to these issues is whether FKBP51 significantly populates a similar conformation in the absence of the inhibitor. NMR relaxation analysis of systematically designed FK1 domain variants can provide a fruitful approach to characterizing such transient conformational substrates (25, 26).

**Experimental Procedures**

**Protein Preparation**—Gene sequences for the wild type and sequence variants of the FK1 domain of human FK506-binding protein FKBP51 (Glu20-Glu140) were chemically synthesized (Genscript), with codon optimization for expression in *Escherichia coli*. The gene sequences were cloned into the expression vector pET11a and the plasmids were transformed into the BL21Star(DE3) strain (Invitrogen) for expression. The protein expression and purification procedure for the FKBP51 domain was carried out as previously described (20). All isotopically labeled samples were prepared via protein expression in minimal medium containing 0.1% 15NH4Cl as nitrogen source. All protein samples were concentrated via centrifugal ultrafiltration and then equilibrated into a pH 6.50 buffer containing 25 mM sodium phosphate, 2 mM dithiothreitol, and 2 mM tris(2-carboxyethyl)phosphine by a series of centrifugal concentration steps to a final concentration of 0.5 mg/ml protein.

**NMR Spectroscopy**—NMR relaxation data were collected on a Bruker Avance III 600 MHz spectrometer and a Bruker Avance II 900 MHz spectrometer at 25 °C. HSQC-based T1, T1p, and heteronuclear NOE experiments were carried out as described by Lakomek *et al.* (27) using delays and spin lock field strengths for the T1p experiments as previously described for the wild type and L119P variants of the FK1 domain for FKBP51 (20). T1p data sets were collected at multiple frequency offsets (four at 60.8 MHz 15N and five at 91.2 MHz 13N).
To maximize the sensitivity for detecting variations in the conformational exchange line broadening effects between the wild type and the mutational variants, the transverse relaxation rates were analyzed in differential mode. For residues that do not exhibit conformational exchange line broadening, the backbone $^{15}$N $R_2$ values are generally dominated by the dynamics of global molecular tumbling, which depends upon sample viscosity. The median $R_2$ values were used to normalize the relaxation effects arising from slightly differing sample temperatures and protein concentrations.

The robustness of this normalization approach is directly indicated by the $\Delta R_2$ values for the residues not exhibiting substantial differential line broadening because these $\Delta R_2$ values incorporate the experimental errors of the individual relaxation measurements as well as the discrepancies that arise from a lack of equivalence for the conformational/orientational dynamics of the protein samples (20). As a means of further enhancing the robustness of the differential $R_2$ analysis, except where explicitly noted, $R_2$ relaxation values less than 25 s$^{-1}$ were excluded from the $\Delta R_2$ calculations if the root mean square deviation fit to the optimal decay constant was larger than 1.5% of the initial intensity.

**X-ray Crystallography**—Initial crystallization conditions for the FK1 domain of the K121G variant was established using the Hampton Research Crystal Screen I and II. Large crystals were grown at room temperature in hanging drops. 2 $\mu$l of protein solution at 65 mg/ml concentration was mixed with an equal volume of reservoir solution containing 26% PEG 3350, 0.1 M HEPES, pH 7.5, 0.2 M ammonium acetate, and 5% isopropyl alcohol, and equilibrated overnight. The crystals belong to space group $C_2$ with cell parameters: $a = 69.435$ Å, $b = 31.990$ Å, $c = 57.435$ Å, $\beta = 119.03^\circ$. There is one molecule per asymmetric unit, with a crystal solvent content of 42%. Prior to data collection, crystals were transferred to a reservoir solution containing crystallization buffer supplemented with 25% glycerol, and then flash-cooled under a nitrogen stream at 100 K, and stored in liquid nitrogen. Diffraction data were collected at 100 K using beamline X25 of the National Synchrotron Light Source (Brookhaven National Laboratory). These data were processed and scaled using HKL2000 (28). Using the full resolution range (to 1.2 Å) with the high resolution structure of FKBP51 (PDB code 3O5Q (22)) as a search model, only one clear solution was found with the PHASER molecular replacement program within the PHENIX suite (29). The $R_{factor}$ of the initial model was 48.6%. Structural refinement was carried out using PHENIX (30). Model rebuilding was carried out using Coot (31). Figures of crystallographic structures were generated using Chimera software (32) and PyMol ((33)).

**Results**

**Structural Analysis of the Interface between the $\beta_4$-$\beta_5$ Loop and the $\beta_4$+$\beta_{3a}$ Strands in the FK1 Domain of FKBP51**—For the 22 residues spanning Cys-107 at the end of the $\beta_4$ strand to Leu-128 at the beginning of the $\beta_5$ strand (Fig. 1), human FKBP51 and FKBP52 differ only at residues 119 and 124 (L119P and S124P). The five residues of the $\beta_{3a}$ strand (Lys$^{35}$-Ser$^{38}$) are completely conserved not only for FKBP51 and FKBP52 but for the close structural homologs FKBP12 and FKBP12.6 (Lys$^{35}$-Ser$^{38}$). Beyond these two segments, the FK1 domains of FKBP51 and FKBP52 exhibit only a 61% sequence identity, highly suggestive of a substantial functional role for this region of the protein.

The tip of the $\beta_4$-$\beta_5$ loop is positioned by a set of hydrophobic interactions involving the C$^{\beta}$ from residue 121 and the side chains of Ile-122 and Pro-123, which are packed against the phenyl ring of Phe-67 in the $\beta_{3a}$ strand and the side chain of Leu-61 at the end of the $\beta_5$ strand (Fig. 2A). This set of interacting residues is conserved in FKBP52, FKBP12, FKBP12.6, and in many of the other 18 FKBP domains within the human genome (34).

In hindsight, an early indication of the functional significance for FKBP51 and FKBP52 of the interface between the $\beta_4$-$\beta_5$ loop and the underlying $\beta_2$ and $\beta_{3a}$ strands was the widespread use of a F67D,D68V double mutant as a presumed indicator for the participation of the peptidyl prolyl isomerase activity for these proteins in their biological functions, in particular for steroid receptor activity (15, 35–38). More recently, in the context of their analysis of the effects of the L119P and P119L variants of FKBP51 and FKBP52 in the transcriptional activity of the steroid hormone receptor discussed above, Smith and colleagues (16) compared the F67D,D68V double mutant to other mutations that lay within the catalytic cleft to conclude that the steroid receptor activity does not depend upon prolyl isomerization catalysis.

In the effort to better understand how structural and dynamic effects propagate across the hydrophobic interface between the $\beta_4$-$\beta_5$ loop and the $\beta_4$ and $\beta_{3a}$ strands, we have...
structure throughout the rest of this interface remained closely similar to that observed in the highest resolution structure reported for the wild type protein (22). The absence of discernible structural perturbation arising from the K121G substitution enhances the reliability with which our NMR relaxation results for the residue 121 side chain substitutions could be interpreted.

The Exchange Line Broadening of the $\beta_4$-$\beta_5$ Loop and the $\beta_2$+$\beta_3$о Strands Arise from Distinct Conformational Transitions Which Are Energetically Coupled—Each of the three standard NMR relaxation experiments, $R_1$, $R_2$ (or $R_{1\infty}$), and heteronuclear NOE, are sensitive to conformational motion that is faster than the molecular rotational correlation time (~10 ns for smaller globular proteins). Previous studies on the wild type FK1 domain of FKBP51 revealed three segments in which both the $^{15}$N $R_1$ relaxation rates and the heteronuclear NOE values were reduced, relative to the rest of protein backbone, indicative of motion in the picosecond-nanosecond time frame. However, for two of these three segments, the $\beta_3\approx\beta_3\beta_4$ loop and the tip of the long $\beta_2\approx\beta_5$ loop, the $^{13}$N $R_2$ relaxation rates are increased (Fig. 3A). These increased rates reflect the fact that the $R_2$ transverse relaxation experiment is also sensitive to motion in the conformational exchange line broadening regime (microsecond-millisecond). The residues at the tip of the $\beta_2\approx\beta_5$ loop have also been reported to have elevated crystallographic $B$-factors (22), which is consistent with the common observation that the orientational disorder implied by residues exhibiting reduced relaxation rates and NOE values is often reflected in local crystallographic disorder.

Upon introducing the K121G mutation, the conformational exchange line broadening for the $\beta_2\approx\beta_5$ loop of FKBP51 is almost completely eliminated (Fig. 3B). Interestingly, the small

### Table 1

| Crystallographic data collection, refinement, and model details (PDB code 4R0X) |
|-----------------------------------------------|
| **Statistics** |
| Data collection |
| Resolution range (Å) | 27–1.24 (1.27–1.24) |
| No. of unique reflections | 30,633 |
| Redundancy | 6.4 (3.1) |
| Completeness (%) | 97.6 (74.2) |
| Average $I/\sigma(I)$ | 47.1 (4.4) |
| $R_{merge}$ (%) | 4.4 (30.1) |
| Refinement |
| Resolution limits (Å) | 27–1.24 |
| No. reflections | 30,633 |
| Minimum $F/\sigma F$ | 1.39 |
| $R_{work}$ (%) | 15.8 |
| $R_{free}$ (%) | 16.7 |
| Non-H atoms |
| Protein | 953 |
| Water | 199 |
| Average B (Å$^2$) | 17.7 |
| All atoms | 15.3 |
| Solvent | 29.1 |
| Wilson B (Å$^2$) | 12.4 |
| Geometry |
| Root mean square deviation bond length (Å) | 0.005 |
| Root mean square deviation bond angle (°) | 1.034 |
| Ramachandran Plot |
| Favored | 97.5 |
| Allowed | 2.5 |
| Outliers | 0 |

**FIGURE 2.** The hydrophobic packing interactions between the $\beta_4$-$\beta_5$ loop and the $\beta_2$+$\beta_3$о strands in the wild type and K121G variant of the FK1 domain of FKBP51. A, the packing interactions between Ile-122 and Pro-123 of the $\beta_4$-$\beta_5$ loop, Leu-61 of the $\beta_3$о strand, and Phe-67 of the $\beta_2$о strand are conserved among FKBP51 and FKBP52 as well as among FKBP12 and FKBP12.6 family members as illustrated in the high resolution structures of FKBP51 (yellow, PDB code 3OSQ (22)). These packing interactions are closely similar to that observed in the highest resolution structure reported for the wild type protein (22). The crystal structure throughout the rest of this interface remained closely similar to that observed in the highest resolution structure reported for the wild type protein (22). The absence of discernible structural perturbation arising from the K121G substitution enhances the reliability with which our NMR relaxation results for the residue 121 side chain substitutions could be interpreted.

**Coupling of Conformational Transitions in FKBP51**

Given that the C$^B$ atom of Lys-121 is packed tightly against the C$^B$ atom of Phe-67 at the site of the inter-strand bifurcated hydrogen bonding discussed above, we wished to analyze how alterations in that steric interaction might modulate the conformational dynamics on either side of the interface. Because removal of the C$^B$ atom at residue 121 would eliminate this steric interaction, we solved the crystal structure of the K121G variant of FKBP51 to a resolution of 1.24 Å (Table 1) to determine whether the glycine substitution might introduce variations into the backbone conformation that could complicate the structural interpretation of the NMR relaxation measurements. The targeted steric interaction was eliminated as expected (Fig. 2), whereas the crystallographic...
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FIGURE 3. $^{15}$N transverse relaxation measurements for the backbone amide resonances in the FK1 domain of wild type and $\beta_4\beta_5$ loop variants of FKBP51. Transverse ($R_2$) relaxation rates at 600 MHz $^1$H (black) and 900 MHz $^1$H (red) are shown for the wild type (A) and K121G variant (B) of the FK1 domain. For the differential $R_2$ comparison at 600 MHz $^1$H (C) and 900 MHz (D), the median $R_2$ values for the non-exchange broadened residues of the wild type (blue), K121A (green), and K121G (red) variant data sets were scaled to that of the L119P variant to correct for small variations in the global molecular correlation time due to minor differences in sample temperature and concentration. Given the increased experimental uncertainty of the fast relaxing resonances in the $\beta_4\beta_5\beta_3a$ loop (Ser-70, Arg-73, and Glu-75), these residues were not included in the differential analysis. For the residues outside of segments 57–77 and 108–128, the median average differences for $\Delta R_2$ ($s^{-1}$) were 0.083 and 0.091 for K121G, 0.088 and 0.113 for K121A, and 0.093 and 0.090 for wild type at 600 MHz and 900 MHz, respectively.

A degree of exchange line broadening in the $\beta_2$ and $\beta_3a$ strands previously reported for the wild type protein (20) is significantly enhanced in the K121G variant.

These effects are more clearly illustrated when the transverse relaxation rates for the samples are directly analyzed in differential mode. In the present case (Fig. 3, C and D), the L119P data set was used as reference due to the strong attenuation of exchange line broadening in the $\beta_3\beta_5$ loop of this variant (20). In comparing protein samples for which it is warranted to assume that the differences in internal motion affecting the $R_2$ values occur exclusively within the conformational exchange line broadening time regime, the resultant cancellation of errors provided approximately a 3-fold increase in precision for the data of this study, relative to that obtained by applying the standard model-free analysis approach of Lipari and Szabo (42) to the individual protein data sets. In illustration, for the 72 backbone amide resonances outside of segments 57–77 and 108–128, which contain the residues exhibiting systematic differences in line broadening, the median average difference for the $\Delta R_2$ values of the K121G and L119P variants at 600 (Fig. 3C) and 900 MHz (Fig. 3D) were ~0.8% of the median $R_2$ values. The line broadening effects in the $\beta_2$ and $\beta_3a$ strands arising from the K121G substitution are up to 80-fold larger than this median average difference for the $\Delta R_2$ values outside of the two affected segments.

For the conformational exchange broadening both in the $\beta_4\beta_5$ loop and within the $\beta_2$ and $\beta_3a$ strands, the effects of the K121A substitution were intermediate between those of the K121G variant and the wild type protein. In contrast to the suppression of exchange line broadening in both the $\beta_4\beta_5$ loop and $\beta_2$ and $\beta_3a$ strands caused by the L119P variant, the K121G and K121A substitutions enhance the exchange line broadening in the $\beta_2$ and $\beta_3a$ strands, whereas at the same time suppressing line broadening in the $\beta_4\beta_5$ loop. This opposite pattern of response to the substitutions at residue 121 indicates that the differential exchange line broadening of the $\beta_2$ and $\beta_3a$ strands does not reflect the propagation of dynamics from the $\beta_4\beta_5$ loop, but rather a distinct conformational transition underlies the exchange line broadening of $\beta_2$ and $\beta_3a$ strands.

Comparison of the $\Delta R_2$ values at 600 and 900 MHz indicates an approximate 2.25-fold increase for residues in both the $\beta_2$ and $\beta_3a$ strands and the $\beta_4\beta_5$ loop (Fig. 3, C and D). Variation of conformational exchange line broadening with the square of the magnetic field strength is indicative of these conformational transitions occurring in the submillisecond time frame approaching the fast exchange limit (21), as had previously been reported for the $\beta_4\beta_5$ loop in FKBP12 (43–45).

For a two-state transition near the fast exchange time regime, the magnitude of the exchange line broadening is proportional to $p_A p_B \Omega^2/k_{ex}$, where $p_A$ and $p_B$ are the state populations, $\Omega$ is the difference in $^{15}$N resonance frequency for the two states, and $k_{ex} = k_{AB} + k_{BA}$ is the conformational exchange rate. The fact that the exchange line broadening effects for the various residues within the $\beta_4\beta_5$ loop scale proportionally in going from K121G to K121A to the wild type protein strongly indicates that the differential $^{15}$N chemical shifts and therefore presumably the conformation of the interchanging states are largely unaffected by these substitutions. As a result, the marked decrease in exchange line broadening within the $\beta_4\beta_5$ loop in going from wild type to K121A to K121G variants arises from a large decrease in the population of the minor conformational state and/or from a large increase in the rate of exchange. Given the increase in flexibility that could be expected from decreasing the steric bulk of the side chain at residue 121, it is anticipated that an increased rate of conformational exchange likely dominates these line broadening effects.

The packing of the $\beta_4\beta_5$ loop against the $\beta_2$ and $\beta_3a$ strands plays a significant role in determining the dynamics of the exchange line broadening transition for residues in those strands. The line broadening effects for residues within the $\beta_2$ and $\beta_3a$ strands also scale nearly proportionally going from the
wild type to the K121A and K121G variants, consistent with the conformations of interchanging states being largely unaffected by these substitutions. In contrast to the $\beta_2$-$\beta_{3a}$ loop, the magnitude of the exchange line broadening in the $\beta_2$ and $\beta_{3a}$ strands increases in going from wild type to the K121A and K121G variants. This increase predominately reflects either a decreased rate for the conformational transition of the $\beta_2$ and $\beta_{3a}$ strands or an increase in the population of the minor conformational state for that transition. As the K121G and K121A variants reduce the packing interaction between the $\beta_2$-$\beta_{3a}$ loop and the $\beta_{3a}$ strand, an increase in the minor state population offers a more plausible explanation for the increased exchange line broadening.

Selective Suppression of the $\beta_2$+$\beta_{3a}$ Strand Exchange Line Broadening Transition in FKBP51—As noted above, the introduction of the FKBP51-like P119L mutation into the FK1 domain of FKBP52 caused the $\beta_2$-$\beta_{3a}$ loop to exhibit a FKBP51-like pattern of exchange line broadening effects, albeit with a 5-fold lower intensity at the various residues along that loop. Introducing a second FKBP51-like mutation P124S into the 5 loop differs significantly from that observed for wild type FKBP51. It may be noted that the bifurcated main chain hydrogen bond disruption of the FKBP51-like P119L mutation into the FK1 domain of FKBP52 caused the $\beta_2$-$\beta_{3a}$ loop to exhibit a FKBP51-like pattern of exchange line broadening to a level of 60% that of the wild type FKBP51. For the wild type protein (Fig. 3A), is only modestly reduced from that observed for the L119P variant (Fig. 3A).

4LAV, and 4LAW (48), a hydrogen bond between the $\beta_2$ and $\beta_{3a}$ strands in the FK1 domain of FKBP52 surrounding Thr-58. Thr-58 is the only residue displayed that differs from the sequence of FKBP51. As illustrated in the 1.90-Å resolution crystal structure (PDB code 1Q1C (47)), the hydrogen bonding between the side chain hydroxyl of Thr-58 and the O* of Ser-69 indirectly stabilizes the hydrogen bond between the Ser-69 O* and the carbonyl oxygen of Tyr-57, which is not observed in the highest resolution structures of FKBP51.

The L119P variant only partially suppresses the exchange line broadening in the $\beta_2$ and $\beta_{3a}$ strands (20). To test the premise that introducing the K58T substitution into FKBP51 might stabilize the $\beta_2$-$\beta_{3a}$ strand interactions seen in the crystal structures of the unliganded protein, $^{15}$N relaxation measurements were carried out on this variant. Qualitatively, the $^{15}$N $R_2$ data for the K58T-substituted FKBP51 (Fig. 6A) offers no apparent evidence for conformational exchange line broadening in the $\beta_2$ and $\beta_{3a}$ strands. More rigorously, the model-free analysis (42, 49) of the $^{15}$N $T_1$, $T_{1p}$, and heteronuclear NOE data for K58T variant predicted exchange broadening at both magnetic fields only for Lys-60, and these predicted values are only slightly above the estimated experimental uncertainty. Interestingly, the K58T substitution also results in a 3-fold reduction in the much larger exchange line broadening effects within the $\beta_{3a}$-$\beta_{3a}$ loop (Fig. 6A), suggesting that the conformational transition that gives rise to those line broadening effects also involves the disruption of the hydrogen bonding interaction between the side chains of Thr-58 and Ser-69. At the same time, the exchange line broadening in the $\beta_2$-$\beta_{3a}$ loop of the K58T variant (Fig. 6A) is only modestly reduced from that observed for the wild type protein (Fig. 3A).

$^{15}$N Chemical Shift Analysis of the Exchange Line Broadening Transition in the $\beta_2$ and $\beta_{3a}$ Strands—Given the suppression of exchange line broadening observed for the K58T variant of FKBP51 in the $\beta_2$ and $\beta_{3a}$ strands, this protein provides an appropriate reference state for comparing the differences in exchange broadening for this segment in the wild type and K121A and K121G variants. The largest exchange line broadening effects at 600 (Fig. 6B) and 900 MHz (Fig. 6C) were...
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FIGURE 6. $^{15}$N transverse relaxation in the K58T variant of FKBP51 and structural interpretation of the $\Delta R_2$ values for wild type and the K121A and K121G variants. $R_2$, relaxation rates at 600 MHz $^1$H (black) and 900 MHz $^1$H (red) are shown for the K58T variant of the FK1 domain (A). For the differential $R_2$ comparison within the $\beta_2$ and $\beta_{3a}$ strands at 600 (B) and 900 (C) MHz, the median $R_2$ values for the wild type (blue), K121A (green), and K121G (red) data sets were scaled to that of the K58T variant to correct for small variations in the global molecular correlation time. A subset of the Asp$^{68}$ relaxation data are denoted with open bars indicating root mean square deviation fits to the optimal decay constants larger than the 1.5% cutoff used for the other $\Delta R_2$ values. Overlap with the resonance of Val-94 precluded the use of the Lys-60 data for wild type and the K121A variant. SPARTA+$^*$-based differential $^{15}$N chemical shift values were predicted for the iFit1-bound (black) and iFit4-bound (gray) domains, as compared with the reference FK506-bound FK1 domain of FKBP51 (D).

observed for the amides of Lys-60, Lys-66, Phe-67, and Asp-68, strongly suggestive of a transition involving the bifurcated main chain hydrogen bond (Fig. 4). In such a transition, the substantial exchange line broadening of the Lys-60 amide might reflect not only changes in the local backbone geometry of the $\beta_2$ strand, it could also reflect alterations in the hydrogen bonding of the Gly-59 carbonyl oxygen, which serves as the acceptor for the bifurcated hydrogen bonds to the $\beta_{3a}$ strand. Disruption of hydrogen bonding at a peptide $C=O$ group is known to give rise to changes in the chemical shift for the peptide nitrogen resonance of several ppm (50).

Early studies of FKBP12 replaced Phe-36 with valine so as to introduce a hydrophobic pocket along the side of the packing interface between the $\beta_2$-$\beta_3$ loop and the underlying $\beta_{3a}$ strand (51, 52). This F36V variant was then used to develop a series of selective inhibitors, including Shield1 (53), which have been applied to numerous mammalian cellular expression studies (e.g. Refs. 54 and 55). Introducing the analogous F67V mutation to the FK1 domains of FKBP51 and FKBP52, Hausch and colleagues (23) recently reported the development of FKBP51-selective inhibitors using Shield1 as their initial scaffold. In contrast to earlier studies on the F36V variant of FKBP12, which exhibited minimal structural perturbation upon inhibitor binding (51), Hausch and colleagues (23) found that their iFit inhibitors bound to a conformation of wild type FKBP51 and FKBP52 in which the Phe-67 side chain is reoriented away from its typical position packed underneath the tip of the $\beta_2$-$\beta_3$ loop (Fig. 2).

As illustrated for the complex with the iFit1 inhibitor (PDB code 4TW6 (23)) (Fig. 7), the resultant backbone conformation of the $\beta_2$ and $\beta_{3a}$ strands entails a flipping of the Phe-67 peptide linkage. In contrast to Phe-67 and Asp-68 forming a bifurcated hydrogen bond interaction with the carbonyl of Gly-59, as seen in the reference FK506-inhibited structure (PDB code 3O5R (22)), the amide of Asp-68 in the iFit1-inhibited structure now forms a linear hydrogen bond with the carbonyl oxygen of Gly-59. The resultant rearrangement in the $\beta_{3a}$ strand causes a major repositioning of the $\beta_{1a}$-$\beta_{3a}$ bond vector of Phe-67 so that the phenyl ring is now directed out toward the solvent phase (Fig. 7B). The binding of the iFit1 inhibitor is also accompanied by an appreciable shift in position for the tip of the $\beta_4$-$\beta_5$ loop, which is moved away from the $\beta_{3a}$ strand.

Because the amide of Lys-66 is also oriented away from the $\beta_2$ strand, the flipping of the Phe-67 peptide linkage in the iFit1-inhibited structure results in the kinked $\beta_{3a}$ strand conformation being shifted by one residue toward the $\beta_2$-$\beta_{3a}$ loop (Fig. 7). Otherwise, the conformation of the $\beta_2$-$\beta_{3a}$ loop and the $\beta_2$ strand in the iFit1-inhibited structure roughly follows that of the reference FK506-inhibited protein. The iFit4-inhibited structure (PDB code 4TW7 (23)) differs more strongly from the reference FK506-inhibited protein. In this case, a more extended canonical antiparallel hydrogen bonding pattern is formed, and the “kink” has been further shifted so as to contribute to an enlarged $\beta_2$-$\beta_{3a}$ loop.

Conformational exchange line broadening data provide a measure of the relative magnitude of the differential $^{15}$N chemical shift values for the residues affected by this transition. Although these differential $^{15}$N chemical shift values alone will generally be insufficient to provide de novo structural determination, given plausible structural models for the transition, experimental $\Delta R_2$ values can provide a valuable test for the reliability of those structural models.

If the exchange line broadening effects in the $\beta_2$ and $\beta_{3a}$ strands arise from a two-state transition between conformations similar to the crystal structures of the FK506- and iFit-
inhibited proteins, as displayed in Fig. 7, prediction of the $^{15}$N chemical shifts for those two conformational states would provide an estimate for the relative magnitude of the exchange line broadening for the residues of these $\beta$ strands. A number of empirical algorithms have been developed for predicting NMR chemical shifts from protein structures (56–59). Although the capability of these algorithms to predict individual chemical shift values for residues within a given protein is still fairly modest, the robustness of this approach can be significantly improved when applied in a differential mode that provides for the cancellation of various modeling uncertainties, as recently illustrated in the analysis of the $\beta_2$-$\beta_3$ loop transition in FKBP12 (40).

Using the SPARTA+ program of Shen and Bax (56), the $^{15}$N chemical shifts were predicted for the iFit1- and iFit4-inhibited crystal structures of the FKBP51 FK1 domain as well as for the reference FK506-inhibited structure. In addition to the dependence of chemical shift on the local peptide backbone conformation, the SPARTA+ algorithm (56) uses the Haigh-Mallion model (60, 61) for estimating aromatic ring current effects and also incorporates electric field effects for the $^1$H resonances.

Changes in the backbone conformation thoroughly dominate the predicted differences in $^{15}$N chemical shift for these liganded FK1 domains. Despite the marked reorientation of the phenyl ring of Phe-67 upon the binding of the iFit inhibitors, the predicted differential contributions from the aromatic ring current effects are quite small. In particular, for the residues exhibiting the largest differential exchange line broadening (Lys-60, Lys-66, Phe-67, and Asp-68 in Fig. 6, B and C) the maximal ring current contribution to the differential $^{15}$N chemical shift is only 0.1 ppm (Fig. 6D).

With respect to the FK506-inhibited reference structure, the iFit1- and iFit4-inhibited structures yield similar patterns of predicted differential $^{15}$N chemical shifts for the $\beta_2$ and $\beta_{3a}$ strands (Fig. 6D), consistent with their qualitatively similar conformations. The correlation coefficient for differential $^{15}$N chemical shifts over these 11 residues is 0.93 between the iFit1- and iFit4-inhibited structures.

Particularly for the iFit1-inhibited structure, the predicted differential $^{15}$N shifts correlate rather well with the observed $\Delta R_2$ values in the $\beta_2$ and $\beta_{3a}$ strands as well as in the intervening loop. Averaging over the $\Delta R_2$ values for the 900 MHz datasets (Fig. 6C) of the wild type, K121A, and K121G proteins (data for Lys-60 and Leu-61 normalized by the average for each protein variant), the correlation coefficient with respect to the differential $^{15}$N shifts predicted from the iFit1-inhibited structure was also 0.93. Thus within the structural sensitivity of these predicted differential $^{15}$N chemical shifts, the transient conformation that gives rise to the exchange line broadening in the $\beta_2$ and $\beta_{3a}$ strands would appear to be as similar to the iFit1-inhibited crystal structure as that structure is to the iFit4-inhibited structure.

**Discussion**

The correspondence between the crystal structure-based prediction of differential $^{15}$N chemical shifts and the observed exchange line broadening within the $\beta_2$ and $\beta_{3a}$ strands of the unliganded FK1 domain of FKBP51 strongly supports the expectation that this line broadening arises from a transition to a significantly populated conformation that is similar to what is observed in the crystal structures of the iFit-inhibited domain. As such, conformational selection is surely the more appropriate mechanism to assume for the binding interaction of FKBP51 with the iFit family of inhibitors until more detailed thermodynamic and kinetics studies are obtained. More generally, this exchange line broadening analysis for the $\beta_2$ and $\beta_{3a}$ strands in the FK1 domain of FKBP51 provides further demonstration of the utility of NMR relaxation analysis in identifying sites of transient conformational transitions that are potentially amenable to systematic drug targeting.

Hausch and colleagues (23) have proposed that the altered conformation that they observed in their crystal structures of the iFit-inhibited FK1 domains is biochemically relevant. One line of potential support for that hypothesis comes from their demonstration that the introduction of either one of two
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FKBP51-like substitutions, T58K or W60K, into FKBP52 yields a variant that binds the fluorescent iFit-FL ligand more tightly than does the wild type FKBP52 domain by over 100-fold, thus binding nearly as tightly as does FKBP51 (23). Conversely, introducing the K58T or the K60W mutation into FKBP51 markedly reduces the binding affinity for this ligand, consistent with a corresponding alteration in the stability of the conformation in the crystal structures of the iFit-inhibited protein.

In a set of experiments designed to test the regulation of the glucocorticoid receptor, Haush and colleagues (23) used the brain barrier-permeable analog SAFit2 in mouse studies to inhibit the endogenous FKBP51. They found a reduced corticosterone level at the peak of the circadian cycle and an enhanced suppression of corticosterone levels following the combined treatment with dexamethasone and corticotrophin-releasing factor, results that closely mimic the behavior of FKBP51 null mice. Although indicating that the signaling functions of FKBP51 can be selectively inhibited in vivo, these studies do not address the issue of whether the conformational transition utilized in binding the inhibitor also participates in the physiological functioning of this signaling pathway.

If a conformational transition involving the rearrangement of the Phe-67 side chain and the bifurcated main chain hydrogen bonding interactions of the $\beta_2$ and $\beta_{3a}$ strands is critical for mediating steroid receptor interactions, and furthermore, single point mutations can efficiently interchange the conformational sampling of the $\beta_{3a}$ strand in both FKBP51 and FKBP52, it is interesting that only mutations in the $\beta_4$-$\beta_5$ loop were repeatedly isolated in the hormone-induced transcription selection studies of Smith and colleagues (16). In that earlier study, a total of 25 gain-of-function mutants were isolated from eight independent error-prone PCR libraries of FKBP51 variants. In addition to five independent occurrences of the L119P mutation discussed above, at least three independent isolates were found for both the FKBP52-like S124P variant and the nonhomologous A116V variant, which in one isolate was found combined with the L119P mutation. This A116V,L119P double mutant yielded transcription activity levels fully equivalent to those of wild type FKBP52.

Although the results of Smith and colleagues (16) do not exclude the possibility that such a conformational transition in the $\beta_{3a}$ strand participates in mediating steroid receptor signaling, these results do strongly indicate that such a conformational plasticity-based mechanism should incorporate the additional transition of the $\beta_4$-$\beta_5$ loop. The present study demonstrates the energetic coupling between the transition centered in the $\beta_4$-$\beta_5$ loop and a second transition centered in the $\beta_{3a}$ strand. The most physically plausible interpretation of the pattern of mutationally induced variations in the exchange line broadening behavior indicates that an increase in conformational flexibility near the tip of the $\beta_4$-$\beta_5$ loop gives rise to an increased population of the reoriented conformation of the $\beta_{3a}$ strand, fully consistent with the requirement of enabling the side chain of Phe-67 to escape from its burial within the tightly packed interface (Fig. 7).

A model for the differential interaction with the steroid receptor can be posited in which the transition to an alternate conformation in the $\beta_4$-$\beta_5$ loop shifts the equilibrium between the Phe-67-in and the Phe-67-out conformations of the $\beta_{3a}$ strand so as to favor the latter conformation (Fig. 8). The suggestion that the tip of the $\beta_4$-$\beta_5$ loop might serve as a latch for preferentially stabilizing the Phe-67-in conformation is in accord with the observation that the binding of the iFit1 inhibitor is accompanied by the $\beta_4$-$\beta_5$ loop moving away from the $\beta_{3a}$ strand (Fig. 7B).

Because the alternate $\beta_4$-$\beta_5$ loop conformation and the Phe-67-out conformation are both more readily populated in the FKBP51 domain, that protein more favorably adopts this combination of local conformations, which is assumed to more strongly interact with the steroid receptor complex in a low affinity state for hormone binding. For FKBP52, Pro-119 inhibits the transition of the $\beta_4$-$\beta_5$ loop to an alternate conformation, whereas Thr-58 (and perhaps Thr-60) inhibits the transition to the Phe-67-out conformation. Note that even when the FKBP51 sequence for the $\beta_4$-$\beta_5$ loop is incorporated into FKBP52 (i.e. P119L,P124S), no exchange line broadening of the $\beta_{3a}$ strand is observed (20). Hence interactions in both the $\beta_4$-$\beta_5$ loop and the $\beta_4$-$\beta_{3a}$ strands tend to lock FKBP52 into a...
conformation which preferentially binds to a high affinity state of the steroid receptor.

Currently, there is no detailed structural model for the conformational transition that underlies the exchange line broadening observed within the $\beta_1$-$\beta_2$ loop in the FK1 domain of FKBP51. However, previous studies on both FKBP12 and FKBP12.6 can provide useful insights into this process. The pattern of conformational exchange line broadening for the residues along the $\beta_4$-$\beta_5$ loop in the FK1 domain of FKBP51 is similar to that observed for FKBP12 (39). Using a combination of structure-based $\text{^{15}N}$ chemical shift predictions and mutations that selectively modulate the $\beta_4$-$\beta_5$ loop transition in FKBP12, we were able to deduce that a central structural element of that transition involves a shift in the backbone $\phi$ torsion angle for Gly-89 from the positive value, as observed in all reported crystal structures, to a negative $\phi$ angle (40). Given that the homologous residue of FKBP51 is Pro-120, which is covalently constrained to a negative $\phi$ torsion angle, the detailed structural mechanism of the exchange line broadening transition must necessarily differ in this case.

Our recent report of the first crystal structures of FKBP12.6 in the unliganded state (62) may provide a more useful initial model for interpreting structural aspects of this exchange line broadening transition in FKBP51. Structures of unliganded FKBP12.6 were obtained from two distinct crystal forms in which the $\beta_4$-$\beta_5$ loop conformations primarily differ from each other due to a flipping of the $(\psi_{\gamma_{99}}\phi_{\gamma_{90}})$ main chain torsion angles. In the P3,21 (PDB code 4IQC) crystal form a side chain methyl group of Val-90 at the tip of the $\beta_4$-$\beta_5$ loop is oriented toward the catalytic cleft as is the case for the homologous Ile-90 side chain in all reported crystal structures of FKBP12. In contrast, The P2$_1$ (PDB code 4IQ2) crystal form has a Val-90 methyl group directed away from the catalytic cleft. If a similar transition were to occur in FKBP51, the side chain of the homologous Lys-121 (with its C$^\beta$ packed snugly against the C$^\beta$ of Phe-67 as illustrated in Fig. 2) would undergo a substantial reorientation that could contribute to the differing interactions of FKBP51 with the low and high affinity states of the steroid receptor. It should be noted that the cis peptide linkage of Pro-120 in the FK1 domain of FKBP51 results in crystal structures in which the $(\psi_{\delta_{120}}\phi_{\kappa_{121}})$ torsion angles differ appreciably from homologous $(\psi_{\gamma_{99}}\phi_{\gamma_{90}})$ angles of either unliganded FKBP12.6 structure so that an analogous transition must differ in detail. Nevertheless, the transition observed from the crystal structures of unliganded FKBP12.6 may provide a useful initial model for future experimental studies.

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References

1. Tai, P. K., Maeda, Y., Nakao, K., Wakim, N. G., Duhring, J. L., and Faber, L. E. (1986) A 59-kilodalton protein associated with progestin, estrogen, androgen and glucocorticoid receptors. Biochemistry 25, 5269–5275
2. Silverstein, A. M., Galigniana, M. D., Kanelakis, K. C., Radany, C., Renoir, J. M., and Pratt, W. B. (1999) Different regions of the immunophilin FKBP52 determine its association with the glucocorticoid receptor, hsp90, and cytoplasmic dynein. J. Biol. Chem. 274, 36890–36896
3. Sanchez, E. R. (1990) HSP56: a novel heat shock protein associated with untransformed steroid receptor complexes. J. Biol. Chem. 265, 22067–22070
4. Smith, D. F., Faber, L. E., and Toft, D. O. (1990) Purification of unactivated progesterone receptor and identification of novel receptor-associated proteins. J. Biol. Chem. 265, 3996–4003
5. Denny, W. B., Valentine, D. L., Reynolds, P. D., Smith, D. F., and Scammell, J. G. (2000) Squirrel monkey immunophilin FKBP51 is a potent inhibitor of glucocorticoid receptor binding. Endocrinology 141, 4107–4113
6. Wochnik, G. M., Rüegg, J., Abel, G. A., Schmidt, U., Holsboer, F., and Rein, T. (2005) FK506-binding proteins 51 and 52 differentially regulate dynine interaction and nuclear translocation of the glucocorticoid receptor in mammalian cells. J. Biol. Chem. 280, 4609–4616
7. Davies, T. H., Ning, Y. M., and Sánchez, E. R. (2002) A new first step in activation of steroid receptors: hormone-induced switching of FKBP51 and FKBP52 immunophilins. J. Biol. Chem. 277, 4597–4600
8. Baughman, G., Wiederrecht, G. J., Chang, F., Martin, M. M., and Bourgeois, S. (1997) Tissue distribution and abundance of human FKBP51, an FK506-binding protein that can mediate calcineurin inhibition. Biochem. Biophys. Res. Commun. 232, 437–443
9. Binder, E. B., Salyakina, D., Lichtner, P., Wochnik, G. M., Ising, M., Pütz, B., Papilo, S., Seaman, S., Lucase, S., Kohli, M. A., Nickel, T., Künzel, H. E., Fuchs, B., Majer, M., Pfennig, A., Kern, N., Brunner, I., Modell, S., Baghai, T., Deiml, T., Zill, P., Bondy, B., Rupprecht, R., Messer, T., Köhnlein, O., Dabritz, H., Brückl, T., Müller, N., Pfister, H., Lieb, R., Mueller, J. C., Löhnmussar, E., Strom, T. M., Bettecken, T., Meitinger, T., Uhr, M., Rein, T., Holsboer, F., and Muller-Myskoss, B. (2004) Polymorphisms in FKBP5 are associated with recurrence of depressive episodes and rapid response to antidepressant treatment. Nat. Genet. 36, 1319–1325
10. Galigniana, N. M., Ballmer, L. T., Toneatto, J., Erlejman, A. G., Lagadari, M., and Galigniana, M. D. (2012) Regulation of the glucocorticoid response to stress-related disorders by the Hsp90-binding immunophilin FKBP51. J. Neurochem. 122, 4–18
11. Pei, H., Li, L., Fridley, B. L., Jenkins, G. D., Kalarri, K. R., Lingle, W., Petersen, G., Lou, Z., and Wang, L. (2009) FKBP51 affects cancer cell response to chemotherapy by negatively regulating Akt. Cancer Cell 16, 259–266
12. Fabian, A. K., März, A., Neimanis, S., Biondi, R. M., Kozany, C., and Hausch, F. (2013) InterAKTions with FKBP5s: mutational and pharmacological exploration. PLoS ONE 8, e57508
13. Stechschulte, L. A., Hinds, T. D., Jr., Khuder, S. S., Shou, W., Najjar, S. M., and Sanchez, E. R. (2014) FKBP51 controls cellular adipogenesis through P38 kinase-mediated phosphorylation of GRα and PPARγ. Mol. Endocrinol. 28, 1265–1275
14. Stechschulte, L. A., Hinds, T. D., Jr., Ghanem, S. S., Shou, W., Najjar, S. M., and Sanchez, E. R. (2014) FKBP51 reciprocally regulates GRα and PPARγ activation via the AKT-P38 pathway. Mol. Endocrinol. 28, 1254–1264
15. Riggs, D. L., Roberts, P. J., Chirillo, S. C., Cheung-Flynn, J., Prapapanich, V., Ratajczak, T., Gaber, R., Picard, D., and Smith, D. F. (2003) The Hsp90-binding peptidylprolyl isomerase FKBP52 potentiates glucocorticoid signaling in vivo. EMBO J. 22, 1158–1167
16. Riggs, D. L., Cox, M. B., Tardif, H. L., Hessling, M., Buchner, J., and Smith, D. F. (2007) Noncatalytic role of the FKBP52 peptidyl-prolyl isomerase domain in the regulation of steroid hormone signaling. Mol. Cell. Biol. 27, 8658–8669
17. Pfaff, S. J., and Fletterick, R. J. (2010) Hormone binding and co-regulator binding to the glucocorticoid receptor are allosterically coupled. J. Biol. Chem. 285, 15256–15267
18. Treble, P. J., Woolven, J. M., Saunders, K. A., Simpson, K. D., Farrow,
Coupling of Conformational Transitions in FKBP51

S. N., Matthews, L. C., and Ray, D. W. (2013) A ligand-specific kinetic switch regulates glucocorticoid receptor trafficking and function. J. Cell Sci. 126, 3159–3169

Kirschke, E., Goswami, D., Southworth, D., Griffin, P. R., and Agard, D. A. (2014) Glucocorticoid receptor function regulated by coordinated action of the Hsp90 and Hsp70 chaperone cycles. Cell 157, 1685–1697

Mustafi, S. M., LeMaster, D. M., and Hernández, G. (2014) Differential conformational dynamics in the closely homologous FK506-binding domains of FKBP51 and FKBP52. Biochem. J. 461, 115–123

Palmer, A. G., 3rd. (2014) Chemical exchange in biomacromolecules: past, present and future. J. Am. Chem. Soc. 136, 2867–2877

Boehr, D. D., Nussinov, R., and Wright, P. E. (2009) The role of dynamic conformational ensembles in biomolecular recognition. Nat. Chem. Biol. 5, 789–796

Sekhar, A., and Kay, L. E. (2013) NMR paves the way for atomic level broadening defines the NMR chemical shift time scale. J. Am. Chem. Soc. 135, 8348–8353

Kirschke, E., Goswami, D., Southworth, D., Griffin, P. R., and Agard, D. A. (2014) Structural basis of conformational transitions in the active site and 80’s loop in the FK506-binding protein FKBP12. Biochem. J. 458, 525–536

Anderson, J. S., Mustafi, S. M., Hernández, G., and LeMaster, D. M. (2014) Statistical allosteric coupling to the active site indole ring flip equilibria in the FK506-binding domain. Biophys. Chem. 192, 41–48

Lipari, G., and Szabo, A. (1982) Model-free approach to the interpretation of nuclear magnetic resonance relaxation in macromolecules: 1. theory and range of validity. J. Am. Chem. Soc. 104, 4546–4559

Mustafi, S. M., Chen, H., Li, H., Lemaster, D. M., and Hernández, G. (2013) Analyzing the visible conformational substates of the FK506-binding protein FKBP12. Biochem. J. 453, 371–380

Mustafi, S. M., Brecher, M., Zhang, J., Li H., Lemaster, D. M., and Hernández, G. (2014) Structural basis of conformational transitions in the active site and 80’s loop in the FK506-binding protein FKBP12. Biochem. J. 458, 525–536

Andersen, J. S., Mustafi, S. M., Hernández, G., and LeMaster, D. M. (2014) Statistical allosteric coupling to the active site indole ring flip equilibria in the FK506-binding domain. Biophys. Chem. 192, 41–48

Lipari, G., and Szabo, A. (1982) Model-free approach to the interpretation of nuclear magnetic resonance relaxation in macromolecules: 1. theory and range of validity. J. Am. Chem. Soc. 104, 4546–4559

Brath, U., Akke, M., Yang, D., Kay, L. E., and Mulder, F. A. (2006) Functional dynamics of human FKBP12 revealed by methyl 13C rotating frame relaxation dispersion NMR spectroscopy. J. Am. Chem. Soc. 128, 5718–5727

Brath, U., and Akke, M. (2009) Differential responses of the backbone and side-chain conformational dynamics in FKBP12 upon binding the transition-state analog FK506: implications for transition-state stabilization and target protein recognition. J. Mol. Biol. 387, 233–244

Sapienza, P. J., Mauldin, R. V., and Lee, A. L. (2011) Multi-timescale dynamics study of FKBP12 along the rapamycin-mTOR binding coordinate. J. Mol. Biol. 405, 378–394

Li, P., Ding, Y., Wu, B., Shu, C., Shen, B., and Rao, Z. (2003) Structure of the N-terminal domain of human FKBP52. Acta Crystallogr. D Biol. Crystallogr. 59, 16–22

Bracher, A., Kozany, C., Häkle, A., Wild, P., Zacharias, M., and Hausch, F. (2013) Crystal structures of the free and ligand-bound FK1–FK2 domain segment of FKBP52 reveal a flexible intra-domain hinge. J. Mol. Biol. 425, 4134–4144

Mandel, A. M., Akke, M., and Palmer, A. G. 3rd. (1996) Dynamics of ribonuclease H: temperature dependence of motions on multiple time scales. Biochemistry 35, 16009–16023

Tang, S., and Case, D. A. (2011) Calculation of chemical shift anisotropy in proteins. J. Biomol. NMR 51, 303–312

Clackson, T., Yang, W., Rozamus, L. W., Hatada, M., Amara, J. F., Rollins, C. T., Stevenson, L. F., Magari, S. R., Wood, S. A., Courage, N. L., Lu, X., Cerasoli, F. Jr., Gilman, M., and Holt, D. A. (1998) Redesigning an FKBP-ligand interface to generate chemical dimers with novel specificity. Proc. Natl. Acad. Sci. U.S.A. 95, 10437–10442

Yang, W., Rozamus, L. W., Narula, S., Rollins, C. T., Yuan, R., Andrade, L. J., Ram, M. K., Phillips, T. B., van Schravendijk, M. R., Dalgarno, D., Clackson, T., and Holt, D. A. (2000) Investigating protein-ligand interactions with a mutant FKBP possessing a designed specificity pocket. J. Med. Chem. 43, 1135–1142

Banaszynski, L. A., Chen, L. C., Maynard-Smith, L. A., Ooi, A. G., and Wandless, T. J. (2006) A rapid, reversible, and tunable method to regulate protein function in living cells using synthetic small molecules. J. Biol. Chem. 281, 31328–31336

Feng, H., Zeng, Y., Whitesell, L., and Katsanis, E. (2001) Stressed apoptotic tumor cells express heat shock proteins and elicit tumor-specific immunity. Blood 97, 3505–3512

Shen, Y., and Bax, A. (2010) SPARTA+: a modest improvement in empirical NMR chemical shift prediction. J. Biomol. NMR 48, 13–22

Han, B., Liu, Y., Ginzinger, S. W., and Wishart, D. S. (2011) SHIFTX2: significantly improved protein chemical shift prediction. J. Biomol. NMR 50, 43–57

Kohlhoff, K. J., Robustelli, P., Cavalli, A., Salvatella, X., and Vendruscolo, M. (2009) Fast and accurate predictions of protein NMR chemical shifts from interatomic distances. J. Am. Chem. Soc. 131, 13894–13895
59. Moon, S., and Case, D. A. (2007) A new model for chemical shifts of amide hydrogens in proteins. J. Biomol. NMR 38, 139–150
60. Haigh, C. W., and Mallion, R. B. (1979) Ring current theories in nuclear magnetic resonance. Prog. Nucl. Magn. Reson. Spectrosc. 13, 303–344
61. Case, D. A. (1995) Calibration of ring-current effects in proteins and nucleic acids. J. Biomol. NMR 6, 341–346
62. Chen, H., Mustafi, S. M., LeMaster, D. M., Li, Z., Héroux, A., Li, H., and Hernández, G. (2014) Crystal structure and conformational flexibility of the unligated FK506-binding protein FKB12.6. Acta Crystallogr. D Biol. Crystallogr. 70, 636–646
63. Blackburn, E. A., and Walkinshaw, M. D. (2011) Targeting FKBP isoforms with small-molecule ligands. Curr. Opin. Pharmacol. 11, 365–371