Atomistic simulations and network-based modeling of the Hsp90-Cdc37 chaperone binding with Cdk4 client protein: A mechanism of chaperoning kinase clients by exploiting weak spots of intrinsically dynamic kinase domains

Josh Czemeres¹, Kurt Buse¹, Gennady M. Verkhivker¹,²*

¹ Department of Computational and Data Sciences, Schmid College of Science and Technology, Chapman University, Orange, California, United States of America, ² Department of Biomedical and Pharmaceutical Sciences, Chapman University School of Pharmacy, Irvine, California, United States of America

*verkhivk@chapman.edu

Abstract

A fundamental role of the Hsp90 and Cdc37 chaperones in mediating conformational development and activation of diverse protein kinase clients is essential in signal transduction. There has been increasing evidence that the Hsp90-Cdc37 system executes its chaperoning duties by recognizing conformational instability of kinase clients and modulating their folding landscapes. The recent cryo-electron microscopy structure of the Hsp90-Cdc37-Cdk4 kinase complex has provided a framework for dissecting regulatory principles underlying differentiation and recruitment of protein kinase clients to the chaperone machinery. In this work, we have combined atomistic simulations with protein stability and network-based rigidity decomposition analyses to characterize dynamic factors underlying allosteric mechanism of the chaperone-kinase cycle and identify regulatory hotspots that control client recognition. Through comprehensive characterization of conformational dynamics and systematic identification of stabilization centers in the unbound and client-bound Hsp90 forms, we have simulated key stages of the allosteric mechanism, in which Hsp90 binding can induce instability and partial unfolding of Cdk4 client. Conformational landscapes of the Hsp90 and Cdk4 structures suggested that client binding can trigger coordinated dynamic changes and induce global rigidification of the Hsp90 inter-domain regions that is coupled with a concomitant increase in conformational flexibility of the kinase client. This process is allosteric in nature and can involve reciprocal dynamic exchanges that exert global effect on stability of the Hsp90 dimer, while promoting client instability. The network-based rigidity analysis and emulation of thermal unfolding of the Cdk4-cyclin D complex and Hsp90-Cdc37-Cdk4 complex revealed weak spots of kinase instability that are present in the native Cdk4 structure and are targeted by the chaperone during client recruitment. Our findings suggested that this mechanism may be exploited by the Hsp90-Cdc37 chaperone to recruit and protect intrinsically dynamic kinase clients from degradation. The results of this
investigation are discussed and interpreted in the context of diverse experimental data, offering new insights into mechanisms of chaperone regulation and binding.

Introduction

The 90 kDa heat-shock proteins Hsp90s belong to a class of highly abundant and evolutionary conserved molecular chaperones that are present in the cytosol of bacteria and in all eukaryotic organisms [1–6]. These chaperones manage late stages of conformational development, maturation and folding for a wide array of client proteins, including protein kinases and transcription factors that constitute a variety of signal transduction pathways and implicated in cell proliferation, differentiation, apoptosis and immune response [7–18]. The Hsp90 chaperone operates as a homodimer with a modular architecture in which each monomer consists of three distinct domains: an N-terminal domain (Hsp90-NTD) that harbors the ATP binding site and a mobile lid motif that mediates ATP-dependent dimerization of the NTDs, a middle domain (Hsp90-MD), which is implicated in binding of client proteins, and a C-terminal domain (Hsp90-CTD) that is required for constitutive dimerization [12–18]. Structural and biophysical studies have identified different conformational states of Hsp90 [19–25] that are associated with the main stages of the Hsp90-ATPase cycle. Hydrogen/deuterium exchange mass spectrometry (HX-MS) studies, electron microscopy (EM) and small-angle X-ray scattering (SAXS) experiments have characterized the thermodynamics and kinetics of structural transitions during the ATPase cycle, showing that these changes reflect stochastic fluctuations between preexisting conformational states, and are only weakly coupled to the nucleotide binding for the eukaryotic cytosolic Hsp90 [26–32]. In contrast to spontaneous conformational changes featured by eukaryotic Hsp90s, the ratchet mechanism of the ‘cochaperone-independent’ bacterial HtpG is strictly controlled by the nucleotide binding [31]. HX-MS studies of conformational dynamics for the HtpG, human Hsp90β, and yeast Hsp90 chaperones have unveiled that eukaryotic Hsp90s in their apo forms are considerably more flexible than the HtpG chaperones [32]. Functional assays of yeast Hsp90-NTD and Hsp90-MD mutants with reported effects on the ATPase activity were undertaken to show that the proper timing of conformational transitions is vital for client recruitment and processing by the chaperone [33]. According to this study, a client-specific residence time in the open conformational states is required for the normal cycle progression, and the excessive stabilization and prolonged occupation of the closed states could be detrimental for the client cycle. By incorporating specific fluorescence probes into yeast Hsp90 and employing single-molecule spectroscopy, kinetics of conformational changes in this chaperone has been recently dissected, revealing cooperativity of local and global motions during the ATPase cycle [34].

The integration of various cochaperones into the Hsp90 machinery can modulate progression of the Hsp90-ATPase cycle and ensure timely execution of protein client loading, activation, and release stages [35–42]. Importantly, cytosolic Hsp90 is the only form of the Hsp90 chaperones that are cochaperone-dependent for functional activities. The family of Hsp90 chaperones can be also regulated by post-translational modifications, which modulate dynamics and interactions with clients and cochaperones [43–49]. Cdc37 is a highly specialized cochaperone that in coordination with Hsp90 can facilitate conformational maturation, protein folding and acquisition of functional states for a large and diverse clientele of protein kinases [50–54]. The initial functional studies of protein kinase motifs involved in binding with the Hsp90-Cdc37 chaperone have revealed that multiple segments from the N-terminal
and C-terminal kinase lobes cooperate in the client recruitment process [55–59]. A high-throughput study of the Hsp90-client interactions using luminescence-based mammalian interactome mapping has provided a first quantitative analysis of the Hsp90-client interactions, revealing that Cdc37 can distinguish kinase clients by recognizing their conformational instability, whereas cooperative Hsp90-Cdc37 binding must be invoked for thermodynamic sorting of strong and weak kinase clients [60]. HX-MS studies and functional assays of structurally similar client and nonclient kinases have confirmed this mechanism, demonstrating that kinase dependence on the Hsp90-Cdc37 chaperone is correlated with the degree of unfolding cooperativity, tryptophan accessibility and client compactness that favor exposure of the regulatory regions in the catalytic domain [61].

Using mutational analysis, this study showed that the interaction strength between Hsp90-Cdc37 chaperone and Src kinases can span a continuum spectrum of binding affinities from the weak client c-Src to the constitutive strong client v-Src kinase. These findings have suggested that similar molecular principles may underlie chaperone-kinase interactions for both weak and strong clients, and the propensity to expose proper interaction surfaces during recruitment to the chaperone may ultimately determine the interaction strength between a kinase and the Hsp90-Cdc37 system [61]. Recent studies have reaffirmed a central role of Cdc37 in chaperoning of kinase clients as silencing of Cdc37 function in cancer cells dramatically reduces binding of oncogenic kinase clients with Hsp90 [62]. By investigating the effects of Cdc37 mutants that disrupt the Hsp90-Cdc37 interactions on client binding, it was recognized that restricted Hsp90-Cdc37 interactions would not present an impediment for effective chaperoning of a wide range of kinase clients [63]. NMR spectroscopy mapping and fluorescence assays of the Hsp90-Cdc37 interactions with protein kinase clients have shown that these complexes could form cooperatively through an allosteric cross-talk between multiple interaction sites [64]. Molecular mechanisms underlying client recognition and thermodynamic sorting by the Hsp90-Cdc37 chaperone have been further explored by NMR approaches [65]. These illuminating experiments have established that Cdc37 can recognize kinase clients through a series of successive stages, in which the N-terminal domain of Cdc37 (Cdc37-NTD) may perform primary screening of kinase clients and induce partial disorder and the enhanced heterogeneity of kinase ensembles, while the C-terminal domain regions (Cdc37-M/C) would provide additional protection of partially unfolded client states. Accordingly, Cdc37 may expand its 'duties' as a chaperone adaptor and serve a primary 'sensor' of kinase instability that triggers a cascade of allosteric changes that promotes local unfolding and effective sorting of client substrates with the aid of Hsp90 [65].

Structural and thermodynamic characterization of the Hsp90-Cdc37 binding with protein kinase clients have been constrained by difficulties associated with the dynamic changes in client and chaperone conformations and a transient character of chaperone-client interactions. Multiple flexible regions in the Hsp90 and Cdc37 chaperones were implicated as potential binding sites for client interactions, suggesting that the intermolecular binding interfaces undergo structural and dynamic remodeling during client binding and release cycle [66, 67]. The initial EM reconstruction of the Hsp90-Cdc37-Cdk4 kinase complex unveiled an asymmetric assembly and a broad network of interactions formed by the Hsp90 dimer with Cdc37 cochaperone and Cdk4 [68]. The EM-based analysis of the Hsp90-Cdc37-Cdk4 complex offered insight into the stoichiometry, topology and shape of this multi-protein assembly, but the atomistic details of the binding interactions remained hidden. The cryo-EM structure of the human Hsp90-Cdc37-Cdk4 kinase complex [69] marked a fundamental breakthrough in the field, revealing a complex and highly intertwined arrangement in which the Hsp90 dimer and Cdc37 trap a partially disordered kinase state (Fig 1). In this structure, Hsp90 dimer assumes a symmetrical closed conformation, which closely resembles an ATP-bound closed
state of yeast Hsp90 [19]. A partially unfolded N-lobe is implanted inside the Hsp90 dimer and stabilized by the interactions with Hsp90-MD and Hsp90-CTD (Fig 1). A number of unique features and several major intermolecular interfaces that characterize the underlying topology of the Hsp90-Cdc37-Cdk4 complex indicated that allosteric interactions between binding sites are delicately tuned to enable client recognition and progression of the chaperone cycle. Based on structural characterization of the Hsp90-client interactions, Agard and colleagues have proposed a model of the chaperone cycle [69] according to which Cdc37 initially sequesters an open kinase conformation and the resulting intermediate binds to a client-loading open form of the Hsp90 dimer. During the functional cycle, the Hsp90 chaperone progresses from a
relaxed form to a closed ATP-bound state that effectively traps a partially unfolded kinase client, leading to the formation of the Hsp90-Cdc37-Cdk4 complex. Upon ATP hydrolysis, Hsp90 opens up and releases the folded kinase client from the chaperone system. Based on their pioneering work, Agard and colleagues have also inferred that dynamic Hsp90-client interactions may have evolved to enable rapid and diverse kinase responses in signaling networks, where the chaperone role is to protect unstable kinases from degradation and suppress an undesirable basal activity [70]. Importantly, this study has indicated that even mature kinases can maintain chaperone dependence, as Hsp90-addicted kinase clients could frequently sample a partially unfolded state even after folding. Collectively, the body of structural and functional investigations has provided evidence that Cdc37 can serve as a recognition scaffold for unstable kinase clients, but requires cooperation with Hsp90 to properly execute and complete cycle of client maturation, folding and release. However, the molecular details of this highly dynamic binding mechanism that operates through allosteric coordination of several spatially separated binding interfaces remain largely elusive. In addition, the structure of the chaperone-kinase complex provided only a single snapshot of a highly dynamic assembly formed through cooperative allosteric interactions. Current understanding of allosteric communication and long-range signaling that underlie the Hsp90-Cdc37 interactions with client kinases remains rather simplistic and incomplete to explain diverse experimental data on mutations affecting allosteric regulation and client binding.

Computational studies have also elucidated molecular details of Hsp90 dynamics and binding with cochaperones and inhibitors to quantify mechanisms of allosteric regulation in Hsp90. Our initial studies have provided evidence for allosteric communications in Hsp90 by showing that coordinated chaperone movements enable efficient signal transmission between the NTD and CTD binding sites [79–81]. Using these early insights, we combined computational and experimental approaches in structural characterization of the C-terminal binding site of Hsp90 and discovery of allosteric inhibitors [82–84].Atomistic molecular dynamics (MD) simulations and coarse-grained approaches have systematically examined conformational dynamics and collective motions in different Hsp90 states [85, 86]. By combining MD simulations and structure-based network analysis, we have characterized conformational dynamics and allosteric interactions in the Hsp90 complexes with various cochaperones and client proteins [87–90]. These studies have shown that the residue interaction networks in the Hsp90 complexes with p53 [89] and client recruiter cochaperones Cdc37, Sgt1 and Rar1 [90] form small-world topologies in which key mediating centers of the global communications determine functional hotspots of chaperone regulation. The network-centric modeling of the Hsp90 interactions has suggested that allosteric coupling between key functional centers is required in regulation of the inter-domain communications, control of ATP hydrolysis, and protein client binding [88–90]. Computational design and functional assay experiments of mutations that control allosteric points of Hsp90 have characterized structural and dynamic signatures of allosteric Hsp90 switches [91]. A cross-talk between allosteric ligand binding and dynamic changes in the Hsp90 has been explored to characterize mechanisms of allosteric activation in the chaperone [92]. Collectively, computational studies indicated that the interactions of Hsp90 with cochaperones and client proteins are highly dynamic and can elicit a cascade of conformational changes through modulation of structural stability and flexibility in the interacting proteins.

In the current work, we performed all-atom molecular dynamics (MD) simulations of the Hsp90 and Cdk4 structures in different functional states. Multiple independent MD simulations were carried out for the Cdk4-cyclin D1/D3 complexes, a monomeric unbound form of Cdk4, a client-free Hsp90 dimer and the Hsp90-Cdc37-Cdk4 complex. Atomistic simulations and distance fluctuation analysis of MD trajectories were combined with the energy analysis of
binding interfaces to examine stability of different functional states and identify regulatory hotspots that may control client recognition. We present evidence supporting an allosteric mechanism in which client binding can trigger coordinated dynamic changes and induce global rigidification of the Hsp90 inter-domain regions that is coupled with a concomitant increase in conformational flexibility of the kinase client. By combining conformational dynamics with the rigidity decomposition analysis and network-based emulation of thermal unfolding, our study suggested that Hsp90-Cdc37 chaperone may exploit differential stabilization of the kinase lobes and target weak spots of highly dynamic Cdk4 client to induce and protect a non-native, partially unfolded kinase state. The results of this investigation are discussed and interpreted in the context of diverse experimental data, offering new insights into mechanisms of chaperone regulation and binding.

Results and discussion

Atomistic simulations of the Hsp90 chaperone in different functional forms: Client-induced Modulation of conformational ensembles couples local and global dynamic changes

Using an ensemble-based model of allosteric regulation [93–97], we examined several stages of the chaperone-client cycle by simulating conformational dynamics of the Hsp90 chaperone and Cdk4 structures in their different unbound and client-bound functional states. The central hypothesis of this analysis assumes a mechanism of client recognition by the chaperone system in which modulation of structural and dynamic changes in the interacting proteins controls progression of the chaperone and kinase life cycles. The cryo-EM structure of the human Hsp90-Cdc37-Cdk4 kinase complex [69] showed a complex and intertwined arrangement of the interacting modules (Fig 1). In this structure, Hsp90 dimer assumes a closed conformation and the open structure of the kinase is in a partially unfolded state, with its two lobes on separate sides of Hsp90 and with the intervening region in an elongated nonnative conformation. The partially unfolded N-lobe region of the kinase client is threaded through the cleft between the Hsp90 monomers and stabilized by hydrophobic contacts with the client-binding site that mimic the native interactions between the kinase lobes. The crystal structures of phosphorylated Cdk4 bound to cyclin D revealed a dynamic Cdk/Src-like inactive kinase conformation in these complexes, showing that cyclin binding and phosphorylation are still not sufficient to trigger activation conformational transitions. Importantly, the activation mechanism of Cdk4 client is distinct from nonclient members of Cdk family [60] and requires synchronization of multiple events including cyclin binding, phosphorylation and substrate binding to achieve full activation [98,99]. It remains unclear why chaperone-induced trapping of an open Cdk4 conformation can benefit its activation. As a possible explanation, it was suggested that a lobe-separated, open Cdk4 conformation could create favorable thermodynamic and kinetic conditions for efficient ATP binding, which in turn would stabilize the kinase active state and trigger subsequent release of the folded kinase from chaperone system [100–102]. It was also argued that the increased dynamics of the open kinase state can be beneficial for rapid access and binding of substrates that are essential for Cdk4 activation [102].

To address these questions, we simulated conformational dynamics of the Hsp90 chaperone and Cdk4 structures in their native and client-bound functional states. We tested a hypothesis that Cdk4 catalytic domain may be predisposed for recruitment to the chaperone system by featuring an intrinsically dynamic structure with the excessively flexible N-lobe regions—a common attribute of kinase clients that may be exploited by the chaperone to induce partial unfolding of kinase client upon recruitment to the Hsp90-Cdc37-Cdk4 complex. First, we compared conformational dynamics of the unbound Hsp90 dimer and client-bound Hsp90 in
the Hsp90−Cdc37−Cdk4 complex. In this analysis, we examined whether binding could alter conformational dynamics of the Hsp90 dimer and cause local or global redistribution of conformational mobility in functional regions of the binding partners. Conformational mobility of protein residues was evaluated by using the mean square residue fluctuations and computed B-factors. We specifically focused on changes in the conformational variations of the Hsp90 regulatory motifs, the inter-domain and intermolecular interfaces (Fig 2). The increased stabilization of the Hsp90 dimer in the chaperone-client complex propagated from the interfacial regions to the Hsp90-MD and Hsp90-CTD regions (Fig 2A and 2B). The Hsp90-MD residues in the core of the complex and the three-helix bundle (residues 398−453) exhibited the reduced thermal fluctuations, as these regions were stabilized by the intra- and intermolecular interactions. Consistent with the HX-MS experiments [32], conformational dynamics showed the

https://doi.org/10.1371/journal.pone.0190267.g002

Fig 2. Conformational dynamics and PCA modes in the unbound and client-bound Hsp90 dimers. Conformational dynamics profiles of the Hsp90-monomer (A) and Hsp90-B monomer (B) in the unbound forms (shown in brown filled bars) and in the complex (shown in colored lines). Residue-based conformational mobility profiles in the complex are shown as lines with filled squares and colored as follows: Hsp90-NTD (residues 1−215) in blue, Hsp90-MD (residues 216−552) in red, and Hsp90-CTD (residues 553−690) in green. (C, D) The normalized squared displacement of the Hsp90 residues averaged over first three PCA components are shown for the unbound (in brown lines) and bound Hsp90 dimer (in red lines). The slow mode profile for Hsp90- monomer a (panel C) and for Hsp90-monomer B (panel D).
increased stability of the Hsp90-MD clusters 420–429, 472–484 and the MD-CTD interfacial region (residues 513–519) (Fig 2A and 2B). The intermolecular binding interfaces displayed reduced fluctuations, including the Hsp90-MD segment (residues K399, K402, R405, and K406) involved in interactions with Cdc37-NTD and Hsp90 residues F341, L343, F344, L611 and M602 that interact with the disordered kinase region (Fig 2A and 2B). Of special significance were the markedly reduced conformational fluctuations and stabilization of the recognition Src loop (residues 341-FDLFENKKKK-350), and the Hsp90-CTD helix 15 (residues 599–611) that displayed highly mobility in the unbound Hsp90 dimer [32]. The hydrophobic motif of the amphipathic Src-loop maintained stabilizing contacts with the Cdc37-M/C domain and kinase N-lobe during simulations and remained largely immobilized in the complex. A noticeable redistribution of conformational dynamics was also observed in the Hsp90-CTD helix 15, where the extensive intra- and intermolecular contacts formed by M602, L611, and R612 residues promoted stabilization of the Hsp90-CTD regions (Fig 2A and 2B). Our findings revealed that conformational dynamics of the Hsp90 near the Hsp90-CTD dimerization regions and binding interfaces with Cdk4 was considerably suppressed. Importantly, however, client binding induced structural stabilization that propagated beyond the binding interface residues. The integration of the Cdk4 client to the Hsp90-Cdc37 system incurred global changes in the dynamics of the Hsp90 dimer by narrowing the accessible conformational space and reducing flexibility of the MD and CTD regions in both monomers. The quenching of local dynamics in the MD and CTD domains was balanced by the increased flexibility of the Hsp90-NTD regions and strengthening of the inter-domain couplings (Fig 2A and 2B). These coordinated dynamic changes can be functionally significant to facilitate client binding and ensure proper dwell time of the client-bound conformational state during the chaperone cycle. Overall, local quenching of subdomain dynamics in the MD and CTD regions of Hsp90 can be counterbalanced by strengthening of the inter-domain correlations and inter-molecular couplings. A similar pattern of dynamic changes induced by allosteric activators was detected in other protein systems that are primarily regulated by dynamically-driven allostery [103–105].

To characterize functional motions and collective dynamics of the Hsp90 chaperone, we used Principal Component Analysis (PCA) [106,107] of MD trajectories and compared the slow mode profiles averaged over first three PCA components for the unbound and bound forms of Hsp90 (Fig 2C and 2D). The maxima along the slow mode profiles correspond to regions undergoing global structural movements, while the local minima describe immobilized hinge positions that coordinate collective changes. The slow mode shapes for the unbound and client-bound forms of the Hsp90 dimer revealed a considerable similarity, supporting the notion that conservation of major hinge centers and preexisting pathways of functional changes are determined by the native protein topology. In the unbound form, the slow mode profiles revealed several minima in the Hsp90-MD corresponding to hydrophobic residues P287, W289, F341, L343, Y356, F376, L369, I370, I400, and L401 (Fig 2C and 2D). Structural mapping of the Hsp90 hinge clusters pointed to their strategic location near the inter-domain regions (S1 Fig). In particular, the inter-domain coupling in the Hsp90 dimer are determined by the stable hinge cluster formed at the NTD-MD interface by F208 from the NTD and MD residues P287, W289, Y356 and V360. These observations are consistent with the experimentally observed critical role of this hydrophobic cluster to stabilization of the NTD-MD interface in the Hsp90 chaperones [108].

Structural maps of collective dynamics driven by the slowest PCA modes in the unbound (Fig 3A) and client-bound Hsp90 dimers (Fig 3B) highlighted client-induced partial redistribution and migration of the hinge centers in the complex. We observed relocation of major hinge clusters from the NTD and NTD-MD regions in the unbound Hsp90 dimer to the MD-CTD regions in the Hsp90-Cdc37-Cdk4 complex (Fig 3). In the Hsp90-Cdc37-Cdk4
complex, the hinge centers in the Hsp90-MD and MD-CTD inter-domain regions (F341, L343, Y356, I370, and F376) expanded and formed interacting local clusters that further strengthened stability of these coordinating sites. Several hinge residues in the Hsp90-MD were organized in the cluster L369-I370-V381-L401-I404 located at the binding interface with Cdk4-NTD (S1 Fig). In the MD-CTD inter-domain region, hinge residues W598, M602, M606, L611, and R612 were colocalized and formed a hinge center that mediated the intermolecular coupling with the client kinase (S1 Fig). Importantly, the mobility of the recognition Src-loop (residues 341-FDLEF-345) and MD-CTD interfacial regions (residues 598–612) was markedly reduced in the slow modes, giving rise to the formation of a major hinge center at the interface with the client kinase.

Exploring conformational dynamics of the Cdk4 client: Differential stabilization of kinase lobes and highly flexible N-lobe are intrinsic characteristics of dynamic kinase clients

To characterize structural and dynamic changes in the Cdk4 client, we explored conformational landscape of the native Cdk4 domain by simulating crystal structures of Cdk4/cyclin D1
and Cdk4/cyclin D3 complexes. MD simulations of the Cdk4-cyclin D3 crystal structure revealed a considerable conformational flexibility of the inactive Cdk4 conformation that was especially evident in the N-lobe regions, including the αC-helix (residues 50–67), αC-β4 loop (residues 67–74) and β4-β5 sheet (residues 75–95) (Fig 4A). Due to the increased flexibility of the N-lobe regions proximal to the αC-helix/αC-β4-loop, we observed local movements of the αC-helix, often deviating from the fully inactive 'out' position. These results are consistent with the HX-MS studies that suggested that the increased exposure and greater backbone hydrogen exchange near activation regions is a unifying characteristic of kinase clients [61]. In the Cdk4-cyclin D3 complex (pdb id 3G33), the Gly-rich β3-αC loop experienced considerable thermal fluctuations (Fig 4A). The elevated mobility in this region could remove steric constraints on positional preferences of the adjacent αC-helix which explains conformational

Fig 4. Conformational dynamics of Cdk4 in different functional states. (A) Conformational dynamics profile of Cdk4 catalytic domain in the complex with cyclin D3 (pdb id 3G33) is shown in brown bars and serves a reference for comparison with mobility profiles in other functional states. Conformational dynamics profile of the unbound, monomeric Cdk4 form is shown in maroon lines. Cdk4 mobility in the Hsp90-Cdc37-Cdk4 complex is shown for N-lobe in blue lines and C-lobe in red lines. (B) The normalized squared displacement of the Cdk4 domain residues averaged over first three PCA components are shown for the Cdk4-cyclin D3 complex (in brown lines) and for the Hsp90-Cdc37-Cdk4 complex (in red lines). (C) An overlay of representative Cdk4 conformations from the MD ensemble of the partially disordered Cdk4 client in the Hsp90-Cdc37-Cdk4 complex. The ordered portion of the N-lobe is shown in cyan ribbons, a disordered region of the N-lobe is in green ribbons, and the ordered C-lobe is in red ribbons. Notice a significant conformational mobility and inflated nature of the ordered regions, while disordered region is shielded by the Hsp90 interactions.
variations of the αC-helix between the inactive and active-like conformations. MD simulations also showed that the C-lobe core became less compact as the central αE-helix (residues 113–134) and αF-helix (residues 193–210) displayed appreciable thermal fluctuations. Significant dynamic changes were also detected in inter-lobe regions, including the αD-helix (residues 99–107) and a small proline-rich loop (108-PPPGLP-113) connecting the αD-helix and αE-helix. These findings indicated that the intrinsic dynamics of Cdk4 can result in the reduced allosteric coupling between functional regions and lower rigidity of the kinase fold. These factors can determine client status of Cdk4 and create a favorable dynamic environment for chaperone intervention. Hence, Cdk4 can be predisposed for recruitment to the chaperone system by featuring an intrinsically dynamic catalytic domain.

We also analyzed conformational dynamics of Cdk4 in the Hsp90-Cdc37-Cdk4 complex. The client kinase is held by the Hsp90-Cdc37 in a spatially confined, yet highly dynamic ensemble. The results revealed appreciable fluctuations of Cdk4, particularly in partially unfolded N-lobe regions β4 (residues 75–85), β5 (residues 86–93) and αC-β3 loop (46-NGGGGGGL-54) (Fig 4A). A comparison of slow modes profiles for the Cdk4 client revealed conservation of hinge residues located near the inter-lobe region that is anchored by Cdc37-NTD (Fig 4B). Overall, the ensemble of Cdk4 conformations displayed signatures of a swollen globule-like state [109], where the folded C-lobe and ordered N-lobe region of Cdk4 client were mobile showing and inflated structure with packing interactions broken. At the same time, the unfolded portion of the N-lobe is shielded and partially stabilized by the Hsp90 interactions (Fig 4C).

To quantify the extent of protection of the kinase residues in the complex, we also evaluated the residue depth (RD) values by measuring the average residue exposure in the MD ensemble [110–112]. The results showed that the hydrophobic regions near the αC-helix (residues 50–67), αC-β4 loop (residues 67–74) and a β4-β5 sheet (residues 75–95) are indeed protected in the complex (S2 Fig). Hence, integration of Cdk4 client into the Hsp90 system results in spatial confinement of the dynamic Cdk4 conformational ensemble that is shielded by stabilizing contacts of the Hsp90 binding region. This can facilitate the release of water molecules from the interacting proteins and favor dynamic ensemble that is reminiscent of so-called dry molten globule intermediate that can occur at the initial stage of unfolding [109]. A similar compaction of conformational ensembles in client proteins upon recruitment to the chaperone was observed in the NMR studies of chaperone Spy and its soluble client protein Im7 [113], and for clients bound to the periplasmic trimeric chaperone Skp [114,115]. Noteworthy, Hsp90 provides a significantly larger surface area for interactions with substrate proteins than other chaperones [116,117]. It was proposed that client recognition by the Hsp90 can be based on distributing weak hydrophobic interactions over a fairly large surface that is supported by some electrostatic interactions. Intriguingly, our findings that a flexible Src-loop in the Hsp90-MD plays an important role in the dynamic exchange with the client kinase corroborated with several structural studies suggesting that this chaperone region is involved in binding with many partially disordered protein clients. In particular, NMR-based structural analysis of the Hsp90 interactions with the intrinsically disordered Tau protein revealed the binding site near the amphipathic Src-loop of Hsp90, suggesting that similar binding principles can be exploited by the chaperone in recruitment of distinct client proteins [118].

In general, by comparing conformational dynamics of the chaperone and client kinase in different functional states, we found that global propagation of rigidity in the Hsp90 binding regions can be coupled with the instability of the kinase client. A mechanism of reversible entropy transfer where the ordering of the chaperone flexible regions is accompanied by reciprocal unfolding or increased dynamics of the bound substrate was previously recognized as a distinct mode of chaperone function that is particularly well-suited for the chaperone cycles of
repeated client recruitment and release [119,120]. Our results indicated that a similar mechanism may be also operational during client recognition and integration into the Hsp90-Cdc37 chaperone complex.

Distance fluctuations analysis of the conformational ensembles was used to probe structural stability and allosteric communication propensities of the Hsp90 and Cdk4 residues in the unbound and client-bound chaperone forms. We computed the residue-based distance fluctuation force constants that measure the energy cost of the residue deformation during simulations [121–123]. In this approach, structural rigidity of functionally important regions is associated with their high stability indexes and small fluctuations in their average inter-residue distances. Our previous studies showed that distance fluctuations profiles and respective force constant indexes can be related with allosteric residue propensities, as the mean square fluctuations between a pair of residues provides an efficient measure of the signal commute time [87–90]. According to this model, stable residues that display small fluctuations in their distances to other residues correspond to effectively communicating rigid sites and can serve as allosteric hotspots (or ‘signal dispatchers’). We first compared the distance fluctuation force constant profiles in the unbound and bound forms of the Hsp90 dimer (Fig 5A and 5B). This analysis revealed client-induced differential changes in the stability of the Hsp90 monomers. While the stability of the Hsp90-MD residues in the first monomer was moderately decreased (Fig 5A), a significant stabilization of the Hsp90-MD regions was seen in the second monomer B (Fig 5B), reflecting strengthening of both the inter-domain interactions and intermolecular contacts with Cdc37 and Cdk4 binding partners. Notably, functional hinge centers corresponded to the sharp peaks in the force constant distribution. Of particular interest was the increased distance fluctuation force constants and corresponding rigidification of the recognition Src-loop residues 341-FDLF-344 that are involved in the intermolecular contacts with the disordered kinase regions. More importantly, the predicted allosteric centers in the Hsp90-MD (L369, I370, V381, L401, and I404) and Hsp90-CTD regions (W598, M602, M606, L611, R612) formed large stable clusters that rigidified structural environment beyond the local binding interfaces.

A global expansion of stable allosteric clusters induced by client binding was observed in the Hsp90-MD and Hsp90-CTD regions that also solidified the MD-CTD inter-domain couplings (Fig 5A and 5B). In addition, we registered large force constant values for the Cdc37-NTD residues that are involved in binding with the Hsp90-MD and anchor Cdk4 C-lobe (Fig 5C). On the other hand, the force constant values for Cdk4 N-lobe residues were reduced as compared to the native Cdk4 structure (Fig 5D). The observed redistribution of allosteric hotspots supported our findings that major centers that control signal transmission in the complex migrated to the Hsp90-MD/CTD regions and Cdc37-NTD interfacial residues. Moreover, a global rigidification and the enhanced mediating capabilities of these Hsp90 and Cdc37 regions can be contrasted with the increased instability of Cdk4 residues. These results supported a model of reciprocal dynamic changes between Hsp90 and Cdk4 in which client binding can induce expansion of allosteric centers in the Hsp90 MD-CTD regions that stabilize the Hsp90 recognition interface. At the same time, Hsp90-kinase interactions protect a heterogeneous Cdk4 ensemble in a spatially confined inflated state. Distance fluctuation analysis also revealed differential destabilization (or polarization) of the kinase lobes, where a more dynamic N-lobe can be contrasted with more stable C-lobe. This effect was seen in the folded Cdk4-cyclin D complex, monomeric form of Cdk4, and in the Hsp90-Cdc37-Cdk4 complex (Fig 5D). These results are consistent with our recent studies in which we showed that divergences in the activation mechanisms of Cdk2 (nonclient) and Cdk4 (strong client) may be associated with differences in the relative stability and cooperativity of the kinase lobes in these Cdk proteins [124]. Interestingly, client binding can exploit differential stability of the kinase
lobes by further increasing the disparity between highly dynamic N-lobe (tolerant to perturbations) and relatively stable C-lobe (more susceptible to perturbations). Conformational dynamics showed that the most flexible regions in the native Cdk4 structures correspond to the αC-β3 loop (46-NGGGGGL-54), β4 (residues 75–85), and β5 strands (residues 86–93) of the N-lobe. Strikingly, these regions are targeted and become partially unfolded upon recruitment to the Hsp90-Cdc37 chaperone. Our results suggested that the observed effect may reflect the intrinsic dynamic preferences of kinase clients, in which these N-lobe regions may be especially vulnerable to perturbations and form weak spots of kinase instability. These findings are consistent with similar effects seen in oncogenic kinases that are typically chaperone clients, where activation mutations are mainly assembled in a more flexible N-lobe that allows to readily modulating activation transitions [125,126]. Structural separation of stable

https://doi.org/10.1371/journal.pone.0190267.g005

Fig 5. The distance fluctuations analysis and residue-based force constant profiles of the unbound and client-bound Hsp90 forms. The distance fluctuation force constant profiles of the Hsp90-monomer (A), Hsp90-B monomer (B) and Cdk4 kinase client (C) in the unbound forms (shown in grey light bars) and in the complex (shown in domain-colored bars). Residue-based force constant profiles in the complex are shown in bars and colored as follows: Hsp90-NTD (residues 1–215) in blue, Hsp90-MD (residues 216–552) in red, and Hsp90-CTD (residues 553–690) in maroon. (C) The Cdc37-NTD (residues 1–147) is shown in blue, Cdc37-M/C (residues 148–260) in red. (D) Cdk4-N lobe (residues 1–99) is shown in blue, and Cdk4 C-lobe (residues 100–295) is in red.
and flexible regions in the chaperone-kinase complex highlight the notion that protein polarity can be required to afford evolution of new functions without sacrificing fold stability [127–129].

Energetic analysis of protein-protein binding interfaces in the Hsp90-Cdc37-Cdk4 complex differentiates hotspot residues and quantifies functional role of the Cdc37-M/C domain

We also performed alanine scanning of the binding interfaces in the Hsp90-Cdc37-Cdk4 complex by computing changes in protein-protein binding free energies induced by mutations of the interfacial residues (Figs 6 and 7). The central objective of this analysis was to determine energetic hotspots of protein binding and stability in the complex and quantify role of these functional residues in the allosteric mechanism of the chaperone-kinase cycle. In this analysis, we employed and compared three different computational methodologies that included

![Fig 6. Alanine scanning and free energy analysis of the binding interfaces in the Hsp90-Cdc37-Cdk4 complex using BeAtMuSiC approach.](https://doi.org/10.1371/journal.pone.0190267.g006)
coarse-grained, knowledge-based BeAtMuSiC predictor of binding free energy changes \([130–132]\), empirical force field approach FoldX with the all-atom representation of protein structure \([133–137]\), and the molecular mechanics Poisson–Boltzmann surface area (MM-PBSA) method \([138, 139]\), in which all-atom MD simulations were utilized to determine the ensemble-averaged changes in the binding affinity of the interacting proteins \([140–142]\). The knowledge-based BeAtMuSiC predictor utilizes a combination of 13 statistical potentials weighed by solvent accessibility and evaluates the effect of single missense mutations on the strength of interfacial interactions and the folding free energy of the complex \([130]\). We opted for this approach as a primary tool in our computations of mutation-induced binding free energy.
changes based on its known efficiency and robustness in reproducing experimental data. Indeed, BeAtMuSiC was amongst the top performing tools during community-wide evaluation of methods for predicting the effect of mutations on protein-protein interactions [131], where it produced a robust correlation with the experimental data from the SKEMPI database [132].

First, we present a detailed energetic analysis of the binding interfaces in the Hsp90-Cdc37-Cdk4 complex obtained from BeAtMuSiC predictions (Fig 6). The free energy profiles revealed a group of hotspot residues in the Hsp90-MD and Hsp90-CTD regions that contribute decisively to binding with Cdc37 (Fig 6A) and Cdk4 protein partners (Fig 6B). The Hsp90 binding interfaces with Cdc37 and Cdk4 are dominated by strong interactions of the Hsp90-MD hotspot residues (F341, L343, F344, K399, I400, L401, K402, K406) and Hsp90-CTD hotspots (M602, L611, R612) (Fig 6A and 6B). The hotspot residues often anchor local interaction clusters that are interconnected and stabilize binding interfaces in the complex (Fig 7A and 7C). One of these hydrophobic clusters (I404-V381-I408-L369-1370-I400-I401), which is centered around I400 and L401 hotspots, is linked with a group of interfacial lysine residues (K399, K402, K406, and K411) that form salt bridge interactions with Cdc37-NTD residues (Fig 7A and 7B). This cluster closely overlaps with an allosteric center in the Hsp90-MD (L369, I370, V381, L401, and I404). Another energetic hotspot is formed by cluster F376-L343-F433-N436 that anchored the Src-loop positions (R338, F341, L343, and F344) with the Cdk4 N-lobe (Fig 7C). Of particular significance was emergence of the Hsp90-CTD residues (M602, L611, R612) as prominent energetic hotspots of the Hsp90-Cdk4 binding interface. These observations corroborated with the dynamics analysis, showing that conformational flexibility of the Hsp90-CTD binding interface with Cdk4 was largely restricted due to favorable interactions made by these residues. A key energetic hotspot of the Cdc37 binding interface with Cdk4 corresponded to Cdc37-NTD motif 20-HPN-22 that mimics the shape and the interaction pattern of the αC-β4 loop (Cdk4 N-lobe) with the C-lobe (Fig 7D). According to our analysis, structural and energetic mimicry provided by the HPN motif may favor displacement of the αC-β4 loop and Cdc37-induced stabilization of the lobe-separated, open kinase state. A strong energetic contribution to binding affinity was detected for several residues in the Cdc37-M/C domain (R166, R167 and Q208) (Fig 6C and 6D). Structural mapping of the Cdc37-M/C interfaces highlighted the decisive role of the Cdc37 hotspots (R245, Q246, R166, and R167) in binding with Hsp90 and Cdk4 respectively (Fig 7E). Interestingly, the predicted Cdc37-M/C hotspots R166, R167 and Q208 were previously reported as important interfacial sites in the NMR study of the complex between human Cdc37 (residues 148–276) and human Hsp90-NTD [143,144]. A biochemical analysis of the full-length human Hsp90-Cdc37 complex in living cells showed that mutations in the Cdc37 (M164A, R167A, L205A, and Q208A) could reduce the Hsp90-Cdc37 interactions by as much as 70–95%. [145]. In the context of our analysis, these residues may correspond to conserved hotspots that can be exploited by Cdc37 in binding with different partners and in different structural contexts. In the Hsp90-Cdk4 interface, several Cdk4 residues (K88, E94) from a partially disordered segment anchored strong electrostatic interactions with E345 (Hsp90-MD) and R612 (Hsp90-CTD) that can impose spatial constraints on the dynamic ensemble of open Cdk4 conformations (Fig 6E and 6F). We argue that spatial compaction of the heterogeneous client ensemble can speed up conformational search within the funnel of globule-like states and thereby facilitate process of client refolding and release from the chaperone system.

Our results supported the notion that while Cdc37-NTD may control initial recognition and screening of client folds, a subsequent thermodynamic separation of strong and weak kinase clients may be determined through modulation of binding affinity by the Cdc37-M/C
The energetic analysis of the Hsp90-Cdc37 interactions with the kinase client also supported a model in which binding may promote flexibility of the client substrate while simultaneously inducing stabilization of the flexible recognition regions in Hsp90. This mechanism may underlie allosteric cooperativity between Cdc37 domains and Hsp90 during recognition and recruitment of kinase clients. To further validate energetic predictions, we compared the BeAtMuSiC results (Figs 6 and 7) with the FoldX method (Fig 8A) and MM-PBSA approach (Fig 8B). The results displayed a strong positive correlation between binding free energy changes computed by different tools, also pointing to similar group of residues as energetic hotspots. Only a slightly lower correlation coefficient (R = 0.608) was found when BeAtMuSiC predictions were plotted against MM-PBSA results (Fig 8B). A somewhat greater dispersion of binding free energy changes in this comparison was likely due to the ensemble-average estimates in MM-PBSA calculations that were based on MD simulation trajectory of the complex. In general, a comparison of three different energetic methods produced consistent results, highlighting efficiency of knowledge-based BeAtMuSiC predictor in the analysis of binding free energies of protein-protein complexes. A detailed quantitative analysis of the binding interfaces in the Hsp90-Cdc37-Cdk4 complex obtained from MM-PBSA calculations (S3 Fig) demonstrated a close correspondence with BeAtMuSiC predictions. Noteworthy, these methods produced similar distribution peaks and predicted the same binding energy hotspots, suggesting that these salient signatures of the binding interfaces may be fairly robust and largely independent on the details of the energy force field. In particular, MM-PBSA calculations reaffirmed an important role of the Hsp90-CTD residues (M602, L611, R612) as key energetic hotspots of the Hsp90-Cdk4 binding interface (S3 Fig). Additionally, the MM-PBSA profiles highlighted a significant contribution of the Cdc37-M/C domain (R166, R167) in binding with Cdk4 client and strong energetic effect of the Hsp90 interactions.
with the disordered kinase region. In general, the energetic analysis quantified contribution of different binding interfaces and reinforced the notion that client recognition by the Hsp90-Cdc37 chaperone system may involve a significant dynamic and energetic exchange between binding partners near the extended hydrophobic interfaces.

A model of chaperoning kinase clients by targeting weak spots of intrinsically dynamic kinase domains: Insights from network-based rigidity analysis and thermal unfolding simulations

Using FIRST approach [146–150] and the Python-based Constraint Network Analysis (CNA) interface [151,152] we also performed network-based rigidity decomposition of the Hsp90 and Cdk4 structures in different functional states. This analysis particularly quantified changes in the rigidity and flexibility of Cdk4 kinase domain in the unbound form, Cdk4-cyclin D complex and Hsp90-Cdc37-Cdk4 complex. The underlying hypothesis behind this approach is that a) differential stabilization of the N-lobe and C-lobe in the client kinase may be intrinsically present in the Cdk4-cyclin D complex and b) Hsp90-Cdc37 chaperone may exploit intrinsic flexibility of the N-lobe and presence or weak spots to amplify instability and induce a partially unfolded kinase state that is protected by the chaperone. In the FIRST approach, thermal unfolding of protein structures is emulated by gradually removing noncovalent constraints from the constrained network of bonds connecting residue nodes in the protein structure. This procedure can determine whether a bond is flexible or rigid and decompose the constraint network into rigid clusters and flexible regions. During unfolding, the weak constraints are removed first while stronger interactions are sustained longer, leading to progressive decomposition into rigid and flexible regions. We monitored the evolution of the ‘giant’ rigid cluster that disintegrates and breaks apart into a number of smaller rigid clusters during unfolding phase transition. Rigidity decomposition algorithm identifies unfolding nuclei as residues that break away from the giant rigid cluster near the transition point and become flexible. These sites are considered to be ‘weak spots’ that could initiate unfolding. The higher the frequency of a particular residue to be a weak spot, the greater the likelihood that the corresponding site corresponds to a highly flexible region. We incorporated MD-based ensembles of Cdk4 conformations in different states and used 500 representative samples for each of the trajectories to conduct thermal unfolding simulation on an ensemble of networks created from an ensemble of input structures [151,152]. Using this approach, the frequencies for all residues to become weak spots at the unfolding transition point were computed.

The rigidity profile of the unbound Hsp90 dimer showed that the most stable residues resided in the ATP binding and near the NTD-MD interfaces (Fig 9A and 9B). The distribution peaks corresponded to the nucleotide binding site residues in the NTD (Y33, E42, L43) and catalytic loop in the Hsp90-MD (residues 382–402) with a particularly strong spike at a critical catalytic residue R392 (R380 in yeast Hsp90) that coordinates the γ-phosphate of ATP. Another peak corresponded to a hydrophobic cluster at the NTD-MD interface that is formed by residues W289, P287, F304, Y356 and F208. This interaction cluster is conserved in the Hsp90 proteins and structurally proximal to another group of rigid sites in the Hsp90 dimer (L388, I390 and R392). Together, these clusters of rigid residues are responsible for cross-subunit NTD interactions and stabilization of the catalytic site environment [123]. Notably, the most rigid regions in the unbound Hsp90 dimer are assembled near the nucleotide binding site and the NTD-MD interface (Fig 9A and 9B). Structural mapping of highly stable sites highlighted decomposition of rigidity and flexibility in the Hsp90 dimer (Fig 9C) and Hsp90-Cdc37-Cdk4 complex (Fig 9D). In the Hsp90 dimer, the distribution of rigid residues near the inter-domain interfaces reflected stabilization of the nucleotide binding site,
Coordinated dimerization of the Hsp90-NTDs and association of the Hsp90-NTD with the Hsp90-MD. Central to this analysis is the global propagation of rigidity in the chaperone complex, where the rigidity of the NTD regions was reduced, while and structural environment in the MD-CTD inter-domain regions was considerably rigidified (Fig 9B). During chaperone cycle, upon ATP hydrolysis, Hsp90 dimer is expected to open NTDs with Cdc37-Cdk4 still bound—a necessary step that enables kinase to refold before leaving the complex [65, 69].

We monitored changes in the rigidity and flexibility of Cdk4 client (Fig 10) in the Cdk4-cyclin D3 complex, Cdk4 unbound form and in the Hsp90-Cdc37-Cdk4 complex. Strikingly, even in the complex with cyclin, we observed polarization of kinase lobes, with a mostly rigid C-lobe and a more dynamic N-lobe. The weak spots in the Cdk4-cyclin D3 complex and Cdk4...
The unbound form corresponded to the intrinsically flexible β3-αC loop and β4-β5 strand regions (Fig 10A and 10B). These regions are targeted by the Hsp90-Cdc37 and become partially unfolded in the Hsp90-Cdc37-Cdk4 (Fig 10C). The β3-αC loop plays an important role in modulating dynamics of the adjacent regulatory αC-helix, thus exerting tight control over kinase activity [153]. According to our findings, these N-lobe regions may serve as initiation sites for kinase unfolding and present weak spots of kinase instability that are targeted by the chaperone. Structural mapping of unfolding nuclei residues showed the evolution and gradual expansion of the weak spot clusters (Fig 10D and 10F). Of special significance was the emergence of localized weak spots in the Cdk4-cyclin D3 complex, where the β3-αC loop and β4-β5 were the most flexible regions (Fig 10D). In the cyclin-free, monomeric form of Cdk4, we
observed a gradual expansion of the weak spot regions (Fig 10E), where large fractions of the N-lobe formed flexible clusters, including the G-loop, $\beta$3-$\alpha$C loop, $\beta$4-$\beta$5, and $\beta$6-$\beta$7 strands. Interestingly, in the monomeric Cdk4 form, the entire layered $\beta$-sheet region of the N-lobe becomes flexible near the unfolding transition point. It is worth noting that the instability of the $\beta$-sheet regions is believed to be a common dynamic characteristic shared by many kinase clients of the Hsp90-Cdc37 chaperone [60]. In all functional forms of Cdk4, the distribution of rigid and flexible regions is not uniform as the C-lobe harbors most of the rigid residues, while the N-lobe residues belonged to flexible regions (Fig 10D and 10F). The important finding of this analysis is evidence for differential stabilization of the kinase lobes in Cdk4 client and the emergence of the primary unfolding weak spot near the $\beta$3-$\alpha$C loop and $\beta$4-$\beta$5 strand of the N-lobe. These characteristic features of the Cdk4 client may be intrinsically present in different functional forms of the kinase client. Consistent with the cryo-electron microscopy structure of the Hsp90-Cdc37-Cdk4 kinase complex [69], network emulation of thermal unfolding showed that these segments would correspond to the weak spots of Cdk4 client. We argue that Cdc37 and Hsp90 can recognize and exploit this signature of the kinase client by targeting the predicted weak spots of kinase instability to unfold and trap the N-lobe of Cdk4 during kinase recruitment.

Our results suggest a mechanism for client kinase recognition by the Hsp90-Cdc37 machine. We propose that differential stabilization of kinase lobes is sensed by Cdc37 cochaperone that initially binds and splits kinase at the lobes, creating platform for Hsp90 intervention. At the next stage, weak spots of the N-lobe, that are most vulnerable to structural perturbations in the dynamic kinase client, can be recognized by the Hsp90 binding site regions. Through a cross-talk between Hsp90 and Cdc37 binding interfaces, the chaperone system effectively traps partially unfolded kinase conformations and allow for conformational remodeling of the client that is initiated from the weak spot sites. The bound clients are held by chaperones in a spatially confined dynamic ensemble, which may increase the chances of refolding by facilitating intramolecular interactions. According to our findings, robust recognition of weak kinase spots and reciprocal dynamic exchange at the Hsp90-kinase interface may represent a plausible mechanistic scenario underlying chaperone-kinase interactions. In the proposed model of chaperone-kinase functional cycle, allosteric interactions between Hsp90 and Cdc37 may facilitate conformational and energetic polarization of the kinase lobes, inducing partial unfolding that is thermodynamically controlled through an entropy transfer between Hsp90 and client kinase. This mechanism may underlie allosteric cooperativity Cdc37 and Hsp90 during initial screening of client folds and subsequent thermodynamic sorting of strong and weak clients, which may be controlled by the Hsp90. The results of this study may help to explain how allosterically regulated machines such as Hsp90-Cdc37 can fulfill their chaperoning function by selectively remodeling the folding energy landscapes of their diverse protein clients.

Materials and methods

MD simulations

All-atom MD simulations have been performed for the structure of the human Hsp90-Cdc37-Cdk4 complex (pdb id 5FWM, 5FWK, 5FWL) [69], Cdk4/cyclin D1 complex (PDB 2W9FZ) [98], Cdk4/cyclin D3 complex (PDB 3G33) [99]. We carried out five independent 500 ns for the human Hsp90-Cdc37-Cdk4 complex and ten independent 500 ns for the Cdk4-cyclin D complexes. The conformational ensembles and computed properties of the complexes were generated using aggregation of all independent trajectories. All structures were obtained from the Protein Data Bank [154]. Structure preparation process included
several main steps previously reported in detail in our studies of molecular chaperones [155,156]. In brief, in this protocol, hydrogen atoms, missing residues and protonation states were assigned using the WHATIF program [157]. The missing segments of in the Hsp90-Cdc37-Cdk4 complex and the kinase crystal structures were modeled using the ArchPRED program [158]. MD simulations were performed using CHARMM22 force field in the NAMD 2.6 package [159]. The simulation steps and specifics of the MD protocol were fully consistent with the workflow previously detailed in our studies of molecular chaperones [155,156]. In brief, the following protocol preceded the 500 ns production stage of simulations. All atoms of the complex were first restrained at their crystal structure positions with a force constant of 10 Kcal mol$^{-1}$ Å$^{-2}$. The system was subjected to the following simulation annealing to ensure the proper equilibration. The temperature was increased from 0K to 500K at a rate of 1K per 1ps and was kept at 500K for 500ps. The temperature was then decreased from 500 K to 300K at a rate of 1K per 1ps and was kept at 300K for additional 500ps. An NPT production simulation was then run on each of the equilibrated structures for 500 ns keeping the temperature at 300 K and constant pressure of 1 atm.

**Distance fluctuations analysis**

In this protein mechanics-based approach [121–123], we evaluated residue stability and flexibility through distance fluctuation analysis of the conformational ensembles for studied systems. The fluctuations of the mean distance between a given residue and all other residues in the ensemble were converted into residue stability indexes that measured the energy cost of the residue deformation during simulations. In this model, the high stability indexes would be associated with residues that display small fluctuations in their distances to all other residues, while small values of this stability parameter would point to more flexible sites that experience large deviations of their inter-residue distances. We computed the fluctuations of the mean distance between each atom within a given residue and the atoms that belong to the remaining residues of the protein. The force constant for each residue is calculated by averaging the distances between the residues over the MD trajectory using the following expression:

$$k_i = \frac{3k_B T}{\langle (d_i - \langle d_i \rangle)^2 \rangle}$$

$$d_i = \langle d_{ij} \rangle_j$$

$d_{ij}$ is the instantaneous distance between residue $i$ and residue $j$, $k_B$ is the Boltzmann constant, $T = 300$K. $\langle \rangle$ denotes an average taken over the MD simulation trajectory and $d_i = \langle d_{ij} \rangle_j$ is the average distance from residue $i$ to all other atoms $j$ in the protein. The sum over $j$ implies the exclusion of the atoms that belong to the residue $i$. The interactions between the $C_a$ atom of residue $i$ and the $C_a$ atom of the neighboring residues $i-1$ and $i+1$ are excluded in the calculation since the corresponding distances are nearly constant. The inverse of these fluctuations yields an effective force constant $k_i$ that describes the ease of moving an atom with respect to the protein structure. The average distance fluctuations profiles defined residue stability indexes and characterized the distribution of stable and flexible regions in the protein structures. Allosteric communication propensities of protein residues are associated with their mechanistic properties and mean square deformations of the inter-residue distances that could be related to the inter-residue commute time [87–90]. In this model, an efficient
allosteric communication proceeds through coupling of spatially separated rigid hinge sites that transmit allosteric structural changes.

**Protein-protein binding free energy calculations**

We employed three independent approaches to compute protein-protein binding energy changes induced by alanine mutations of the interfacial residues in the Hsp90-Cdc37-Cdk4 complex. These approaches included coarse-grained BeAtMuSiC predictor of binding free energy changes [130–132], empirical force field approach FoldX with the all-atom representation of protein structure [133–137], and MM-PBSA method [138–142] that utilizes all-atom MD simulations. In these models, the binding free energy of protein-protein complex is expressed as difference in the folding free energy of the complex and folding free energies of the two protein binding partners:

$$\Delta G_{\text{bind}} = G^{\text{com}} - G^A - G^B$$  \hspace{1cm} (3)

The change of the binding energy due to a mutation can be calculated then as the following:

$$\Delta \Delta G_{\text{bind}} = \Delta G_{\text{bind}}^{\text{mut}} - \Delta G_{\text{bind}}^{\text{wt}}$$  \hspace{1cm} (4)

The BeatMusic approach employs a combination of statistical potentials that describe pairwise inter-residue distances, backbone torsion angles and solvent accessibilities derived from known protein structures [160]. In this model, the change in the binding free energy of protein-protein complex upon mutation is expressed as follows:

$$\Delta \Delta G_{\text{bind}} = \sum_{i=1}^{13} \alpha(SA) \Delta \Delta W_i + \alpha_{14}(SA) \Delta V_+ + \alpha_{15}(SA) \Delta V_- + \alpha_{16}(SA)$$  \hspace{1cm} (5)

where $\Delta \Delta W_i$ describe the free energy change induced by 13 different statistical potentials, the terms $\Delta V_+$ and $\Delta V_-$ account for the packing defects, and the weights $\alpha(SA)$ are empirical functions of the solvent accessibility of the mutated residue identified from experimental folding free energy changes induced by mutations [161].

To obtain ensemble-based estimates of binding free energy changes and permit multiple runs of BeAtMuSiC predictor for independent conformations from MD trajectory, we employed Apache Jmeter open source software Java application (http://jmeter.apache.org) designed to simulate multiple requests to a target server from a user (or group of users). This software is used in bioinformatics applications to configure test plans and perform load testing for batches of multiple server requests, store and save all responses, and calculate statistics [162]. A number of threads were initiated by Apache Jmeter application to process multiple BeAtMuSiC predictor requests to the server, store and analyze results and assess server performance.

Alanine scanning of protein binding interface residues in the Hsp90-Cdc37-Cdk4 structure was also performed using FoldX approach [133–137]. In this model, the structure of the WT Hsp90-Cdc37-Cdk4 complex was optimized by the RepairPDB function within FoldX [136]. The conformations of the alanine mutant complexes were generated from the optimized WT structures by the BuildModel function within FoldX. To calculate binding free energies by FoldX, we used Eq (3) and calculated the folding free energy of the Hsp90-Cdc37-Cdk4 complex and each protein separately for the mutant and WT structures. The change of the binding energy due to a mutation is then calculated according to Eq (4).

Binding energies were also calculated based on the MM-PBSA method that combines the molecular mechanics force field with the Poisson–Boltzmann continuum representation of the...
The total free energy $\Delta G_{\text{bind}}$ in MM-PBSA is expressed as the sum of the gas phase contribution $\Delta G_{\text{MM}}$, the solvation free energy $\Delta G_{\text{solv}}$, and an entropic contribution $-T\Delta S$:

$$
\Delta G_{\text{bind}} = <\Delta G_{\text{MM}}> + <\Delta G_{\text{solv}}> - T\Delta S
$$

The brackets $<>$ denote an average of these contributions calculated over the MD trajectories. The gas-phase contribution $<\Delta G_{\text{MM}}>$ to the binding free energy is the difference in the molecular mechanics energy of the complex and the isolated proteins. This contribution is the sum of the differences in the internal energies $\Delta E_{\text{intra}}$, the van der Waals interaction energy $\Delta E_{\text{vdw}}$, and the electrostatic interaction energy $\Delta E_{\text{elec}}$:

$$
<\Delta G_{\text{MM}}> = \Delta E_{\text{intra}} + \Delta E_{\text{vdw}} + \Delta E_{\text{elec}}
$$

where $E_{\text{bond}}$ is the energy of the bonded terms (bonds, angles, dihedral angles, and improper angles) of a given molecule; $E_{\text{vdw}}$ is the van der Waals energy of the molecule; and $E_{\text{elec}}$ is the electrostatic energy of the molecule. These contributions are calculated according to the CHARMM22 force field and describe the molecular mechanical energy in the gas-phase.

The solvation free energy $\Delta G_{\text{solv}}$ is the difference between the solvation energy of the complex and solvation free energies of the isolated protein and ligand:

$$
\Delta G_{\text{solv}} = G_{\text{solv}}^{\text{complex}} - G_{\text{solv}}^{\text{A}} - G_{\text{solv}}^{\text{B}}
$$

$$
\Delta G_{\text{solv}} = \Delta G_{\text{solv}}^{\text{np}} + \Delta G_{\text{solv}}^{\text{pol}}
$$

All energy terms are calculated using MD trajectories of the Hsp90-Cdc37-Cdk4 complex, which is followed by separation of the complexes into individual protein structures and subsequent minimization of the isolated molecules. All energy terms were calculated for 1,000 frames regularly separated by 500 ps along the 500ns trajectory performed for the complex. Due to large size of the studied system and presence of partially disordered Cdk4 protein in the complex, the entropy calculations are very computationally demanding and can be prone to significant statistical errors [142]. The entropy terms for the binding free energy changes may also cancel each other later assuming that protein structures do not experience significant rearrangements upon mutations. Based on these considerations, the entropy calculations were not performed in our binding free energy calculations.

**Network-based modeling of protein rigidity and thermal unfolding**

We utilized FIRST (Floppy Inclusion and Rigid Substructure Topography) approach [146-150] and the Python-based Constraint Network Analysis (CNA) interfaces [151,152] to build a network of the covalent and noncovalent bond constraints in the protein. In the FIRST approach, hydrogen bonds, salt bridges, and hydrophobic contacts define noncovalent bond constraints that are calculated with an empirical energy function. A hydrogen bond energy $E_{\text{hb}}$ is calculated using a geometry-based empirical function [163] and only hydrogen bonds with an energy below cutoff $E_{\text{cuthb}} = -1.0 \text{ kcal/mol}$ are included in the network. Hydrophobic contacts are considered between all carbon and sulfur atoms separated by a distance less than the sum of their van der Waals radii (1.7 Å for C and 1.8 Å for S) plus a temperature-independent $D_{\text{cutdp}} = 0.25$ Å. In the FIRST approach, rigidity changes are monitored by a gradual removal of hydrogen bonds in the order of increasing strength, keeping all covalent and hydrophobic
interactions and repeating the rigidity analysis at each step, thus decomposing protein structure into rigid and flexible regions. This algorithm determines whether a bond is flexible or rigid and decomposes the constraint network into rigid clusters and flexible regions. A rigid cluster is a set of residue nodes that move together as a rigid body, whereas residues that are not a component of a rigid cluster are assigned to a flexible region. During this process, the weak constraints are removed first while stronger interactions are sustained longer, leading to progressive decomposition into rigid and flexible regions. Using this model, the rigidity index ($r_i$) is calculated for each residue, which is defined by the energy value $E_{\text{cubie}}$ when the noncovalent constraints that involve a given residue are removed from any rigid cluster of the set of rigid clusters. In $C_\alpha$ atom-based representation of a residue adopted here, the average of the two $r_i$ values of the two backbone bonds is taken. In this model, the lower the index value $r_i$ the longer a given residue stays as a part of a rigid cluster during the thermal unfolding simulation, which is defining characteristic of residue rigidity. Thermal unfolding in the FIRST approach is implemented by emulating temperature-dependent unfolding trajectories. During unfolding, non-covalent constraints corresponding to weaker interactions that dissolve at low temperatures are removed from the network first, and each new network is then again decomposed into rigid and flexible clusters. By proceeding from a rigid network at low temperature to a flexible network at high temperature, unfolding phase transitions can be observed, at which point a giant rigid cluster in the network breaks apart into smaller rigid clusters. The identification of weak spots is performed using the CNA software package [151,152]. In this procedure, rigid cluster decompositions immediately before and after folded-unfolded transition are compared, and residues whose $C_\alpha$ atoms are part of the giant cluster before the transition, and leave the giant cluster after transition are identified as locally weak spots in the constraint network. A residue is considered flexible if its $C_\alpha$ atom is either in a flexible region or part of a small rigid cluster of less than four atoms. We employed MD-based ensembles of Cdk4 conformations obtained from multiple simulations of the Cdk4-cyclin D3 complex, unbound Cdk4 form, and Hsp90-Cdc37-Cdk4 complex. 500 representative samples for each of the trajectories were used to perform thermal unfolding simulation on an ensemble of networks created from an ensemble of input structures [151,152]. The frequencies of Cdk4 residues to become weak spots are computed and averaged over conformational ensembles for each of these functional forms. The frequency of all residues being predicted as a weak spot throughout the ensemble is counted and, finally, all weak spots are assigned a rank according to the decreasing order of their frequency.

Supporting information

**S1 Fig. Structural mapping of hinge clusters in the Hsp90-Cdc37-Cdk4 complex.** The hinge clusters are mapped onto the conformational mobility profiles along the slow collective modes. Two alternative 180º views of the Hsp90-Cdc37-Cdk4 structure are presented (A,B). Conformational dynamics profiles were computed by averaging protein motions in the space of five lowest frequency modes. The hinge residues are shown in spheres and colored according to their mobility in the slow modes. The color gradient from blue to red indicates the decreasing structural rigidity (or increasing conformational mobility) of the protein residues. (C) The hinge cluster near the NTD-MD inter-domain region is formed by conserved residues P287, W289, P287, F304, Y356 and F208 residues. A close proximity of this hinge center to the catalytic residue R392 and ATP is highlighted. R392 and ATP are shown in atom-based colored spheres. (D) The hinge clusters formed in the central core of the Hsp90-MD: F376-L343-F433-N436 and L369-I370-V381-L401-I404-V381. (E) The hinge centers near the MD-CTD...
interface are formed by clusters W598-M606-R612 and L455-I441-C521-L525-L533.

TIF

S2 Fig. The residue depth analysis of the Hsp90-Cdc37-Cdk4 complex. The residue depth profiles of the Hsp90-A monomer (A), Hsp90-B monomer (B), Cdc37 (C) and Cdk4 proteins (D). The residue depth values are obtained from MD-based averaging over the conformational ensemble. The distributions are shown in domain-colored bars. The Hsp90-NTD (residues 1–215) is colored in red bars, Hsp90-MD (residues 216–552) is in blue bars, and Hsp90-CTD (residues 553–690) is in green bars. The Cdc37-NTD (residues 1–147) are colored in red bars, The Cdc37-M/C domain residues (residues 148–260) are in blue bars. The Cdk4 N-lobe (residues 1–99) is in red bars and Cdk4 C-lobe (residues 100–295) is in blue bars.

TIF

S3 Fig. Alanine scanning and free energy analysis of the binding interfaces in the Hsp90-Cdc37-Cdk4 complex using MM-PBSA approach. Binding free energy changes obtained through alanine scanning of the interfacial residues in the Hsp90-Cdc37-Cdk4 complex using MM-PBSA approach. Mutation-induced binding free energy changes ΔΔG of the Hsp90 residues involved in the Hsp90-Cdc37 interface (A) and Hsp90-Cdk4 interface (B). Binding free energy changes ΔΔG of the Cdc37 residues forming interfacial contacts with Hsp90 (C) and Cdk4 (D). Binding free energy changes ΔΔG of the Cdk4 residues contributing to the Cdk4-Hsp90 interface (E) and Cdk4-Cdc37 interface (F). If the free energy change between a mutant and the wild type (WT) proteins ΔΔG = ΔG (MT) - ΔG (WT) > 0, the mutation is considered to be destabilizing. The distributions are shown in red-colored bars and highlight only binding interface residues.

TIF

Author Contributions
Conceptualization: Gennady M. Verkhivker.
Data curation: Josh Czemeres, Kurt Buse, Gennady M. Verkhivker.
Formal analysis: Josh Czemeres, Kurt Buse, Gennady M. Verkhivker.
Funding acquisition: Gennady M. Verkhivker.
Investigation: Josh Czemeres, Gennady M. Verkhivker.
Methodology: Gennady M. Verkhivker.
Project administration: Gennady M. Verkhivker.
Resources: Gennady M. Verkhivker.
Software: Gennady M. Verkhivker.
Supervision: Gennady M. Verkhivker.
Validation: Josh Czemeres, Kurt Buse.
Visualization: Josh Czemeres, Kurt Buse.
Writing – original draft: Gennady M. Verkhivker.
Writing – review & editing: Gennady M. Verkhivker.

References
1. Pearl LH, Prodromou C. Structure, function, and mechanism of the Hsp90 molecular chaperone. Adv Protein Chem. 2001; 59: 157–186. PMID: 11868271
2. Richter K, Buchner J. Hsp90: chaperoning signal transduction. J Cell Physiol. 2001; 188: 281–290. https://doi.org/10.1002/jcp.1131 PMID: 11473354

3. Young JC, Moorefi I, Hartl FU. Hsp90: a specialized but essential protein-folding tool. J Cell Biol. 2001; 154: 267–273. https://doi.org/10.1083/jcb.200104079 PMID: 11470816

4. Picard D. Heat-shock protein 90, a chaperone for folding and regulation. Cell Mol Life Sci. 2002; 59: 1640–1648. PMID: 12475174

5. Young JC, Agashe VR, Siegers K, Hartl UF. Pathways of chaperone-mediated protein folding in the cytosol. Nat Rev Mol Cell Biol. 2004; 5: 781–791. https://doi.org/10.1038/nrm1492 PMID: 15459659

6. Lindquist SL. HSP90 and the chaperoning of cancer. Nat Rev Cancer. 2005; 5: 761–772. https://doi.org/10.1038/nrc1716 PMID: 16175177

7. McClellan AJ, Xia Y, Deutschbauer AM, Davis RW, Gerstein M, Frydman J. Diverse cellular functions of the Hsp90 molecular chaperone uncovered using systems approaches. Cell. 2007; 131: 121–135. https://doi.org/10.1016/j.cell.2007.07.036 PMID: 17923092

8. Zhao R, Houry WA. Molecular interaction network of the Hsp90 chaperone system. Adv Exp Med Biol. 2007; 594: 27–36. https://doi.org/10.1007/978-0-387-39975-1_3 PMID: 17205672

9. Krukenberg KA, Street TO, Lavery LA, Agard DA. Conformational dynamics of the molecular chaperone Hsp90. Q Rev Biophys. 2011; 44: 229–340. https://doi.org/10.1017/S0033581511000314 PMID: 21414251

10. Jackson SE. Hsp90: structure and function. Top Curr Chem. 2013; 328: 155–240. https://doi.org/10.1007/128_2012_356 PMID: 22955504

11. Li J, Buchner J. Structure, function and regulation of the Hsp90 machinery. Biomed J. 2013; 36: 106–117. https://doi.org/10.4103/2319-4170.113230 PMID: 2306880

12. Prodromou C. Mechanisms of Hsp90 regulation. Biochem J. 2016; 473: 2439–2452. https://doi.org/10.1042/BCJ20160005 PMID: 27515256

13. Pearl LH. Review: The HSP90 molecular chaperone-an enigmatic ATPase. Biopolymers. 2016; 105: 594–607. https://doi.org/10.1002/bip.22835 PMID: 26991466

14. Ali MM, Roe SM, Vaughan CK, Meyer P, Panaretou B, Piper PW, et al. Crystal structure of an Hsp90-nucleotide-p23/Sba1 closed chaperone complex. Nature. 2006; 440: 1013–1017. https://doi.org/10.1038/nature04716 PMID: 16625188

15. Shiau AK, Harris SF, Southworth DR, Agard DA. Structural Analysis of E. coli Hsp90 reveals dramatic nucleotide-dependent conformational rearrangements. Cell. 2006; 127: 329–340. https://doi.org/10.1016/j.cell.2006.09.027 PMID: 17055434
25. Dollins DE, Warren JJ, Immormino RM, Gewirth DT. Structures of GRP94-nucleotide complexes reveals mechanistic differences between the hsp90 chaperones. Mol Cell. 2007; 28: 41–56. https://doi.org/10.1016/j.molcel.2007.08.024 PMID: 17936703

26. Graf C, Stankiewicz M, Kramer G, Mayer MP. Spatially and kinetically resolved changes in the conformational dynamics of the Hsp90 chaperone machine. EMBO J. 2009; 28: 602–613. https://doi.org/10.1038/emboj.2008.306 PMID: 19165152

27. Mickler M, Hessling M, Ratzke C, Buchner J, Hugel T. The large conformational changes of Hsp90 are only weakly coupled to ATP hydrolysis. Nat Struct Mol Biol. 2009; 16: 281–286. https://doi.org/10.1038/nstruct.1557 PMID: 19234469

28. Hessling M, Richter K, Buchner J. Dissection of the ATP-induced conformational cycle of the molecular chaperone Hsp90. Nat Struct Mol Biol. 2009; 16: 287–293. https://doi.org/10.1038/nsmb.1565 PMID: 19234467

29. Ratzke C, Mickler M, Hellenkamp B, Buchner J, Hugel T. Dynamics of heat shock protein 90 C-terminal dimerization is an important part of its conformational cycle. Proc Natl Acad Sci USA. 2010; 107: 16101–16106. https://doi.org/10.1073/pnas.1009916107 PMID: 20736353

30. Ratzke C, Berkemeier F, Hugel T. Heat shock protein 90’s mechanochemical cycle is dominated by thermal fluctuations. Proc Natl Acad Sci USA. 2012; 109: 161–166. https://doi.org/10.1073/pnas.1107930108 PMID: 22184223

31. Ratzke C, Nguyen MN, Mayer MP, Hugel T. From a ratchet mechanism to random fluctuations evolution of Hsp90’s mechanochemical cycle. J Mol Biol. 2012; 423: 462–471. https://doi.org/10.1016/j.jmb.2012.07.026 PMID: 22878379

32. Graf C, Lee CT, Eva Meier-Andrejsek L, Nguyen MT, Mayer MP. Differences in conformational dynamics within the Hsp90 chaperone family reveal mechanistic insights. Front Mol Biosci. 2014; 1: 4. https://doi.org/10.3389/fmbio.2014.00004 PMID: 25988145

33. Zierer BK, Rübbelke M, Tippel F, Madl T, Schopf FH, Rutz DA, et al. Importance of cycle timing for the thermal fluctuation. Proc Natl Acad Sci USA. 2012; 109: 161–166. https://doi.org/10.1073/pnas.1107930108 PMID: 22184223

34. Schulze A, Beliu G, Helmerich DA, Schubert J, Pearl LH, Prodromou C, et al. Cooperation of local motions in the Hsp90 molecular chaperone ATPase mechanism. Nat Chem Biol. 2016; 12: 628–635. https://doi.org/10.1038/nchembio.2111 PMID: 27322067

35. Johnson JL. Evolution and function of diverse Hsp90 homologs and co-chaperone proteins. Biochim Biophys Acta. 2012; 1823: 607–613. https://doi.org/10.1016/j.bbamcr.2011.09.020 PMID: 22008467

36. Prodromou C. The ‘active life’ of Hsp90 complexes. Biochim Biophys Acta. 2012; 1823: 624–635. https://doi.org/10.1016/j.bbamcr.2011.09.003 PMID: 21951723

37. Li J, Soroka J, Buchner J. The Hsp90 chaperone machinery: conformational dynamics and regulation by cochaperones. Biochim Biophys Acta. 2012; 1823: 624–635. https://doi.org/10.1016/j.bbamcr.2011.09.003 PMID: 21951723

38. Makhnevych T, Houry WA. The role of Hsp90 in protein complex assembly. Biochim Biophys Acta. 2012; 1823: 674–682. https://doi.org/10.1016/j.bbamcr.2011.09.001 PMID: 21945180

39. Röhi A, Rohrbarg J, Buchner J. The chaperone Hsp90: changing partners for demanding clients. Trends Biochem Sci. 2013; 38: 253–262. https://doi.org/10.1016/j.tibs.2013.02.003 PMID: 23507089

40. Mayer MP, Le Breton L. Hsp90: breaking the symmetry. Mol Cell. 2015; 58: 8–20. https://doi.org/10.1016/j.molcel.2015.02.022 PMID: 25839432

41. Verma S, Goyal S, Jamal S, Singh A, Grover A, Hsp90: Friends, clients and natural foes. Biochimie. 2016; 127: 227–240. https://doi.org/10.1016/j.biochim.2016.05.018 PMID: 27295069

42. Li J, Richter K, Reinstein J, Buchner J. Integration of the accelerator Aha1 in the Hsp90 cochaperone cycle. Nat Struct Mol Biol. 2013; 20: 326–331. https://doi.org/10.1038/nsmb.2502 PMID: 23996352

43. Soroka J, Wandinger SK, Mäusbacher N, Schreiber T, Richter K, Daub H, et al. Conformational switching of the molecular chaperone Hsp90 via regulated phosphorylation. Mol Cell. 2012; 45: 517–528. https://doi.org/10.1016/j.molcel.2011.12.031 PMID: 22365831

44. Mollapour M, Tsutsumi S, Truman AW, Xu W, Vaughan CK, Beeke K, et al. Threonine 22 phosphorylation attenuates Hsp90 interaction with cochaperones and affects its chaperone activity. Mol Cell. 2011; 41: 672–681. https://doi.org/10.1016/j.molcel.2011.02.011 PMID: 21419342

45. Mollapour M, Neckers L. Post-translational modifications of Hsp90 and their contributions to chaperone regulation. Biochim Biophys Acta. 2012; 1823: 648–655. https://doi.org/10.1016/j.bbamcr.2011.07.018 PMID: 21856339

46. Xu W, Mollapour M, Prodromou C, Wang S, Scroggins BT, Palchick Z et al. Dynamic tyrosine phosphorylation modulates cycling of the HSP90-P50(CDC37)-AHA1 chaperone machine. Mol Cell. 2012; 47: 434–443. https://doi.org/10.1016/j.molcel.2012.05.015 PMID: 22727666
47. Woodford MR, Dunn D, Miller JB, Jamal S, Neckers L, Mollapour M. Impact of posttranslational modifications on the anticancer activity of Hsp90 inhibitors. Adv Cancer Res. 2016; 129: 31–50. https://doi.org/10.1016/bs.acr.2015.09.002 PMID: 26916000

48. Lu XA, Wang X, Zhuo W, Jia L, Jiang Y, Fu Y, et al. The regulatory mechanism of a client kinase controlling its own release from Hsp90 chaperone machinery through phosphorylation. Biochem J. 2014; 457: 171–183. https://doi.org/10.1042/BJ20130963 PMID: 24117238

49. Beebe K, Mollapour M, Scroggins B, Prodromou C, Xu W, Tokita M, et al. Posttranslational modification and conformational state of heat shock protein 90 differentially affect binding of chemically diverse small molecule inhibitors, Oncotarget. 2013; 4: 1065–1074. https://doi.org/10.18632/oncotarget.1099 PMID: 23867252

50. MacLean M, Picard D. Cdc37 goes beyond Hsp90 and kinases. Cell Stress Chaperones. 2003; 8: 114–119. PMID: 16627196

51. Pearl LH. Hsp90 and Cdc37—a chaperone cancer conspiracy. Curr Opin Genet Dev. 2005; 15: 55–61. https://doi.org/10.1016/j.gde.2004.12.011 PMID: 15661534

52. Mandal AK, Lee P, Chen JA, Nillegoda N, Heller A, DiStasio S, et al. Cdc37 has distinct roles in protein kinase quality control that protect nascent chains from degradation and promote posttranslational maturation. J Cell Biol. 2007; 176: 319–328. https://doi.org/10.1083/jcb.200604106 PMID: 17242065

53. Mandal AK, Theodoraki MA, Nillegoda NB, Caplan AJ. Role of molecular chaperones in biogenesis of the protein kinome. Methods Mol Biol. 2011; 787: 75–81. https://doi.org/10.1007/978-1-61779-295-3_6 PMID: 21898228

54. Karnitz LM, Felts SJ. Cdc37 regulation of the kinome: when to hold ‘em and when to fold’ em. Sci Signal Transduct Knowl Environ. 2007; 385: pe22.

55. Scroggins BT, Prince T, Shao J, Uma S, Huang W, Guo Y, et al. High affinity binding of Hsp90 is triggered by multiple discrete segments of its kinase clients. Biochemistry. 2003; 42: 12550–12561. https://doi.org/10.1021/bi035001t PMID: 14580201

56. Prince T, Matts RL. Definition of protein kinase sequence motifs that trigger high affinity binding of Hsp90 and Cdc37. J Biol Chem. 2004; 279: 39975–39981. https://doi.org/10.1074/jbc.M406882200 PMID: 15258137

57. Prince T, Sun L, Matts RL. Cdk2: a genuine protein kinase client of Hsp90 and Cdc37. Biochemistry. 2005; 44: 15287–15295. https://doi.org/10.1021/bi051423m PMID: 16285732

58. Citri A, Harari D, Shohat G, Ramakrishnan P, Gan J, Lavi S, et al. Hsp90 recognizes a common surface on client kinases J Biol Chem. 2006; 281: 14361–14369. https://doi.org/10.1074/jbc.M512613200 PMID: 16543424

59. Xu W, Yuan X, Xiang Z, Mimnaugh E, Marcu M, Neckers L. Surface charge and hydrophobicity determine ErbB2 binding to the Hsp90 chaperone complex. Nat Struct Mol Biol. 2005; 12: 120–126. https://doi.org/10.1038/nsmb885 PMID: 15643424

60. Taipale M, Krykbaeva I, Koeva M, Kayatekin C, Karras GI, et al. Quantitative analysis of HSP90-client interactions reveals principles of substrate recognition. Cell. 2012; 150: 987–1001. https://doi.org/10.1016/j.cell.2012.06.047 PMID: 22939624

61. Boczek EE, Reefschlager LG, Dehling M, Struller TJ, Häusler E, Seidl A, et al. Conformational processing of oncogenic v-Src kinase by the molecular chaperone Hsp90. Proc Natl Acad Sci U S A. 2015; 112: E3189–E198. https://doi.org/10.1073/pnas.1424342112 PMID: 26056257

62. Smith JR, Clarke PA, de Billy E, Workman P. Silencing the cochaperone CDC37 destabilizes kinase clients and sensitizes cancer cells to HSP90 inhibitors. Oncogene. 2009; 28: 157–169. https://doi.org/10.1038/onc.2008.380 PMID: 18931700

63. Smith JR, de Billy E, Hobbs S, Powers M, Prodromou C, Pearl L, et al. Restricting direct interaction of CDC37 with HSP90 does not compromise chaperoning of client proteins. Oncogene. 2015; 34: 15–26. https://doi.org/10.1038/onc.2013.519 PMID: 24292678

64. Eckl JM, Scherr MJ, Freiburger L, Daake MA, Sattler M, Richter K. Hsp90 Cdc37 complexes with protein kinases form cooperatively with multiple distinct interaction sites. J Biol Chem. 2015; 290: 30843–30854. https://doi.org/10.1074/jbc.M115.693150 PMID: 26511315

65. Keramisanou D, Aboalroub A, Zhang Z, Liu W, Marshall D, Diviney A, et al. Molecular mechanism of protein kinase recognition and sorting by the Hsp90 kinome-specific cochaperone Cdc37. Mol Cell. 2016; 62: 260–271. https://doi.org/10.1016/j.molcel.2016.04.005 PMID: 27105117

66. Genest O, Reidy M, Street TO, Hoskins JR, Camberg JL, Agard DA, et al. Uncovering a region of heat shock protein 90 important for client binding in E. coli and chaperone function in yeast. Mol Cell. 2013; 49: 464–473. https://doi.org/10.1016/j.molcel.2012.11.017 PMID: 23260660
67. Lorenz OR, Freiburger L, Rutz DA, Krause M, Zierer BK, Alvir S, et al. Modulation of the hsp90 chaperone cycle by a stringent client protein. Mol. Cell. 2014; 53: 941–953. https://doi.org/10.1016/j.molcel.2014.02.003 PMID: 24613341

68. Vaughan CK, Gohlke U, Sobott F, Good VM, Ali MM, Prodromou C, et al. Structure of an Hsp90-Cdc37-Cdk4 complex. Mol. Cell. 2006; 23: 697–707. https://doi.org/10.1016/j.molcel.2006.07.016 PMID: 16949366

69. Verba KA, Wang RY, Arakawa A, Liu Y, Shiouzu M, Yokoyama S, et al. Atomic structure of Hsp90-Cdc37-Cdk4 reveals that Hsp90 traps and stabilizes an unfolded kinase. Science. 2016; 352: 1542–1547. https://doi.org/10.1126/science.aaf5023 PMID: 27339980

70. Verba KA, Agard DA. How Hsp90 and Cdc37 lubricate kinase molecular switches. Trends Biochem Sci. 2017; 42: 799–811. https://doi.org/10.1016/j.tibs.2017.07.002 PMID: 28784328

71. Chen B, Piel WH, Gui L, Bruford E, Monteiro A. The HSP90 family of genes in the human genome: insights into their divergence and evolution. Genomics. 2005; 86: 627–637. https://doi.org/10.1016/j.ygeno.2005.08.012 PMID: 16269234

72. Chen B, Zhong D, Monteiro A. Comparative genomics and evolution of the HSP90 family of genes across all kingdoms of organisms. BMC Genomics. 2006; 7: 156. https://doi.org/10.1186/1471-2164-7-156 PMID: 16780600

73. Synoradzki K, Bieganski P. Middle domain of human Hsp90 isoforms differentially binds Aha1 in human cells and alters Hsp90 activity in yeast. Biochim Biophys Acta. 2015; 1853: 445–452. https://doi.org/10.1016/j.bbamcr.2014.11.026 PMID: 25486457

74. Marzec M, Eletto D, Argon Y. GRP94: An HSP90-like protein specialized for protein folding and quality control in the endoplasmic reticulum. Biochim Biophys Acta. 2012; 1823: 774–787. https://doi.org/10.1016/j.bbamcr.2011.10.013 PMID: 22079671

75. Karagöz GE, Duarte AM, Ippel H, Uetrecht C, Sinnige T, van Rosmalen M, et al. N-terminal domain of human Hsp90 triggers binding to the cochaperone p23. Proc Natl Acad Sci USA. 2011; 108: 580–585. https://doi.org/10.1073/pnas.1011867108 PMID: 21183720

76. Tsutsumi S, Mollapour M, Prodromou C, Lee CT, Panaretou B, Yoshida S, et al. Charged linker sequence modulates eukaryotic heat shock protein 90 (Hsp90) chaperone activity. Proc Natl Acad Sci USA. 2012; 109: 2937–2942. https://doi.org/10.1073/pnas.111441109 PMID: 22315411

77. Hainzl O, Lapina MC, Buchner J, Richter K. The charged linker region is an important regulator of Hsp90 function. J Biol Chem. 2009; 284: 22559–22567. https://doi.org/10.1074/jbc.M109.031658 PMID: 19553666

78. Jahn M, Rehn A, Pelz B, Hellenkamp B, Richter K, Rief M, et al. The charged linker of the molecular chaperone Hsp90 modulates domain contacts and biological function Proc Natl Acad Sci USA. 2014; 111: 17881–17886. https://doi.org/10.1073/pnas.1414073111 PMID: 25488691

79. Colombo G, Morra G, Meli M, Verkhivker G. Understanding ligand-based modulation of the Hsp90 molecular chaperone dynamics at atomic resolution. Proc Natl Acad Sci USA 2008; 105: 7976–7981. https://doi.org/10.1073/pnas.0802879105 PMID: 18511558

80. Morra G, Verkhivker G, Colombo G. Modeling signal propagation mechanisms and ligand-based conformational dynamics of the Hsp90 molecular chaperone full length dimer. PLoS Comput Biol. 2009; 5: e1000323. https://doi.org/10.1371/journal.pcbi.1000323 PMID: 19300478

81. Verkhivker GM, Dixit A. Structural and computational biology of the molecular chaperone Hsp90: from understanding molecular mechanisms to computer-based inhibitor design. Curr Top Med Chem. 2009; 9: 1369–1385. PMID: 19860735

82. Dixit A, Verkhivker GM. Probing molecular mechanisms of the Hsp90 chaperone: Biophysical modeling identifies key regulators of functional dynamics. PLoS One. 2012; 7: e37605. https://doi.org/10.1371/journal.pone.0037605 PMID: 22624053
87. Blacklock K, Verkhivker G. Differential modulation of functional dynamics and allosteric interactions in the Hsp90-cochaperone complexes with p23 and Aha1: a computational study. PLoS One. 2013; 8: e71936. https://doi.org/10.1371/journal.pone.0071936 PMID: 23977182

88. Blacklock K, Verkhivker G. Experimentally guided structural modeling and dynamics analysis of Hsp90-p53 interactions: allosteric regulation of the Hsp90 chaperone by a client protein. J Chem Inf Model. 2013; 53: 2962–2978. https://doi.org/10.1021/ci400434g PMID: 24191708

89. Blacklock K, Verkhivker GM. Allosteric regulation of the Hsp90 dynamics and stability by client recruiter cochaperones: protein structure network modeling. PLoS One. 2014; 9: e86547. https://doi.org/10.1371/journal.pone.0086547 PMID: 24466147

90. Blacklock K, Verkhivker GM. Computational modeling of allosteric regulation in the Hsp90 chaperones: a statistical ensemble analysis of protein structure networks and allosteric communications. PLoS Comput Biol. 2014; 10: e1003679. https://doi.org/10.1371/journal.pcbi.1003679 PMID: 24922508

91. Rehn A, Moroni E, Zierer BK, Tippel F, Morra G, John C, et al. Allosteric regulation points control the conformational dynamics of the molecular chaperone Hsp90. J Mol Biol. 2016; 428: 4559–4571. https://doi.org/10.1016/j.jmb.2016.09.014 PMID: 27663270

92. Vettoretti G, Moroni E, Sattin S, Tao J, Agard DA, Bernardi A, et al. Molecular dynamics simulations reveal the mechanisms of allosteric activation of Hsp90 by designed ligands. Sci Rep. 2016; 6: 23830. https://doi.org/10.1038/srep23830 PMID: 27032695

93. Tsai CJ, Sol AD, Nussinov R. Protein allostery, signal transmission and dynamics: a classification scheme of allosteric mechanisms. Mol Biosyst. 2009; 5: 207–216. https://doi.org/10.1039/b819720b PMID: 19225609

94. Csermely P, Palotai R, Nussinov R. Induced fit, conformational selection and independent dynamic segments: an extended view of binding events. Trends Biochem Sci. 2010; 35: 539–546. https://doi.org/10.1016/j.tibs.2010.04.009 PMID: 20541943

95. Tsai CJ, Nussinov R. A unified view of "how allostery works". PLoS Comput Biol. 2014; 10: e1003394. https://doi.org/10.1371/journal.pcbi.1003394 PMID: 24516370

96. Hilser VJ, Wrabl JO, Motlagh HN. Structural and energetic basis of allostery. Annu Rev Biophys. 2012; 41: 585–609. https://doi.org/10.1146/annurev-biophys-050511-102319 PMID: 22577828

97. Motlagh HN, Wrabl JO, Li J, Hilser VJ. The ensemble nature of allostery. Nature. 2014; 508: 331–339. https://doi.org/10.1038/nature13001 PMID: 24740064

98. Day PJ, Cleasby A, Tickle IJ, O'Reilly M, Coyle JE, Holding FP, et al. Crystal structure of human CDK4 in complex with a D-type cyclin. Proc Natl Acad Sci USA. 2009; 106: 4166–4170. https://doi.org/10.1073/pnas.0809645106 PMID: 19237565

99. Takaki T, Echalier A, Brown NR, Hunt T, Endicott JA, Noble ME. The structure of CDK4/cyclin D3 has implications for models of CDK activation. Proc Natl Acad Sci USA. 2009; 106: 4171–4176. https://doi.org/10.1073/pnas.0809674106 PMID: 19237555

100. Polier S, Samant RS, Clarke PA, Workman P, Prodromou C, Pearl LH. ATP-competitive inhibitors block protein kinase recruitment to the Hsp90-Cdc37 system. Nat Chem Biol. 2013; 9: 307–312. https://doi.org/10.1038/ncb.2524 PMID: 23502424

101. Eckl JM, Daake M, Schwartz S, Richter K. Nucleotide-Free sB-Raf is Preferentially Bound by Hsp90 and Cdc37 In Vitro. J Mol Biol. 2016; 428: 4185–4196. https://doi.org/10.1016/j.jmb.2016.09.002 PMID: 27620500

102. Schopf FH, Biebl MM, Buchner J. The HSP90 chaperone machinery. Nat Rev Mol Cell Biol. 2017; 18: 345–360. https://doi.org/10.1038/nrm.2017.20 PMID: 28429788

103. Guo J, Zhou HX. Dynamically driven protein allostery exhibits disparate responses for fast and slow motions. Biophys J. 2015; 108: 2771–27744. https://doi.org/10.1016/j.bpj.2015.04.035 PMID: 26083915

104. Guo J, Zhou HX. Protein allostery and conformational dynamics. Chem Rev. 2016; 116: 6503–6515. https://doi.org/10.1021/acs.chemrev.5b00590 PMID: 26876046

105. Guo J, Zhou HX. Allosteric activation of SENP1 by SUMO1 β-grasp domain involves a dock-and-coalesce mechanism. Elife. 2016; 5: e18249. https://doi.org/10.7554/eLife.18249 PMID: 27576863

106. Amadell A, Linssen AB, Berendsen HJ. Essential dynamics of proteins. Proteins. 1993; 17: 412–425. https://doi.org/10.1002/pro.100170408 PMID: 8108382

107. Hengler-Wildman KA, Lei M, Thai V, Kerns SJ, Karplus M, Kern D. A hierarchy of timescales in protein dynamics is linked to enzyme catalysis. Nature. 2007; 450: 913–916. https://doi.org/10.1038/nature06407 PMID: 18026087
108. Cunningham CN, Krukenberg KA, Agard DA. Intra- and intermonomer interactions are required to synergistically facilitate ATP hydrolysis in Hsp90. J Biol Chem. 2008; 283: 21170–21178. https://doi.org/10.1074/jbc.M800046200 PMID: 18492664

109. Jha SK, Udgaonkar JB. Direct evidence for a dry molten globule intermediate during the unfolding of a small protein. Proc Natl Acad Sci U S A. 2009; 106: 12289–12294. https://doi.org/10.1073/pnas.0905744106 PMID: 19617531

110. Chakravarty S, Varadarajan R. Residue depth: a novel parameter for the analysis of protein structure and stability. Structure. 1999; 7: 723–722. PMID: 10425675

111. Tan KP, Nguyen TB, Patel S, Varadarajan R, Madhusudhan MS. Depth: a web server to compute depth, cavity sizes, detect potential small-molecule ligand-binding cavities and predict the pKa of ionizable residues in proteins. Nucleic Acids Res. 2013; 41: W314–W321. https://doi.org/10.1093/nar/gkt503 PMID: 23766289

112. Song J, Tan H, Mahmood K, Law RH, Buckle AM, Webb GI, et al. Prodepth: predict residue depth by support vector regression approach from protein sequences only. PLoS One. 2009; 4: e7072. https://doi.org/10.1371/journal.pone.0007072 PMID: 19759917

113. He L, Sharpe T, Mazur A, Hiller S. A molecular mechanism of chaperone-client recognition. Sci Adv. 2016; 2: e1601625. https://doi.org/10.1126/sciadv.1601625 PMID: 28138538

114. Burmann BM, Wang C, Hiller S. Conformation and dynamics of the periplasmic membrane-protein-chaperone complexes OmpX-Skp and tOmpA-Skp. Nat Struct Mol Biol. 2013; 20: 1265–1272. https://doi.org/10.1038/nsmb.2677 PMID: 24077225

115. Callon M, Burmann BM, Hiller S. Structural mapping of a chaperone-substrate interaction surface. Angew Chem Int Ed Engl. 2014; 53: 5069–5072. https://doi.org/10.1002/anie.201310963 PMID: 24700611

116. Karagöz GE, Rüdiger SG. Hsp90 interaction with clients. Trends Biochem Sci. 2015; 40: 117–125. https://doi.org/10.1016/j.tibs.2014.12.002 PMID: 25579468

117. Bardwell JC, Jakob U. Conditional disorder in chaperone action. Trends Biochem Sci. 2012; 37: 517–525. https://doi.org/10.1016/j.tibs.2012.08.006 PMID: 23018052

118. Karagöz GE, Duarte AM, Akoury E, Ippel H, Biernat J, Morán Luengo T, et al. Hsp90-Tau complex reveals molecular basis for specificity in chaperone action. Cell. 2014; 156: 963–974. https://doi.org/10.1016/j.cell.2014.01.037 PMID: 24581495

119. Tompa P, Csermely P. The role of structural disorder in the function of RNA and protein chaperones. FASEB J. 2004; 18: 1169–1175. https://doi.org/10.1096/fj.04-1584rev PMID: 15284216

120. van der Lee R, Buljani M, Lang B, Weatheritt RJ, Daughdrill GW, Dunker AK, et al. Classification of intrinsically disordered regions and proteins. Chem Rev. 2014; 114: 6589–6631. https://doi.org/10.1021/cr400525m PMID: 24773235

121. Navizet I, Cailliez F, Lavery R. Probing protein mechanics: residue-level properties and their use in defining domains. Biophys J. 2004; 87: 1426–1435. https://doi.org/10.1529/biophysj.104.042085 PMID: 15345525

122. Sacquin-Mora S, Lavery R. Investigating the local flexibility of functional residues in hemoproteins. Biophys J. 2006; 90: 2706–2717. https://doi.org/10.1529/biophysj.105.074997 PMID: 16428284

123. Sacquin-Mora S, Laforet E, Lavery R. Locating the active sites of enzymes using mechanical properties. Proteins 2007; 67: 350–359. https://doi.org/10.1002/prot.21353 PMID: 17311346

124. Verkhivker GM. Network-based modelling and percolation analysis of conformational dynamics and activation in the CDK2 and CDK4 proteins: dynamic and energetic polarization of the kinase lobes may determine divergence of the regulatory mechanisms. Mol Biosyst. 2017; https://doi.org/10.1039/c7mb00355b PMID: 28926061

125. Paladino A, Morra G, Colombo G. Structural stability and flexibility direct the selection of activating mutations in epidermal growth factor receptor kinase. J Chem Inf Model. 2015; 55: 1377–1387. https://doi.org/10.1021/acs.jcim.5b00270 PMID: 26121158

126. Kiel C, Benisty H, Lloréns-Rico V, Serrano L. The yin-yang of kinase activation and unfolding explains the peculiarity of Val600 in the activation segment of BRAF. Elife. 2016; 5:e12814. https://doi.org/10.7554/eLife.12814 PMID: 28744778

127. Dellus-Gur E, Toth-Petroczy A, Elias M, Tawfik DS. What makes a protein fold amenable to functional innovation? Fold polarity and stability trade-offs. J Mol Biol. 2013; 425: 2609–2621. https://doi.org/10.1016/j.jmb.2013.03.033 PMID: 23542341

128. Tóth-Pétröczy A, Tawfik DS. The robustness and innovability of protein folds. Curr Opin Struct Biol. 2014; 26: 131–138. https://doi.org/10.1016/j.sbi.2014.06.007 PMID: 25038399

129. Tokuriki N, Tawfik DS. Protein dynamism and evolvability Science. 2009; 324: 203–207. https://doi.org/10.1126/science.1169375 PMID: 19359777
130. Dehouck Y, Kwasiygroch JM, Rooman M, Gilis D. BeAtMuSiC: Prediction of changes in protein-protein binding affinity on mutations. Nucleic Acids Res. 2013; 41: W333–W339. https://doi.org/10.1093/nar/gkt450 PMID: 23723246

131. Moretti R, Fleishman SJ, Agius R, Torchala M, Bates PA, Kastritis PL, et al. Community-wide evaluation of methods for predicting the effect of mutations on protein-protein interactions. Proteins. 2013; 81: 1980–1987. https://doi.org/10.1002/prot.24356 PMID: 23843247

132. Moal IH, Fernandez-Recio J. SKEMPI: a Structural Kinetic and Energetic database of Mutant Protein Interactions and its use in empirical models. Bioinformatics. 2012; 28: 2600–2607. https://doi.org/10.1093/bioinformatics/bts489 PMID: 22859501

133. Guerois R, Nielsen JE, Serrano L. Predicting changes in the stability of proteins and protein complexes: a study of more than 1000 mutations. J Mol Biol. 2002; 320: 369–87. https://doi.org/10.1016/S0022-2836(02)00442-4 PMID: 12079393

134. Tokuriki N, Stricher F, Schymkowitz J, Serrano L, Tawfik DS. The stability effects of protein mutations appear to be universally distributed. J Mol Biol. 2007; 369: 1318–1332. https://doi.org/10.1016/j.jmb.2007.03.069 PMID: 17482644

135. Schymkowitz JW, Rousseau F, Martins IC, Ferkinghoff-Borg J, Stricher F, Serrano L. Prediction of water and metal binding sites and their affinities by using the Fold-X force field. Proc Natl Acad Sci U S A. 2005; 102: 10147–10152. https://doi.org/10.1073/pnas.0501980102 PMID: 16006526

136. Schymkowitz J, Borg J, Stricher F, Nys R, Rousseau F, Serrano L. The FoldX web server: an online force field. Nucleic Acids Res. 2005; 33: W382–W388. https://doi.org/10.1093/nar/gki387 PMID: 15980494

137. Van Durme J, Delgado J, Stricher F, Serrano L, Schymkowitz J, Rousseau F. A graphical interface for the FoldX forcefield. Bioinformatics. 2011; 27: 1711–1712. https://doi.org/10.1093/bioinformatics/btr254 PMID: 21505037

138. Srinivasan J, Cheatham TE, Cieplak P, Kollman PA, Case DA. Continuum solvent studies of the stability of DNA, RNA and phosphoramidate-DNA helices. J Am Chem Soc 1998; 120:9401–9409.

139. Kollman PA, Massova I, Reyes C, Kuhn B, Huo S, Chong L, et al. Calculating structures and free energies of complex molecules: combining molecular mechanics and continuum models. Acc Chem Res. 2000; 33: 889–897. PMID: 11123888

140. Homeyer N, Gohlke H. Free energy calculations by the Molecular Mechanics Poisson-Boltzmann Surface Area Method. Mol Inform. 2012; 31: 114–122. https://doi.org/10.1002/minf.201100135 PMID: 27476956

141. Zhang Z, Wang L, Gao Y, Zhang J, Zhenirovskyy M, Alexov E. Predicting folding free energy changes upon single point mutations. Bioinformatics. 2012; 28: 664–671. https://doi.org/10.1093/bioinformatics/bts005 PMID: 22238268

142. Li M, Petukh M, Alexov E, Panchenko AR. Predicting the impact of missense mutations on protein-protein binding affinity. J Chem Theory Comput. 2014; 10: 1770–1780. https://doi.org/10.1021/ct401022c PMID: 24803870

143. Zhang W, Hirshberg M, McLaughlin SH, Lazar GA, Grossmann JG, Nielsen PR, et al. Biochemical and structural studies of the interaction of Cdc37 with Hsp90. J Mol Biol. 2004; 340: 891–907. https://doi.org/10.1016/j.jmb.2004.05.007 PMID: 15223329

144. Sreeramulu S, Jonker HR, Langer T, Richter C, Lancaster CR, Schwalbe H. The human Cdc37-Hsp90 complex studied by heteronuclear NMR spectroscopy. J Biol Chem. 2009; 284: 3885–3896. https://doi.org/10.1074/jbc.M807120200 PMID: 19073599

145. Jiang Y, Bernard D, Yu Y, Xie Y, Zhang T, Li Y, et al. Split Renilla luciferase protein fragment-assisted complementation (SRL-PFAC) to characterize Hsp90-Cdc37 complex and identify critical residues in protein/protein interactions. J Biol Chem. 2010; 285: 21023–21036. https://doi.org/10.1074/jbc.M110.103390 PMID: 20413594

146. Jacobs DJ, Rader AJ, Kuhn LA, Thorpe MF. Protein flexibility predictions using graph theory. Proteins. 2001; 44: 150–165. PMID: 11391777

147. Jacobs DJ, Hendrickson B. An algorithm for two-dimensional rigidity percolation: The pebble game. J Comput Phys. 1997; 137: 346–365.

148. Chubynsky MV, Thorpe MF. Algorithms for three-dimensional rigidity analysis and a first-order percolation transition. Phys Rev E. 2007; 76: 041135.

149. Rader AJ, Hesperheide BM, Kuhn LA, Thorpe MF. Protein unfolding: rigidity lost. Proc Natl Acad Sci U S A. 2002; 99:3540–3545. https://doi.org/10.1073/pnas.062492699 PMID: 11891336

150. Hesperheide BM, Rader AJ, Thorpe MF, Kuhn LA. Identifying protein folding cores from the evolution of flexible regions during unfolding. J Mol Graph Model. 2002; 21: 195–207. PMID: 12463638
151. Krüger DM, Rathi PC, Pfleger C, Gohlke H. CNA web server: rigidity theory-based thermal unfolding simulations of proteins for linking structure, (thermo-)stability, and function. Nucleic Acids Res. 2013; 41: W340–W348. https://doi.org/10.1093/nar/gkt292 PMID: 23609541

152. Pfleger C, Rathi PC, Klein DL, Radestock S, Gohlke H. Constraint Network Analysis (CNA): a Python software package for efficiently linking biomacromolecular structure, flexibility, (thermo-)stability, and function. J Chem Inf Model. 2013; 53: 1007–1015. https://doi.org/10.1021/ci400044m PMID: 23517239

153. Foster SA, Whalen DM, Özen A, Wongchenko MJ, Yin J, Yen I, et al. Activation Mechanism of Oncogenic Deletion Mutations in BRAF, EGFR, and HER2. Cancer Cell. 2016; 29: 477–493. https://doi.org/10.1016/j.ccell.2016.02.010 PMID: 26996308

154. Berman HM, Westbrook J, Feng Z, Gilliland G, Bhat TN, Weisig H, et al. The Protein Data Bank. Nucleic Acids Res. 2000; 28: 235–242. PMID: 10592235

155. Stetz G, Verkhivker GM. Dancing through life: Molecular dynamics simulations and network-centric modeling of allosteric mechanisms in Hsp70 and Hsp110 chaperone proteins. PLoS One. 2015; 10: e0143752. https://doi.org/10.1371/journal.pone.0143752 PMID: 26619280

156. Hekkelman ML, Te Beek TA, Pettifer SR, Thorne D, Attwood TK, Vriend G. WIWS: A protein structure bioinformatics web service collection. Nucleic Acids Res. 2010; 38: W719–W723. https://doi.org/10.1093/nar/gkq453 PMID: 20501602

157. Fernandez-Fuentes N, Zhai J, Fiser A. ArchPRED: a template based loop structure prediction server. Nucleic Acids Res. 2006; 34: W173–W176. https://doi.org/10.1093/nar/gkl113 PMID: 16844985

158. Anandakrishnan R, Aguilar B, Onufriev AV. H++ 3.0: Automating pK prediction and the preparation of biomolecular structures for atomistic molecular modeling and simulations. Nucleic Acids Res. 2012; 40: W537–W541. https://doi.org/10.1093/nar/gks375 PMID: 22570416

159. Phillips JC, Braun R, Wang W, Gumbart J, Tajkhorshid E, Villa E, et al. Scalable molecular dynamics with NAMD. J Comput Chem. 2005; 26: 1781–1802. https://doi.org/10.1002/jcc.20289 PMID: 16222654

160. Dehouck Y, Gilis D, Rooman M. A new generation of statistical potentials for proteins. Biophys J. 2006; 90: 4010–4017. https://doi.org/10.1529/biophysj.105.079434 PMID: 16533849

161. Dehouck Y, Grosfils A, Folch B, Gilis D, Bogaerts P, Rooman M. Fast and accurate predictions of protein stability changes upon mutations using statistical potentials and neural networks: PoPMuSiC-2.0. Bioinformatics. 2009; 25: 2537–2543. https://doi.org/10.1093/bioinformatics/btp445 PMID: 19654118

162. Jiao X, Sherman BT, Huang da W, Stephens R, Baseler MW, Lane HC, et al. DAVID-WS: a stateful web service to facilitate gene/protein list analysis. Bioinformatics. 2012; 28: 1805–1816. https://doi.org/10.1093/bioinformatics/bts251 PMID: 22543666

163. Dahiyat BI, Gordon DB, Mayo SL. Automated design of the surface positions of protein helices. Protein Sci. 1997; 6: 1333–1337. https://doi.org/10.1002/pro.5560060622 PMID: 9194194