The Molecular Mechanism of Substrate Engagement and Immunosuppressant Inhibition of Calcineurin

Simina Grigoriu1,2, Rachel Bond3, Pilar Cossio4, Jennifer A. Chen3, Nina Ly3, Gerhard Hummer4, Rebecca Page2, Martha S. Cyert3*, Wolfgang Peti4,5*

1 Department of Molecular Pharmacology, Physiology and Biotechnology, Brown University, Providence, Rhode Island, United States of America, 2 Department of Molecular Biology, Cell Biology and Biochemistry, Brown University, Providence, Rhode Island, United States of America, 3 Department of Biology, Stanford University, Stanford, California, United States of America, 4 Laboratory of Chemical Physics, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, Maryland, United States of America, 5 Department of Chemistry, Brown University, Providence, Rhode Island, United States of America

Abstract

Ser/thr phosphatases dephosphorylate their targets with high specificity, yet the structural and sequence determinants of phosphosite recognition are poorly understood. Calcineurin (CN) is a conserved Ca2+/calmodulin-dependent ser/thr phosphatase and the target of immunosuppressants, FK506 and cyclosporin A (CSA). To investigate CN substrate recognition we used X-ray crystallography, biochemistry, modeling, and in vivo experiments to study A238L, a viral protein inhibitor of CN. We show that A238L competitively inhibits CN by occupying a critical substrate recognition site, while leaving the catalytic center fully accessible. Critically, the 1.7 Å structure of the A238L-CN complex reveals how CN recognizes residues in A238L that are analogous to a substrate motif, "LxVP." The structure enabled modeling of a peptide substrate bound to CN, which predicts substrate interactions beyond the catalytic center. Finally, this study establishes that "LxVP" sequences and immunosuppressants bind to the identical site on CN. Thus, FK506, CSA, and A238L all prevent "LxVP"-mediated substrate recognition by CN, highlighting the importance of this interaction for substrate dephosphorylation. Collectively, this work presents the first integrated structural model for substrate selection and dephosphorylation by CN and lays the groundwork for structure-based development of new CN inhibitors.

Citation: Grigoriu S, Bond R, Cossio P, Chen JA, Ly N, et al. (2013) The Molecular Mechanism of Substrate Engagement and Immunosuppressant Inhibition of Calcineurin. PLOS Biol 11(2): e1001492. doi:10.1371/journal.pbio.1001492

Academic Editor: Gregory A. Petsko, Brandeis University, United States of America

Received November 1, 2012; Accepted January 10, 2013; Published February 26, 2013

Introduction

The vast majority of eukaryotic proteins are phosphorylated, and this modification rapidly and reversibly modulates protein dynamics, interactions, activities, localization, and/or stability [1]. This essential regulation is carried out by the opposing activities of a large array of protein kinases, and a surprisingly small cadre of phosphoprotein phosphatases. Despite decades of investigation, basic questions about how these phosphatases act on phosphosites that share little similarity in primary sequence remain unanswered [2]. Here, we unveil a key mechanism of substrate recognition by calcineurin (CN) [3], the highly conserved Ca2+/calmodulin-activated ser/thr phosphatase [also called Protein Phosphatase 2B (PP2B) or Protein Phosphatase 3 (PP3)]; establish that structurally unrelated inhibitors of CN specifically disrupt this interaction; and show that substrates engaged at this site have additional interactions to orient the phosphosite toward the catalytic center of the enzyme. CN is ubiquitously expressed and is particularly abundant in the brain. By dephosphorylating a variety of protein substrates in response to Ca2+ signals, CN regulates development, learning and memory, cardiac function, and the immune response [3]. One of the best-studied activities of CN is its dephosphorylation of the nuclear factor of activated T-cell family of transcription factors (NFATc1-c4), which allows NFAT to translocate to the nucleus where it induces the expression of genes required for T-cell activation [4]. Because inhibition of NFAT signaling suppresses T-cell activation, natural products that specifically inhibit CN, cyclosporin A (CSA) and FK506, are widely prescribed as immunosuppressants to prevent posttransplant organ rejection [5]. CN is a member of the PPP family of protein phosphatases, which also includes PP1 and PP2A, among others. These enzymes contain structurally related catalytic domains, which rely on coordinated metal ions to directly bind phosphate and hydrolyze phosphoserine/phosphothreonine [2]. Despite their similarities, substrate recognition by these phosphatases is distinct, and natural products selectively inhibit each enzyme. In contrast to PP1 and PP2A, whose catalytic subunits combine with different regulatory subunits to create a suite of distinct holoenzymes [2], CN is always

Abbreviations: AID, auto-inhibitory domain; CN, Calcineurin; CNA, calcineurin catalytic A subunit; CNB, calcineurin regulatory B subunit; CSA, cyclosporin A; DFT, density functional theory; ITC, isothermal titration calorimetry; MD, Molecular Dynamics; NFAT, nuclear factor of activated T-cells; pNpp, p-nitrophenyl phosphate; PP1, Protein Phosphatase 1; PP2B, Protein Phosphatase 2B; PP3, Protein Phosphatase 3; PPP, phosphoprotein phosphatase; RII, RII regulatory subunit of PKA.

* E-mail: mcyyert@stanford.edu (MSC); wolfgang_peti@brown.edu (WP)
Author Summary

Transplantation medicine was revolutionized by the introduction of the immunosuppressant drugs cyclosporin A and FK506 that prevent rejection of transplanted organs by the recipient’s immune system. These drugs work by inhibiting calcineurin, a conserved protein phosphatase. Calcineurin regulates the immune response by dephosphorylating and activating the members of the NFAT family of transcription factors, which in turn activate genes required for the antigen-dependent stimulation of T-cells. Despite its biological and clinical importance, we have only a limited understanding of how calcineurin and other protein phosphatases interact with their substrates and target specific phosphorylated residues for dephosphorylation. Here, we determined the structure of calcineurin in complex with A238L, a viral peptide inhibitor of its function. This study shows that the viral peptide inhibits calcineurin not by targeting its active site but rather by occupying two critical substrate-binding regions of calcineurin (distant from each other and from the active site), thereby preventing its interaction with protein substrates. These findings allow us to present the first computational model of calcineurin bound to a phospho-substrate at its active site. Furthermore, by elucidating the structural basis for one particular mode of substrate–calcineurin interaction, this study reveals that both this viral peptide and immunosuppressant drugs inhibit calcineurin by blocking substrate access to a single critical region of the enzyme.

composed of a catalytic A subunit (CNA), bound to a regulatory B subunit (CNB) that binds four Ca\textsuperscript{2+} ions [6]. The C-terminus of CNA also contains a calmodulin-binding domain and an auto-inhibitory domain (AID), which regulate CN activity. Under basal Ca\textsuperscript{2+} conditions, the AID interacts with the catalytic center and prevents dephosphorylation. During signaling, increased Ca\textsuperscript{2+} levels cause Ca\textsuperscript{2+}-loaded calmodulin to bind CNA, which displaces the AID from the catalytic site and stimulates CN phosphatase activity [7].

Despite the critical biological importance of CN, the molecular mechanisms that allow CN to recognize and dephosphorylate specific protein substrates are still not well understood. Some but not all substrates contain a short CN-binding motif, termed “PxDxIT,” for its consensus sequence in NFATc1-c4 [8]. This sequence, which also occurs in scaffold proteins—that is, AKAP79 [9]—binds to CNA at a groove distal from the catalytic center. PxDxIT peptides bind with equal affinities to the inactive or active form of CN [10]. Because they do not occlude or alter the CN active site, they fail to inhibit dephosphorylation of either a model phosphopeptide substrate or a small molecule, p-nitrophenyl phosphate (pNpp) [11,12]. Thus, while this interaction can improve dephosphorylation efficiency by tethering a substrate to CN, it is not essential to the mechanism of dephosphorylation.

There has been limited insight into which substrate features do influence catalysis. Short peptides are not efficiently acted upon by CN; however, phosphopeptides that contain a basic residue at the −3 position relative to the phospho site show 4-fold better dephosphorylation [13]. Examination of one protein substrate, the RII regulatory subunit of PKA, defined a 19 mer as the smallest peptide that was robustly dephosphorylated (K_m = 26 μM; V_max = 1.7 μmol min \(^{-1}\) mg \(^{-1}\)), and showed that N-terminal residues (DLDV) lying 10 amino acids upstream of the phospho site, were critical for substrate recognition [14]. Subsequent studies of the NFAT family (NFATc1–c4) similarly identified a conserved CN-interaction site in these proteins,“ΦLxVP,” which interacts with Ca\textsuperscript{2+}/calmodulin-activated CN, and contributes to efficient dephosphorylation [8]. However, this sequence is significantly displaced from and C-terminal to CN-regulated phosphosites. Thus, the molecular details of the“ΦLxVP”–CN interaction and its role in dephosphorylation are still unclear.

Finally, CN is inhibited by the fungal-derived immunosuppressant drugs CSA and FK506, which bind the immunophilin proteins cyclophilin and FKBP, respectively, and engage CN as drug-immunophilin complexes. Structural analyses revealed that, unlike the AID, these drug-immunophilin complexes do not target the active site, but instead bind in a pocket ~50 Å away at the interface of the CNA/CNB subunits [6,15,16]. However, while pNpp can be readily dephosphorylated by CN/drug-immunophilin complexes, CN phosphoprotein and phosphopeptide substrates cannot [17], suggesting that the presence of the drug-immunophilin complexes impedes substrate/CN binding [12]. The detailed mechanism by which immunosuppressants achieve CN inhibition is still unclear [18,19].

Here, we describe the first high-resolution structure of CN bound to a physiological binding partner: the protein inhibitor A238L, from African swine fever virus (ASFV), a highly virulent double-stranded DNA virus that infects domestic pigs in Africa and Europe. Upon infection, A238L suppresses the host immune response by inhibiting both NFkB and CN. While A238L contains a “PxDxIT”-type anchoring sequence, the molecular mechanism by which A238L inhibits the enzyme is unknown [20]. Our studies demonstrate that A238L competitively inhibits CN and that residues in A238L (“FLCV”) directly compete with substrates for binding to a substrate-recognition cleft, the “ΦLxVP” cleft, in activated CN. Our 1.7 Å crystal structure of the CN-A238L complex reveals the molecular interactions that mediate this key element of substrate recognition and surprisingly shows that A238L effectively inhibits CN not by blocking and occluding the active site, which is fully accessible in the complex, but instead by binding and blocking the “ΦLxVP” substrate recognition groove. This structure enabled molecular dynamics (MD) modeling of a minimal substrate, the RII peptide, bound to CN. The model reveals that interactions at the hydrophobic substrate-recognition groove are augmented by charged interactions at position −3 upstream of the phospho site [13]. These studies provide the first structural insights into active site substrate engagement for any ser/thr phosphatase. Thus, this work elucidates a key mechanism by which CN recognizes substrates and provides structural insights into the presentation of phosphosites to the active site during dephosphorylation. Furthermore, the CN-A238L structure also unequivocally shows that the LxVP sequence and immunosuppressants bind to the identical site on CN. Thus, these studies establish the mechanism of action of these drugs and lay the foundation for renewed efforts in the structure-based targeted design of novel CN inhibitors.

Results

A238L Engages CN Via Both an LxVP and a PxDxIT Docking Motif

To understand how substrates engage CN, we investigated A238L (aa 157–239, Malawi LIL20-1 strain), a protein inhibitor from the African swine fever virus (ASFV) that suppresses the host immune response by inhibiting both NFkB and CN [20]. Two substrate motifs, PxDxIT and LxVP, have been shown to bind to CN [8]. While it was known that a C-terminal fragment of A238L contains a PxDxIT motif that anchors the protein to CN [21], we showed that excess amounts of a peptide that encodes the LxVP...
docking motif from NFATc1, but not a mutant (LxVPc1mut
LAVP→AAA) also disrupted A238L-CN binding (Figure S1A). This establishes that A238L also contains an LxVP motif. The only sequence in A238L157–239 that is similar to the LxVP motif is LCVK (Figure 1A). To verify that this sequence binds CN, we generated a GST fusion protein containing the LCVK sequence from A238L (but lacking the PKIHT sequence) and showed that it forms a complex with recombinant CN that is disrupted by the addition of a peptide encoding the LxVP motif from NFATc1, but not a mutant version of this peptide (Figure 1B). In contrast, a GST fusion protein with a mutated sequence (FLCVK→AAGCA) did not bind CN (Figure 1B). Next, we expressed full-length and truncated forms of A238L in yeast as GST fusions and measured their ability to inhibit expression of a CN-dependent reporter gene, CDRE-lacZ [22]. This analysis showed that a shorter A238L fragment, A238L200–239, which includes both CN binding motifs (PKIHT211 and LCVK232), is sufficient to inhibit CN (Figure 1C). Moreover, this same fragment competitively inhibits dephosphorylation of an RII phosphopeptide substrate by CN with a K_d of 0.37 nM (Figures 1D and S1B; RII contains an LxVP motif) [14]. A238L200–239 also forms a very tight complex with CN, with a dissociation constant (K_d) of 4 nM as determined using isothermal titration calorimetry (ITC) measurements (Figures 1E and S2, and Table 1). In agreement with this result, the CN-A238L complex is a stable trimer (CNA1–370,B1–170–
A238L200–239), hereafter referred to as CN-A238L, as evidenced by the elution of the complex in a single peak during size exclusion chromatography at the expected elution volume for a 67.2 kDa trimeric complex. Taken together, these observations demonstrate that the competitive protein inhibitor A238L200–239, hereafter referred to as A238L, binds tightly to CN using both a PxIxIT-type anchoring sequence and an LxVP motif.

Structure of the CN-A238L Heterodimer/Inhibitor Complex

To elucidate the molecular mechanism of A238L binding and inhibition, we determined the 1.7 Å crystal structure of the CN-A238L heterodimer/inhibitor complex (CNA/CNB/A238L). The structure of the CNA/B heterodimer in complex with A238L is virtually identical to CNA/B heterodimer structures from previous reports [6,9,15,16,23–25]. CN residues 1–13, CNB residues 1–5/161–170, and A238L residues 200–204/235–239 were not visible in the electron density map and thus were not modeled. The absence of electron density for A238L residues 200–204 (N-terminal to the LxVP motif) and A238L residues 235–239 (C-terminal to the LxVP motif) suggests that these regions remain flexible upon complex formation and do not contribute to CN binding. A238L binds CN in a largely extended conformation, stretching from the PxIxIT binding site to the CNA/CNB interface and then looping back along the CNA/B interface to occupy the newly identified LxVP binding pocket (Figure 2A, B). As a consequence, the CN-A238L interaction buries 3,083 Å^2 of solvent accessible surface area (SASA). Because A238L potently inhibits CN, it was predicted that A238L would bind and occlude the CN active site. However, the structure reveals that the CN active site is fully accessible in the CN-A238L complex (Figure 2C, left). Moreover, the key catalytic residues are structurally invariant when compared with those in previously determined CN structures [6,16,24], suggesting that CN in the CN-A238L complex is catalytically active (Figure 2C, right). The ability of CN to dephosphorylate small molecule substrates was confirmed using p-nitrophosphatyl phosphate (pNpp) dephosphorylation assays (Figure 2D). In fact, A238L and A238LPKIHTmut both increased the rate of pNpp hydrolysis compared to untreated CN, consistent with previously reported rate increases by both a peptide containing the LxVP sequence from NFATc1 as well as by the immunosuppressants FK506 and CSA [12,17]. Thus, the CN-A238L complex retains full catalytic activity. Together, these data show that A238L does not inhibit CN by blocking the catalytic center, but instead utilizes an alternative mechanism.

The CN-A238L PxIxIT Interaction

The interaction of A238L with CN at the PxIxIT binding pocket buries 886 Å^2 of SASA and is mediated by A238L residues 206–211 (PKIHT, the PxIxIT motif from A238L), which form a short β-strand that hydrogen bonds with ß14 of CN to extend one of its central β-sheets (Figure 3A–C). As observed for other CN-PxIxITpeptide complexes, hydrophobic contacts are the dominant determinants of specificity in the PKIHT interaction between A238L and CN (Figure 3B). The PxIxIT interaction in the CN-A238L complex is nearly identical to those observed in the CN-PVIVITpeptide and the CN-AKAPpeptide complexes [9,24], demonstrating that PxIxIT sequences, contained in many CN-interacting proteins, likely bind CN in a similar manner (Figure 3C,D). The CN-A238L complex also allows for the comparison of interactions outside the PxIxIT binding groove (Figure 3C). As expected, these interactions are more variable. For example, while the C-terminal residues of both peptides angle down away from CN, the A238L chain continues upwards toward the CNA/B interface. Furthermore, both Pro13PKIHT and Cys213A238L, but not Thr345Kas, bind in a shallow hydrophobic pocket comprised of the methyl groups of Lys513CNB, Gln335CNB, and Asn335CNB, suggesting that this pocket might be engaged by other endogenous substrates to enhance binding (Figure 3C). These results illustrate how sequence variations not only in the PxIxIT sequence itself but also in flanking residues may fine-tune the affinity of PxIxIT sequences for CN [26].

The CN-A238L LxVP Interaction

A short motif, LxVP, mediates interaction of several substrates, as well as A238L, with Ca^2+/calmodulin-activated CN [12]. However, until now, the molecular interactions that mediate LxVP recognition by CN have remained unknown (Figure 4). In the CN-A238L complex, the LxVP motif from A238L, LAVP→AAA, is bound to CN (Figure 4A), revealing the very hydrophobic LxVP binding pocket at the CNA/CNB interface in CN (Figure 4B). When bound to CN, the A238L LCVK residues are extended, burying 728 Å^2 of SASA. Unlike the CN PxIxIT binding pocket, which is comprised of residues only from CNA, the CN LxVP binding pocket contains residues from both CNA and CNB. Leu229A238L, which becomes 92% buried upon complex formation, makes hydrophobic contacts with residues from both CNA and CNB subunits: Trp532CNB and Phe536CNB, which, when mutated, alter binding to the LxVP sequence from NFATc1 [12], as well as Leu115CNB, Met118CNB, and Val119CNB (Figure 4C, left). Similarly, Val231A238L becomes 97% buried when bound to CN, forming hydrophobic interactions with Tyr344CNB, Leu343CNB, Pro334CNB, Trp352CNB, and Leu125CNB (Figure 4C, right). In addition, Cys230A238L (the “x” in LxVP) forms hydrogen bonds with Trp532CNB and Asn122CNB (Figures 4D and S3A). Finally, Lys232A238L, which is a non-canonical residue in the LxVP motif as it has a “K” instead of the expected “P,” does not interact with CN and thus is not important for CN recognition by A238L. Consequently, this work shows that residues Leu229A238L and Val231A238L, which make significant interactions with both CNA and CNB, are the key residues that mediate binding to the LxVP interaction pocket. Notably, Phe228A238L, which is immediately N-terminal to the LxVP
binding motif, also contributes to CN binding (Figure 4A, B), fitting into a deep pocket formed by the loops connecting EF-hands 1 and 2 and EF-hands 3 and 4 of CNB. This results in a 25% increase in the SASA buried at this site (FLCVK buries 911 Å² SASA). Critically, multiple LxVP sites are immediately preceded by an aromatic residue (Phe or Tyr), which may act as a binding strength enhancer, as mutation of this aromatic residue weakens the LxVP interaction [12]. Thus, a subset of LxVP sites, including the one in A238L, is best described as W\textsubscript{LxVP}.

A238L Residues Also Contribute to the LxVP Binding Pocket

The W\textsubscript{LxVP} binding groove is comprised of residues from both CNA and CNB, in the CN-A238L complex. Unexpectedly, however,

Table 1. Thermodynamic parameters and dissociation (K\textsubscript{D}) and inhibition (K\textsubscript{I}) constants for CN\textsubscript{A1–391/B1–170} with A238L\textsubscript{200–239} wild-type, and A238L PxIxIT and LxVP mutants, derived from ITC experiments at 25 °C or enzyme assays performed with RII at 37 °C.

| Complex               | K\textsubscript{D} (nM) | ΔH (kcal mol\textsuperscript{-1}) | ΔS (kcal mol\textsuperscript{-1}) | ΔG (kcal mol\textsuperscript{-1}) | K\textsubscript{I} (nM) |
|-----------------------|-------------------------|---------------------------------|---------------------------------|---------------------------------|-------------------------|
| CN:A238L\textsubscript{WT} | 4±1                     | −40.3±4.9                       | 28.7±4.9                        | −11.5±0.2                       | 0.37±0.03               |
| CN:A238L\textsubscript{PxIxITmut} | 624±26                 | −34.3±0.9                       | 25.9±0.9                        | −8.5±0.0                        | 15±1                    |
| CN:A238L\textsubscript{LxVPmut} | 803±26                 | −14.0±0.1                       | 5.7±0.1                         | −8.3±0.0                        | 7,700±3,000             |

Thermodynamic and dissociation constant data represent mean values ± one s.d. for triplicate measurements except A238L\textsubscript{WT}, which was performed 5 times. Inhibition constants are mean values ± one s.e.m. from three independent experiments.

doi:10.1371/journal.pbio.1001492.t001
a few A238L residues also contribute to the ΦLxVP binding pocket. Specifically, A238L, which projects upwards from the PxIxIT site towards the CNB interface, forms a tight 180° turn at Asn227A238L (Figures 4D and S3A). This kink is essential for WlVP binding, as it redirects the WlVP sequence (FLCVK) back toward its docking site on CN at the CNA/B interface. This enables the A238L residues that immediately precede the WlVP to interact directly with the CN-bound WlVP residues and contribute to the WlVP binding site. Thus, in addition to the multiple interactions observed with residues from CNA and CNB, Val231A238L (the “V” in the LxVP motif) also makes intramolecular hydrophobic contacts with Leu221A238L. In addition, the side chain Gln224A238L hydrogen bonds with the backbone amide of Val231A238L (Figures 4D and S3A). Therefore, residues from all three proteins—CNA, CNB, and A238L—function to keep Val231A238L occluded from solvent in the bound conformation. In addition, although Cys230A238L is the “x” in the LxVP motif, this residue also makes multiple intramolecular interactions that help stabilize the A238L bound conformation. Specifically, the amide nitrogen of Cys230A238L hydrogen bonds with the carbonyl oxygen of Phe228A238L. Cys230A238L also forms thiol hydrogen bonds with the amide nitrogen of S226A238L and the carbonyl of N225A238L. These intramolecular interactions explain why Cys230A238L is still nearly completely buried (70%) in the CN-A238L complex even though its side chain points away from the LxVP docking groove. Thus, although it is not yet known how similar the CN-LxVP interaction of CN-A238L is with that of other LxVP docking motifs from substrates, our structure suggests that residues flanking the LxVP sequence may also modulate the affinity of this motif for CN.

Although the most extensive interactions between A238L and CN occur at the PxIxIT and ΦlVP binding grooves, additional, largely polar intra- and intermolecular interactions outside of these docking sites also contribute to A238L binding. For example, the interactions that stabilize the A238L kink (the 180° tight turn at Asn227A238L) which enables the rest of A238L to point back toward the LxVP
binding pocket) are mediated by an extensive network of more than 10 hydrogen bonds, the center of which is Asn\textsubscript{122}CNB (Figures 4D and 3SA). The side chain amide nitrogen and carbonyl of Asn\textsubscript{122}CNB form hydrogen bonds with the backbone carbonyl and backbone amide nitrogen of two residues that border the kink, L\textsubscript{229}A\textsubscript{238}L and Asn\textsubscript{225}A\textsubscript{238}L, respectively. Similarly, A\textsubscript{238}L residues C-terminal to the PxIxIT motif (212GCEDNVY218) also interact with CNA through main chain/side chain/side chain hydrogen bonds (Figure S3B). Although the two dominant A\textsubscript{238}L:CN PxIxIT and LxVP sequences are important for CN binding, as the PxIxIT and LxVP mutations increase the $K_D$ similarly by $150\times$ and $200\times$, respectively (Table 1 and Figure S2).

Next, we investigated the contribution of the individual A\textsubscript{238}L motifs toward the inhibition of CN. A\textsubscript{238}LPKIIIT\textsubscript{mut}, which still contains the FLCVK sequence, competitively inhibits dephosphorylation of RII by CN with $K_i = 15$ nM (Table 1 and Figure 4E) [14]. In contrast, A\textsubscript{238}LFLCKVK\textsubscript{mut}, which still contains the PKIIIT sequence, inhibited CN very poorly, with $K_i = 7,700$ nM, a 20,000-fold decrease in inhibitor efficacy compared to wild-type A\textsubscript{238}L (Table 1 and Figures 3E and S4). In addition, when expressed in HEK293T cells in which NFAT signaling had been stimulated, both A\textsubscript{238}L mutants reduced the activity of an NFAT-dependent reporter gene. At roughly equal levels of protein expression, neither A\textsubscript{238}LFLCKVK\textsubscript{mut} nor A\textsubscript{238}LPKIIIT\textsubscript{mut} were as effective as wt-A\textsubscript{238}L in NFAT inhibition (Figure 5A, B). Taken together,

**Figure 3. CN-A238L interactions: The PxIxIT substrate binding site.** (A) Close-up view of the CN PxIxIT substrate binding site. A238L PKIIIT is shown as magenta sticks and labeled; CNA is shown as a grey surface. (B) Same view as (A), with a transparent CNA surface. Individual CNA residues that participate in the interaction with PKIIIT\textsubscript{A238L} are shown as grey sticks. (C) Superposition of the PKIIIT\textsubscript{A238L} motif (purple) with a synthetic PVIVIT peptide (orange) and the IAIIIT docking site of AKAP79 (green) bound to CN. Pro13\textsubscript{PVIVIT} and Cys213A238L are shown as sticks. (D) Overlay as in (C) but illustrated as sticks. Corresponding “variable” residues Ile\textsubscript{209}A238L, Val328CNA, and Ile341AKAP79 (the second “x” in PxIxIT) participate in the same hydrophobic interaction with Val328CNA (dotted lines). (E) Secondary plot of $K_i^{app}$ as a function of [RII] for inhibition of CN by A238LFLCKVK\textsubscript{mut}, which retains the PKIIIT site. Data show a linear dependence characteristic of competitive inhibition, with $K_i = 7,700\pm 3,000$ nM. $K_i^{app}$ values were obtained from the nonlinear fit of Figure S4A. Points represent averages ± s.e.m. doi:10.1371/journal.pbio.1001492.g003
these results show that in A238L, FLCVK, but not PKIIIT, is required to competitively inhibit dephosphorylation of the RII phosphopeptide, whereas both sequences contribute to reducing CN-mediated dephosphorylation of substrates, such as NFATs, that contain an LxVP substrate recognition motif, and also require PxIxIT-mediated anchoring to CN. Therefore, A238L does not directly inhibit CN activity by blocking its active site, but instead inhibits CN via a model of steric occlusion where it blocks the access of substrates to key binding sites on CN, similar to the steric occlusion of substrate selection previously reported for Protein Phosphatase 1 [27].

Model of CN-Substrate Dephosphorylation

The CN-A238L structure revealed the molecular interactions that mediate CN recognition of the LxVP motif, and we showed that the RII phosphopeptide requires this interaction for its dephosphorylation; therefore, we used bias-exchange metadynamics MD [28] to generate the first detailed model of a substrate, the RII phosphopeptide, bound to activated CN and poised for dephosphorylation (Figure 6A). The LxVP sequence of the RII peptide (81LxVP100) is separated from the phospho-Ser95 residue by nine amino acids, which, due to the distance between the LxVP-binding site and the catalytic center, limits the potential conformations this peptide can adopt during dephosphorylation. Our model of the CN-RII peptide complex shows that the interaction of RII (81LxVP100) with the CN LxVP site is very similar to that observed in the CN-A238L complex. It also shows that additional electrostatic and hydrophobic interactions stabilize interactions between CN and the intervening RII residues, which function to guide p-Ser95 to the CN active site. In particular, RII residues Asp91 and Arg92 form a number of key side chain hydrogen bonds and salt bridge interactions with CN (Figure 6B). This is consistent with previous work suggesting that Arg in the −3 position is a strong positive determinant of dephosphorylation efficiency by CN [13]. The model also shows that the nine intervening residues between the LxVP and phospho-Ser95 are essential for spanning the distance between the CN LxVP binding and active sites. This demonstrates that substrates with phosphorylated residues that are close to the LxVP site (e.g., within six or fewer residues) are unlikely to be dephosphorylated by CN as they will be unable to reach the active site.

Discussion

Our biochemical and structural studies reveal the mechanism by which A238L, a protein made by African swine fever virus that dampens the host immune response during infection, potently inhibits CN. The structure of CN (CNA1–370 and full-length CNB-subunit) in complex with A238L shows that 30 amino acids of A238L form a tight complex with CN. We conclude that an equivalent complex forms with full-length CN, as neither the presence of the AID nor additionally described inhibitory sequences [29] interfere with the inhibition of CN by A238L in
substrate dephosphorylation by steric occlusion of substrate binding sites has only been directly observed in one other ser/thr protein phosphatase, that of PP1, which, like CN, is one of the key members of the PPP family [27]. Thus, our results establish that this mechanism, whereby enzyme activity is modulated via substrate access rather than through active site inhibition or allostery, is likely utilized by the entire PPP family.

Although the structure of CN bound to a bona fide substrate has yet to be determined, the CN-A238L structure and our CN-RII substrate model significantly advance our understanding of how CN uses two complementary strategies to recognize substrates. First, the PxIxIT anchoring motif is used by protein scaffolds, inhibitors, and some substrates to form a stable interaction with CN by docking to a site that is available regardless of the activation state of the enzyme. For PxIxIT-containing substrates, the strength of this anchoring modulates the Ca\(^{2+}\)-concentration dependence of their dephosphorylation, but does not directly contribute to recognition of phosphosites during the dephosphorylation reaction [26,30]. Second, CN overcomes the limited specificity of its catalytic site by recognizing specific residues, such as “ΦlxVP,” which are distal to the phosphosite. This interaction is likely required for the dephosphorylation of multiple substrates, as it is this site that is targeted by multiple inhibitors, including A238L and the immunosuppressants CSA and FK506 (Figure 6C).

This work also reveals interactions that contribute to phosphosite selection by CN. The “ΦlxVP” binding site is now molecularly defined as a hydrophobic binding surface composed of residues from the CNA and CNB subunits, which becomes accessible after binding of Ca\(^{2+}\)-loaded calmodulin displaces the C-terminal AID domain from the active site [12]. Our structure and the RII model show that the substrate residue dephosphorylated by CN (hereafter referred to as pS/pT) must be a minimum of 9–15 residues away from the Φ lxVP sequence, and be in an extended conformation in order for the pS/pT residue to reach the catalytic site. In fact, the pS/pT residues in the RII peptide and the substrate RCAN1 are only 9 and 10 residues C-terminal to the “Φ” in their LxVP motifs, respectively (Figure 6A, B, D) [14,31]. Our model also shows that additional electrostatic interactions, especially at basic residue at position −3, help orient the phosphosite in RII toward the catalytic center. In addition, the C-terminus of the RII peptide lies in a groove identified in other PPPs, namely PP1, to be a substrate recognition groove (hydrophobic groove in PP1), suggesting that this groove functions in a similar manner in CN and thus likely for the entire family of PPPs [2,32].

While our model provides fundamental insights into phosphosite selection by CN for those substrates in which the phosphosite is 9–10 residues C-terminal to LxVP sequence, other substrate access near the catalytic center is likely also important for substrate recognition. This is because the spacing between experimentally determined “LxVP” and phosphosites residues is quite variable. For example, substrates such as NFAT and KSR2 have pS/pT sites that are 10–100 s of residues away from the known LxVP sequences [33–37]. In these cases, one or several pS/pT sites are found in long, extended sequences that are predicted to be unstructured, suggesting that these regions are dynamic when bound to CN and that interactions at the LxVP and PxIxIT sites tether the substrate to CN allowing the phosphosite(s) to encounter the catalytic site and be dephosphorylated (Figure 6D). A detailed understanding of how phosphosites that are distal from the LxVP and PxIxIT motifs engage the CN catalytic center is an active area of investigation.

Finally, these studies also provide critical new insights into the mechanism by which the immunosuppressants CSA and FK506...
Figure 6. Potential interaction modes of CN substrates/regulators with CN. (A) The CN-RII peptide complex obtained by MD. Colors as in Figure 2A, C. CN is shown in surface representation and the RII peptide in dark green with the LxVP motif (LDVP) and phospho-Ser95 as green sticks. LDVP is bound to the LxVP binding pocket (light green), and phospho-Ser95 is bound in the CN active site (cyan). (B) Electrostatic interactions between CN and the RII peptide. The CN electrostatic surface has positively and negatively charged areas colored blue and red, respectively. The LxVP motif and residues in RII that participate in polar interactions with CN are shown as green sticks. (C) Features of selected CN substrates and regulators, including substrates tested in this work (NFAT, Crz1, and the RII peptide). PxIxIT and LxVP motifs are highlighted in yellow and green, respectively, with intervening residues in grey. Regions containing S-T residues that are dephosphorylated by CN are pink. (D) Potential modes of interaction of CN with various binding partners. CN is shown in grey, with the active site in cyan, the PxIxIT docking site in yellow, and the LxVP docking site in green. CN binding partners are shown in blue, with PxIxIT and LxVP motifs in purple and phosphorylated regions shown as red circles. The residues between the two CN docking motifs, or between one docking motif and regions dephosphorylated by CN, are represented as coils, as they are predicted to be unstructured in solution. A238L is the CN-A238L crystal structure.
doi:10.1371/journal.pbio.1001492.g006
inhibit CN. Our structure reveals that the location of the LxVP substrate-binding site on CN is identical to that of the CSA/FK506 binding site (Figure 7A, B). Furthermore, we identified several contacts common to all of these interactions. In particular, multiple residues from the cyclic immunosuppressant molecules overlap nearly perfectly with key residues of the A238L LxVP sequence, LCVK, especially at Leu229A238L and Val231A238L, and occupy hydrophobic surfaces formed by Trp352 and Phe356 of CNA and Met118 and Val119 of CNB (Figure 7C). These findings indicate that disrupting the interaction between CN and LxVP motifs found within substrates is sufficient for inhibiting the dephosphorylation of LxVP-containing substrates, and explains the molecular mode of action of these drugs. Furthermore, the ability of FK506 and CSA to antagonize all known functions of CN suggests that every substrate contains at least one LxVP motif and that interaction of this motif with CN is essential for substrate dephosphorylation. Consequently, our high-resolution 3-dimensional structure of the CN-A238L complex opens up a new avenue for the development of a novel class of powerful CN inhibitors that selectively and potently bind the CN LxVP binding site. Both CN substrate-binding pockets, but especially the LxVP pocket, present excellent targets for the development of immunosuppressive drugs. Clearly, such drugs will also be an extremely powerful tool for investigating the many regulatory processes, including immune activation and cardiac hypertrophy, that are driven by CN.

**Materials and Methods**

**Proteins**

CNA (residues 1-370 or 1-391), CNB (residues 1-170), and wt- and mutant A238L (residues 200-239) were subcloned into vectors containing either an N-terminal His$_x$- or GST-tags and expressed in E. coli. To form the CN-A238L (CNA, 1-370; CNB, 1-170; A238L, 200-239) complex used for X-ray crystallography, His$_x$-tagged CN was purified over a HisTrap HP column and directly eluted into an at least 2-fold molar excess of previously purified A238L. His$_x$-tag cleavage and subtraction purification were performed, and the complex was further purified by SEC (Superdex 200 26/60) in 20 mM Tris pH 8.0, 50 mM NaCl, 0.5 mM TCEP, and 1.5 mM CaCl$_2$. Additional cloning, protein expression, and protein purification procedures are provided in Text S1.

**Crystallization and Structure Determination**

The CN-A238L complex (~7 mg/ml) formed thin plate crystals in 0.16 M ammonium citrate and 20% (w/v) PEG3350 at 22°C. Crystals were obtained using the sitting drop vapor diffusion method (three-well Inteliplate, Art Robbins), with 0.6 µl drops containing a 1:2 ratio of precipitant solution to protein and 50 µl of precipitant solution in the reservoir. Crystals were cryo-protected using a 10 min soak in mother liquor supplemented with 30% glycerol and immediately flash frozen in liquid N$_2$. Data were collected at the NSLS X25 beamline at Brookhaven National Laboratory. Crystals of CN-A238L formed in space group P2$_1$, with unit cell dimensions $a = 72.69$ Å, $b = 48.98$ Å, $c = 82.44$ Å, and $\beta = 104.4$. Data were indexed, integrated, and scaled with DENZO and SCALEPACK as part of HKL2000 [38]. The structure of CN-A238L was determined to 1.7 Å by molecular replacement using a CN heterodimer molecule (CNA/B; PDBID 1AUI [6]; the auto-inhibitory domain was omitted) as the search model. The final model of the CN-A238L complex was obtained using iterative rounds of refinement in Phenix [39] and model building in Coot [40], with TLS used in the final round of refinement. The asymmetric unit contains one copy of the CN-A238L heterodimer/inhibitor complex. The final structure refined to a final R factor of 15.8% (R$_{free}$ = 17.8%). CN-A238L crystals formed at pH 5, leading to protonation of active site residues and thus displacement of the positively charged active site metal ions, as previously seen by Jin & Harrison, who crystallized CN at pH 4.6 [16]. One hundred percent of all residues are in the allowed region of the Ramachandran diagram. Structure validation and stereochemistry analysis was performed with Molprobity [41]. Details and statistics of data analysis and model building are provided in Table S1 and Text S1. A stereoview of the A238L electron density is in Figure S5.

**MD Simulations**

The CN-A238L structure was used as the starting structure for the MD simulations. The phosphoserine (pSer) was placed in the active site according to the PDB structure 1TCO, together with the expected OH ligand [23]. We used density functional theory (DFT) with the B3LYP functional [42], as implemented in TURBOMOLE [43], to obtain an energy minimized structure of the CN active site, including the partial charges of the active site metal ions and their ligands. During the MD simulations, the active site region was restrained to the DFT-optimized configuration. Initial structures of RII bound to CN were built according to the backbone coordinates of A238L at the LxVP site, and the last eight residues of the auto-inhibitory domain in PDB structure 1AU1 [6]. Ten different starting models for the intervening sequence $^{86}$IPGRFD$^{91}$ were constructed using the ROSETTALoop model tool [44]. The resulting 10 CN-RII complexes were
inserted into rectangular boxes (3 x 3 x 12 nm³) with 22,881 TIP3P water molecules [45] and eight Na⁺ ions each for electroneutrality. MD simulations were performed using GROMACS4.5.3 [46], the Amber ff99SB-ILDN force field [47], particle mesh Ewald summation [48] using a 0.12 nm grid spacing, a time step of 2 fs and a 0.9 nm real-space cutoff at a constant temperature [49] of 300 K and at a pressure [50,51] of one bar. After 2 ns of equilibration of each of the 10 starting models, bias-exchange metadynamics [28] using the PLUMED1.3.0 plugin [52] was used to accelerate the conformational sampling. In 9 of the 10 replicas, the Ψ dihedral angles of RII residues B68, Pro87, Arg96, Phe90, Asp91, Arg92, Arg93, Cys97, and Ala98, respectively, were biased, while replica 10 was kept unbiased. To ensure RII stayed bound to CN during the conformational sampling, the F²-pSer and LxVP interactions were harmonically restrained. Convergence was reached after 1.15 μs (115 ns replica), as judged by the free energy profiles of the biased Ψ. The RII structures of the unbiased replica were clustered according to a 1.2 Å backbone root-mean-square-distance (RMSD) threshold. The most populated cluster contains 58% of all structures, from which a representative was selected with the lowest Coulomb energy. A different clustering technique using reweighting of the biased replicas [53] led to similar results.

Isothermal Titration Calorimetry

ITC experiments were performed at 25°C using a VP-ITC microcalorimeter (GE Healthcare). All protein samples were equilibrated in ITC buffer (20 mM Tris pH 7.5, 150 mM NaCl, 1.5 mM CaCl₂, 0.5 mM TCEP). Wild-type or mutated A238L was titrated into CN A1–391/B1–170. Titrant (10 μM per injection) was injected into the sample cell over a period of 20 s with a 250 s interval between titrations to allow for complete equilibration and baseline recovery. Twenty-eight injections were delivered during each ITC experiment, and the solution in the sample cell was stirred at 307 rpm to ensure rapid mixing. Data were analyzed with one set of sites binding model, based on the 1:1 stoichiometry observed in the crystal structure, using Origin 7.0 (OriginLab).

CN Activity Assays with RII

The rate of RII phosphopeptide dephosphorylation by CN was determined by measuring the total phosphate released over four time points (total 5–20 min). Reaction rates were linear over this time period and constituted less than 1% of product formation. The 50 μl reactions contained assay buffer (50 mM Tris pH 7.4, 100 mM NaCl, 6 mM MgCl₂, 0.5 mM CaCl₂, 0.1% PEG 3250, 0.5 mM DTT), 10 nM CN, wt- or mutant A238L (0–10 μM), and 50–100 μM RII phosphopeptide. Reactions were performed at 37°C and were initiated by the addition of 10× RII and terminated using 100 μl Biomol Green Reagent (Enzo Life Sciences). After color development for 20 min, absorbance was measured at 595 nm and compared to phosphate standards of known pNp concentration. Absorbance units were converted to pNp concentration. Values are summarized in Table 1.

Accession Numbers

The structure factors and coordinates for the CN-A238L complex have been deposited with the Protein Data Bank with accession number 4FOZ.

Supporting Information

Figure S1 A238L binds CN via a PxIxIT and LxVP motif. (A) Recombinant S-tagged A238L [157-239] incubated with GST-CN and CNB co-purifies with GST-CN (lane 5). Incubation with excess peptides encoding the LxVP site from NFATc1 (lane 1) or the high-affinity PxIxIT peptide PVIVIT (lane 3) interferes with A238L-CN binding. Control peptides (lanes 2 and 4) do not interfere with binding. (B) Plot of CN rate as a function of A238L [290-239] concentration at different RII concentrations ranging from 50–1,000 μM. Curve fit obtained by nonlinear regression using the Morrison equation to account for tight binding inhibition. Error bars indicate s.e.m. from three independent experiments. (TIF)

Figure S2 Role of the PxIxIT and LxVP sites in the CN-A238L interaction. Raw isothermal titration calorimetry data (upper panels) and derived binding isotherms plotted versus the molar ratio of titrant fit using a one-site model (lower panels) for CN A1–391/B1–170 titrated with: (A) WT A238L, (B) A238L PxIxIT mutant (PKIIIT mutated to AKAIAA), and (C) A238L LxVP mutant (FLCVK mutated to AACAA). Thermodynamic data and K_D values are summarized in Table 1. (TIF)

Figure S3 A238L-CN polar interactions. (A) Stereo-view of the FLCVK_A238L interface. CN residues participating in the direct interaction are in orange. Close-up view of the FLCVK_A238L interface is shown in (B). A238L residues in the interface are shown in orange, whereas CN residues are colored in gray (TIF)
interaction are shown as grey (CNA) or beige (CNB) sticks, with A238L residues shown in purple. The multiple intramer and intermolecular hydrogen bonds that stabilize the A238L kink are shown as black dotted lines. (B) A238L residues immediately C-terminal to the $^{209}$PKHI$^{211}$ motif, $^{213}$GEGDNV$^{218}$, are illustrated as sticks and labeled. CNA residues that interact with these A238L residues are also shown as sticks (black). Hydrogen bonds/salt bridge interactions are indicated by black dashed lines.

**Figure S4** A238L LxVP motif mutant weakly inhibits RII dephosphorylation. (A) Dose-response plot of CN rate as a function of A238L$^{FLCVK}$ at different RII concentrations ranging from 50–1,000 μM. Curves were fit by nonlinear regression using the Morrison equation. Error bars indicate one s.d. from three independent experiments. (B) Plot of CN rate as a function of [RII]. Data fit the Michaelis-Menten model for competitive inhibition. Points represent averages ± s.d. from three independent experiments. Concentrations of A238L$^{FLCVK}$ are indicated. (TF)

**References**

1. Olsen JV, Blagoev B, Guad F, Macle B, Kumar C, et al. (2006) Global, in vivo, and site-specific phosphorylation dynamics in signaling networks. Cell 127: 635–648.
2. Shi Y (2009) Serine/threonine phosphatases: mechanism through structure. Cell 138: 460–481.
3. Aramburu J, Rao A, Klee CB (2000) Calcineurin: from structure to function. Curr Top Cell Regul 36: 237–295.
4. Crabtree GR, Schreiber SL (2009) SnapShot: Ca$^{2+}$-calcineurin-NFAT signaling. Cell 138: 210–211.
5. Musson RE, Smit NP (2011) Regulatory mechanisms of calcineurin phosphatase activity. Curr Med Chem 18: 301–315.
6. Kissinger CR, Parge HE, Knighton DR, Lewis CT, Pelletier LA, et al. (1995) Crystal structures of human calcineurin and the human FKBP12-FK506-calcineurin complex. Nature 378: 641–644.
7. Yang SA, Klee CB (2000) Low affinity Ca$^{2+}$-binding sites of calcineurin B mediate conformational changes in calcineurin A. Biochemistry 39: 16147–16154.
8. Roy J, Ceyt MS (2009) Cracking the phosphatase code: docking interactions determine substrate specificity. Sci Signal 2: re9.
9. Li H, Pink MD, Murphy JG, Stein A, Dell’acqua ML, et al. (2012) Balanced and site-specific phosphorylation dynamics in signaling networks. Cell 127: 635–648.
10. Stathopoulos AM, Ceyt MS (1997) Calcineurin acts through the CRZ1/TCN1-encoded transcription factor to regulate gene expression in yeast. Genes Dev 11: 3432–3444.
11. Griffith JP, Kin JL, Kim EE, Smitzak MD, Thomson JA, et al. (1995) X-ray structure of calcineurin inhibited by the immunophilin-immunosuppressant FKBP12-FK506 complex. Cell 82: 507–522.
12. Li H, Zhang I, Rao A, Harrison SC, Hogan PG (2007) Structure of calcineurin in complex with PVIVIT peptide: portrait of a low-affinity signalling interaction. Nature 449: 654–661.
13. Blumethal DK, Takai K, Hansen RS, Krebs EG (1986) Dephosphorylation of cyclicAMP-dependent protein kinase regulatory subunit (type II) by calmodulin-dependent protein phosphatase. Determinants of substrate specificity. J Biol Chem 261: 8100–8115.
14. Hua Q, Kim HY, Liu Y, Zhao Y, Mondragon A, et al. (2002) Crystal structure of calcineurin/cyclophilin-cyclosporin shows common but distinct recognition of immunophilin-drug complexes. Proc Natl Acad Sci U S A 99: 12037–12042.
15. Jin L, Harrison SC (2002) Crystal structure of human calcineurin complexed with cyclosporin A and human cyclophilin. Proc Natl Acad Sci U S A 99: 13522–13527.
16. Liu J, Farmer JD, Jr., Lane WS, Friedman J, Weissman I, et al. (1991) Calcineurin is a common target of cyclophilin-cyclosporin A and FKBP-FK506 complexes. Cell 66: 807–817.

**Table S1** Data collection and refinement statistics. (DOCX)

**Text S1** Supplementary materials and methods. (DOCX)

**Acknowledgments**

The authors thank Dr. M. Allaire (National Synchrotron Light Source, NSLS) for his support at NSLS beamline X25. Use of the NSLS at Brookhaven National Laboratory was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under contract no. DE-AC02-98CH10886. MD calculations were performed on the Biowulf computing cluster at NIH. The authors thank Dr. Linda Dixon for providing information and materials relating to A238L and Dan Herschlag, Jagorce Roy, and Evan Guiney for helpful discussion and reading of the manuscript. We thank Dara Dowlatshahi for technical advice and Dr. V. Kaila (National Institutes of Health) for his support with the quantum mechanical calculations of the model substrate.

**Author Contributions**

The author(s) have made the following declarations about their contributions: Conceived and designed the experiments: SG RB PC GH RP MSC WP. Performed the experiments: SG RB NL JC PC RP WP. Analyzed the data: SG RB PC GH RP MSC WP. Wrote the paper: SG RB PC GH RP MSC WP.

---

For the full text, please refer to the original publication.
34. Kafadar KA, Cyert MS (2004) Integration of stress responses: modulation of calcineurin signaling in Saccharomyces cerevisiae by protein kinase A. Eukaryot Cell 3: 1147–1153.

35. Kafadar KA, Zhu H, Snyder M, Cyert MS (2003) Negative regulation of calcineurin signaling by Hir25p, a yeast homolog of casein kinase I. Genes Dev 17: 2698–2708.

36. Okamuro H, Aramburu J, Garcia-Rodriguez C, Viola JP, Raghavan A, et al. (2000) Concerted dephosphorylation of the transcription factor NFAT1 induces a conformational switch that regulates transcriptional activity. Mol Cell 6: 539–550.

37. Sopko R, Huang D, Preston N, Chua G, Papp B, et al. (2006) Mapping pathways and phenotypes by systematic gene overexpression. Mol Cell 21: 319–330.

38. Otwinowski Z, Minor W (1997) Processing of X-ray diffraction data collected in oscillation mode. Methods in Enzym (part A) 276: 307–326.

39. Adams PD, Afonine PV, Bankoczi G, Chen VB, Davis IW, et al. (2010) PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr D Biol Crystallogr 66: 213–221.

40. Emsley P, Cowtan K (2004) Coot: model-building tools for molecular graphics. Acta Crystallogr D Biol Crystallogr 60: 2126–2132.

41. Lovell SC, Davis IW, Arendall WB, 3rd, de Bakker PI, Word JM, et al. (2003) Structure validation by Calpha geometry: phi,psi and Cbeta deviation. Proteins 50: 437–450.

42. Becke AD (1993) Density-functional thermochemistry. III. The role of exact exchange. J Chem Phys 98: 5649–5652.

43. Ahlrichs R, Bar M, Haser M, Horn H, Kolmel C (1989) Electronic-structure calculations on workstation computers: the program system TURBOMOLE. Chem Phys Lett 162: 165–169.

44. Rohl CA, Strauss CE, Chivian D, Baker D (2004) Modeling structurally variable regions in homologous proteins with rosetta. Proteins 55: 636–677.

45. Jorgensen WL, Chandrashekar J, Madura JD, Impey RW, Klein ML (1983) Comparison of simple potential functions for simulation liquid water. J Chem Phys 79: 996–995.

46. Hess B, Kutzner C, Spoel D, Lindahl E (2008) GROMACS 4: algorithms for highly efficient, load-balanced, and scalable molecular simulation. J Chem Theory Comput 4: 435–447.

47. Lindorf-Larsen K, Piana S, Palmo K, Maragakis P, Klepeis JL, et al. (2010) Improved side-chain torsion potentials for the Amber ff99SB protein force field. Proteins 78: 1959–1958.

48. Darden T, York D, Pedersen L.(1993) Particle mesh Ewald: an N-log(N) method for Ewald sums in large systems. J Chem Phys 98: 10089–10092.

49. Hoover WG (1985) Canonical dynamics: equilibrium phase-space distributions. Phys Rev A 31: 1695–1697.

50. Andersen HC (1980) Molecular-Dynamics simulations at constant pressure and/or temperature. J Chem Phys 72: 2384–2393.

51. Parrinello M, Rahman A (1981) Polymorphic transitions in single crystals: a new molecular dynamics method. J Appl Phys 52: 7182–7190.

52. Benomar M, Branduardi D, Busi G, Camilloni C, Peviani D, et al. (2009) PLUMED: a portable plugin for free-energy calculations with molecular dynamics. Comput Phys Commun 180: 1961–1972.

53. Marinelli F, Pietrucci F, Laio A, Piana S (2009) A kinetic model of trp-cage folding from multiple biased molecular dynamics simulations. PLoS Comput Biol 5: e1000452. doi:10.1371/journal.pcbi.1000452

54. Akaike H (1974) A new look at the statistical model identification. IEEE Transactions on Automatic Control 19: 716–723.

55. Tait SW, Reid EB, Greaves DR, Wileman TE, Powell PP (2000) Mechanism of inactivation of NF-kappa B by a viral homologue of I kappa b alpha. Signal-induced release of I kappa b alpha results in binding of the viral homologue to NF-kappa B. J Biol Chem 275: 34656–34664.