Importance of cycle timing for the function of the molecular chaperone Hsp90

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Hsp90 couples ATP hydrolysis to large conformational changes essential for activation of client proteins. The structural transitions involve dimerization of the N-terminal domains and formation of 'closed states' involving the N-terminal and middle domains. Here, we used Hsp90 mutants that modulate ATPase activity and biological function as probes to address the importance of conformational cycling for Hsp90 activity. We found no correlation between the speed of ATP turnover and the *in vivo* activity of Hsp90: some mutants with almost normal ATPase activity were lethal, and some mutants with lower or undetectable ATPase activity were viable. Our analysis showed that it is crucial for Hsp90 to attain and spend time in certain conformational states: a certain dwell time in open states is required for optimal processing of client proteins, whereas a prolonged population of closed states has negative effects. Thus, the timing of conformational transitions is crucial for Hsp90 function and not cycle speed.

Hsp90 is an ATP-dependent molecular chaperone that controls a large set of client proteins¹⁻⁶. Approximately 20% of the proteome appears to directly or indirectly depend on Hsp90 (ref. 7). Hsp90 is a homodimer in which each monomer consists of an N-terminal domain (N domain, which contains the nucleotide-binding pocket⁸) connected via a long charged linker to the middle domain (M domain). The C-terminal domain is responsible for dimerization⁹, which is a prerequisite for efficient ATPase activity^{10–12}. The last five amino acids of Hsp90 form the MEEVD motif, which serves as the binding site for a large class of cochaperones containing a peptide-binding TPR domain¹³.

ATP binding and hydrolysis, which are important for the function of Hsp90 in vivo¹⁴⁻¹⁶, are coupled to a slow conformational cycle in which the Hsp90 dimer progresses from an N-terminally open state to a closed state. The open state is V-shaped and involves dimerization via the C domains and no interaction of the N domains 17. In the crystal structure of Hsp90 in the presence of the nonhydrolyzable ATP analog AMP-PNP, Hsp90 adopts a compact structure in which the N domains are also associated 18. The conformational changes from the open to the N-terminally closed state are rate limiting and precede ATP hydrolysis^{19,20}. A fluorescence resonance energy transfer (FRET)-based kinetic analysis of the conformational cycle has revealed that Hsp90 adopts specific intermediate states 19 before it finally reaches a fully closed state in which ATP hydrolysis occurs. Specifically, after ATP binding, the so-called ATP lid in the Hsp90 N domain changes its position and exposes a hydrophobic N-domain patch that, together with its counterpart in the second monomer, stabilizes the N-terminally closed state¹⁸. Furthermore, this conformational change releases the contacts with the N-terminal helix, which subsequently binds to the N domain of the second subunit within the dimer. Once the N-terminal association is achieved, the Hsp90 N domains associate with the Hsp90 M domains and form the active ATPase. Some of the conformational states appear to be isoenergetic, and random fluctuations between states are possible^{21,22}. Several cochaperones target specific conformations of Hsp90 and thus reduce structural fluctuations; in addition, they modulate specific properties of Hsp90, such as its ATPase activity. Here, we used a set of yeast Hsp90 N- and M-domain mutants with reported effects on biological activity to gain insight into the importance of the timing of conformational transitions for Hsp90 functionality and client-protein processing. Our combined *in vivo* and *in vitro* assays demonstrated that the time spent in open conformations, together with the ability to proceed to the N-terminally closed state, is important for client processing and the essential functions of Hsp90 in yeast.

RESULTS

N- and M-domain mutations affect Hsp90's ATPase activity

The conformational transition from the open to the closed state is rate limiting in the ATPase cycle of Hsp90. We used N- and M-domain mutations with reported effects on the ATP turnover²³ and biological activity of yeast Hsp90 (refs. 12,14,15,18,24–27) (**Fig. 1a** and **Table 1**) to probe the role of specific conformational rearrangements in Hsp90. These mutants affect binding interfaces in the N and M domains that are linked to distinct functional states of Hsp90. According to their ATPase activities, the mutants fell into three categories (**Fig. 1b** and **Supplementary Table 1**). The first exhibited an increased turnover rate: A107N and Δ 8-Hsp90 showed the strongest effect, and T22I showed a slight increase in ATPase activity. The second class

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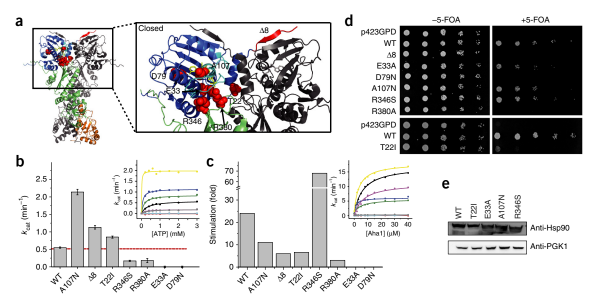


Figure 1 Localization and effects of mutations on ATPase activities. (a) Mutated residues are shown as red spheres in the crystal structure of closed yeast Hsp90 (PDB 2CG9) 18 . Blue, N domain; green, M domain; orange, C domain; cyan, lid structure. (b) Influence of mutations on the ATPase activity of Hsp90. Coupled enzymatic ATPase assay (gray bars). Data are shown as mean \pm s.d. (n = 3 technical replicates). Inset, k_{cat} values of the Hsp90 variants in the presence of various concentrations of ATP: black, WT; yellow, A107N; blue, $\Delta 8$; green, T22I; purple, R346S; gray, R380A; red, E33A; cyan, D79N. (c) Stimulation of Hsp90 ATPase by the cochaperone Aha1, determined by a coupled enzymatic ATPase assay. Fold increases in ATPase activity in the presence of Aha1 are shown. Inset, k_{cat} values of the Hsp90 mutants in the presence of various Aha1 concentrations: black, WT; yellow, A107N; blue, $\Delta 8$; green, T22I; purple, R346S; gray, R380A; red, E33A; cyan, D79N. (d) Viability of the Hsp90 variants deduced by a 5-fluoroorotic acid (5-F0A) shuffling assay. Shuffling experiments were carried out at least three times and represent technical replicates started from independent single yeast colonies. (e) Western blot analysis showing expression of Hsp90 in yeast cells after 5-F0A shuffling. PGK1, loading control.

included mutants with decreased ATPase activity, such as R346S and R380A. The third class included ATPase-deficient mutants, such as E33A and D79N. For most mutants, the $K_{\rm m}$ values for ATP binding were comparable to those of the wild type (WT) (**Supplementary Table 1**). Interestingly, A107N and Δ 8-Hsp90 exhibited a higher affinity for ATP, thus suggesting changes in the ATP-binding pocket.

The altered ATPase activities indicated that the conformational cycle was affected. We used the cochaperone Aha1 as a conformational sensor for structural alterations. Aha1 accelerates the cycle by promoting the formation of the unfavorable closed conformation 19,28,29 . All variants, except E33A and D79N, were stimulated by Aha1, albeit to a different extent (**Fig. 1c** and **Supplementary Table 1**). Interestingly, the R346S variant with inherently low ATPase activity exhibited a WT-like $k_{\rm cat}$ value, thus demonstrating that conformational transitions preceding the formation of closed states are compromised in the absence of Aha1. In contrast, R380A, which exhibited an ATPase activity as slow as that of R346S, was stimulated only slightly by Aha1, thus suggesting that different conformational steps are affected.

Yeast viability does not require Hsp90 ATP hydrolysis

To investigate the biological consequences of the alterations in the conformational cycle of Hsp90, we tested the variants for their ability to support viability in a yeast strain in which both Hsp90 isoforms were deleted, and Hsp90 was provided in a plasmid²⁴. Control experiments confirmed that after loss of the WT-URA plasmid, the strain could not grow on medium lacking uracil (**Supplementary Fig. 1a**). Expression of the variants T22I, E33A, A107N and R346S supported viability similar to that of the WT protein (**Fig. 1d,e**). The expression of the D79N and R380A variants caused lethality, as previously reported^{14,15,27}. Surprisingly, in contrast to the results of previous studies^{14–16}, we found that yeast cells expressing E33A as the sole source of Hsp90 were able to grow, although we detected no ATPase

activity *in vitro*. To further investigate this effect, we tested the ability of the Hsp90 E33A mutant under control of the endogenous promoter to support growth of different yeast strains. We additionally performed tetrad dissections (**Supplementary Fig. 1b,c**). All of these experiments confirmed our initial findings. In contrast, the $\Delta 8$ Hsp90 variant was not able to support yeast viability despite its increased ATPase activity. Finally, although R380A and R346S displayed similar ATPase activities, only R346S supported viability. These results showed that the ability to hydrolyze ATP is not sufficient to confer the essential function of Hsp90.

Hsp90 mutants differ in accumulation of the closed state

To obtain insight into the structural consequences of mutating specific residues, we used small-angle X-ray scattering (SAXS) to determine the shapes of the Hsp90 variants (**Supplementary Fig. 2**). In the absence of nucleotide, WT yeast Hsp90 adopted an open conformation characterized by a broad distribution in the interatomic distance distribution (P(r)) curve (**Fig. 2a** and **Supplementary Table 2**; radius of gyration (R_g) of 64.2 Å; maximum dimension ($D_{\rm max}$) of 240 Å), as previously reported^{30,31}. Most variants tested showed a WT-like shape in the absence of nucleotide. The only exception was the lethal $\Delta 8$ -Hsp90 mutant, which adopted a partly compact state, as indicated by the reduced R_g of 61.9 Å and a redistribution of the P(r) to smaller distances. This mutant appeared to be closed to some extent even in the absence of nucleotide (**Fig. 2a**).

In the presence of ATP- γ S, a slowly hydrolyzed ATP analog, all variants displayed a narrow range of distribution in the P(r) curves, a result indicative of a more compact state (**Fig. 2b**). Δ 8 was further compacted, and the mutant E33A also showed a compact shape. Thus, E33A was able to undergo specific conformational transitions in response to binding ATP- γ S. Compared with the WT protein, the R380A and R346S mutants were less efficient at adopting the closed state.

Table 1 Reported characteristics of analyzed Hsp90 variants

Hsp90 variant	Effects	References
Δ8	Increased ATPase activity; increased N-terminal dimerization in the presence of ATP	23
T22I	Temperature-sensitive mutant of T22, which is covered by the lid in the open state; increased ATPase activity	12,24,25
E33A	Hydrolysis deficient but able to bind ATP; reported lethality in yeast	14,15
D79N	Mutant of D79, which directly contacts the adenine base of the bound ATP; loss of the ability to bind ATP; loss of viability of yeast cells	14,15
A107N	Increased ATPase rate through stabilization of the N-terminally dimerized state	12,18
R346S	Decreased ATPase activity	26
R380A	Mutant of R380, which is located in the catalytic loop of the M domain and orients the γ -phosphate for hydrolysis; decreased ATPase	18,27
	activity in vitro and lethality in yeast	

In the presence of ATP, $\Delta 8$, T22I, A107N and the ATPase-deficient mutant E33A showed a redistribution of distances in the P(r) curves, thus indicating that they adopted a more compact conformation, whereas the WT protein and R346S and R380A variants displayed open conformations (**Fig. 2c**). Thus, the SAXS analysis showed that several of the mutants—particularly $\Delta 8$ and E33A, and to lesser extent T22I and A107N—exhibited scattering properties consistent with the increased presence of closed states, in contrast with the properties of the WT protein.

Structural effects of mutations in the N domain

NMR 1 H, 15 N correlation spectra of the mutations within the N domain (**Supplementary Fig. 3**) indicated mutation-specific conformational changes in the N domain, as evidenced by the chemical shift-changes between WT and mutants (**Fig. 3a,b** and **Supplementary Fig. 4**). The deletion of the first eight amino acids ($\Delta 8$) led to substantial chemical-shift changes for residues in various regions of the N domain (**Fig. 3b**). The other mutant with strongly increased ATPase activity, A107N, exhibited local changes in the lid and in helices $\alpha 1$ and $\alpha 2$. For the variant T22I, the same structural elements as in A107N were affected (**Fig. 3b**). For both variants, various NMR chemical shifts, even those far from the location of the mutated residue, indicated a complex allosteric network of interactions in the N domain. T22 and A107 are located in helix $\alpha 1$ and the lid, respectively, and are involved in N-terminal closure of the full-length Hsp90 dimer (described below).

The two mutations that lacked ATPase activity showed distinct patterns. Whereas the E33A mutation affected all $\alpha\text{-helices},$

D79N had more local effects in the β -sheet and in helix $\alpha 2$, results consistent with direct contacts being made to ATP⁸. The more widespread chemical-shift changes observed for E33A indicated that this residue is important for the overall conformation and dynamics of the N domain.

In the presence of ATP, changes for A107N and T22I were in the vicinity of the nucleotide-binding site (**Fig. 3c**). As expected, E33A but not D79N responded to ATP (**Fig. 3c**). ATP affected the β -sheet, helix $\alpha 2$ and the lid. We observed similar changes in the presence of AMP-PNP (**Supplementary Fig. 5**), thus demonstrating that nucleotide binding is sufficient to induce structural changes in the E33A mutant.

The mutations affect distinct conformational steps

To probe whether the mutations affected the transition to the closed state, we used a FRET system in which the M and N domains in the dimer were labeled with a donor or acceptor dye, respectively¹⁹. After mixing of the components, all Hsp90 variants exchanged subunits and formed FRET complexes (**Supplementary Fig. 6** and **Supplementary Table 3**). Addition of ATP-γS to labeled WT Hsp90 induced N-terminal closing and concomitant increases in FRET efficiency and thus the population of the closed state of Hsp90 (ref. 19) (**Fig. 4a**). All mutants except D79N showed changes in FRET efficiency in response to ATP-γS, thus indicating N-terminal closing (**Fig. 4a** and **Supplementary Table 2**). As compared with the kinetics of the WT, the kinetics was equally fast or was accelerated for mutants with increased ATPase activity (Δ8, T22I and A107N) and was decelerated for variants with

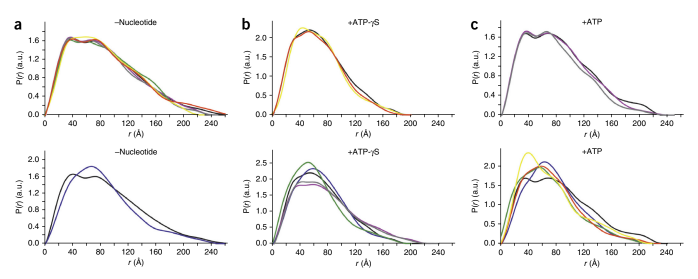


Figure 2 SAXS analysis of Hsp90 variants. (a–c) P(r) curves for WT and mutant Hsp90 in the absence of nucleotide (a) or the presence of 4 mM ATP-γS (b) or ATP (c). Bottom, mutants for which significant differences in the P(r) were observed; top, mutants for which no differences or only minor differences were observed. Black, WT; yellow, A107N; blue, Δ8; green, T22I; purple, R346S; gray, R380A; red, E33A.

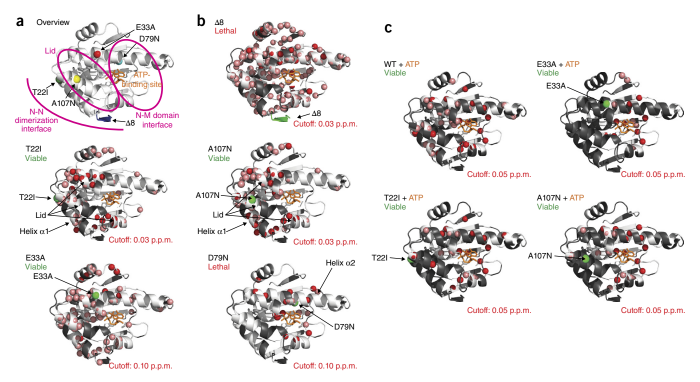


Figure 3 NMR analysis of N-domain mutants. (a) Overview of mutations in the N domain of Hsp90. (b) Amide chemical-shift differences larger than the indicated cutoff value observed in 1 H, 15 N-HSQC spectra of the Hsp90 N-domain mutants compared with the WT protein, as indicated by red spheres on the crystal structure of residues 1–210 of yeast Hsp90 (PDB 1AM1)⁸. Residues with large chemical-shift changes in the mutant could not be unambiguously assigned and are indicated by salmon spheres; residues not assigned in the WT protein are shown in gray. Bound ADP is shown in orange to indicate the nucleotide-binding site. The position of the mutation is indicated by a green sphere. (c) Differences in chemical shifts between the free mutant and the complex with ATP are indicated, as in b. Shifting residues whose endpoints could not be assigned in the complex are indicated by salmon spheres; unassigned residues in the free mutant or overlapping residues, which could not be assigned in the mutant, are in gray.

decreased ATPase rates (R346S and R380A). These findings were in line with the conformational changes being the rate-limiting events during the ATPase cycle^{19,20}.

In the presence of ATP, for WT Hsp90, the closed state is only marginally populated 19 , and thus we observed no change in the FRET signal. This observation was also true for A107N, R346S and D79N. However, for the lethal mutants $\Delta 8$ and R380A and the viable variants T22I and E33A, the acceptor fluorescence increased with the addition of ATP (**Fig. 4b, Supplementary Fig. 6** and **Supplementary Table 3**), thus suggesting that the closed state is more favored in these mutants than in the WT. Interestingly, this effect was observed for mutants with higher as well as lower ATPase activities. Thus, the specific step in the cycle affected by the mutation, not the kinetics of the overall reaction, is important for biological function.

Mutants differ in the accumulation of closed conformations

To gain further insight into the nature of the N-terminally closed state, we performed chase experiments by adding an excess of unlabeled Hsp90 D79N to preformed Hsp90 FRET complexes. Because the disruption of the FRET complexes was strongly influenced by the N-terminal dimerization properties of Hsp90 variants 19 , we were able to test the stability of the closed state (**Fig. 4c**). Without ATP, all variants showed kinetics with comparable half-lives (**Fig. 4c** and **Supplementary Table 4**), thus implying similar C-terminal dimerization. When we used ATP- γ S to induce the N-terminally closed state, all mutants, except D79N, responded with dramatically increased apparent half-lives of the FRET complexes (**Fig. 4c** and **Supplementary Table 4**). In the presence of ATP, the chase with unlabeled Hsp90 resulted in a complex disruption for WT, D79N and R346S, a result

similar to that observed without nucleotide. However, we observed a dramatic increase in the half-lives of the FRET complexes for the lethal variants $\Delta 8$ and R380A, as well as for the ATPase-deficient mutant E33A. The effect was less pronounced for the variants T22I and A107N compared with Hsp90 WT. Thus, several mutants capable of hydrolyzing ATP (R380A, $\Delta 8$, T22I and A107N) displayed accumulation of N-terminally dimerized intermediate states under continuous cycling conditions and ongoing hydrolysis.

To characterize the intermediate states that accumulated, we assessed the binding of the cochaperone Aha1 to the variants in the presence of ATP. Aha1 accelerates conformational changes in Hsp90, thus leading to the closed-1 state¹⁹. As expected, all variants were able to bind Aha1 (**Supplementary Fig. 7a**). When we preincubated the FRET complexes with ATP and Aha1, for all variants except R346S, the chase with unlabeled Hsp90 resulted in a slower complex disruption, as also observed in the absence of Aha1. However, the apparent half-lives of Hsp90 FRET complexes, except for that with D79N, differed from that of the WT in the presence of ATP and Aha1. We observed an additional increase for the variants Δ 8, T22I, R380A and to a lesser extent A107N (**Supplementary Fig. 7b**). Together, these results support the idea that the mutations influenced the conformational changes leading to the closed-1 state.

Mutations affect formation of closed-1 and closed-2 states

We had previously observed two different N-terminally dimerized conformations. In closed-1, the N domains are dimerized, but the N-M interaction is not yet fully formed³². In contrast, the closed-2 state exhibits the N-M interaction³³. From the experiments above, we were unable to deduce which state was populated in the variants.

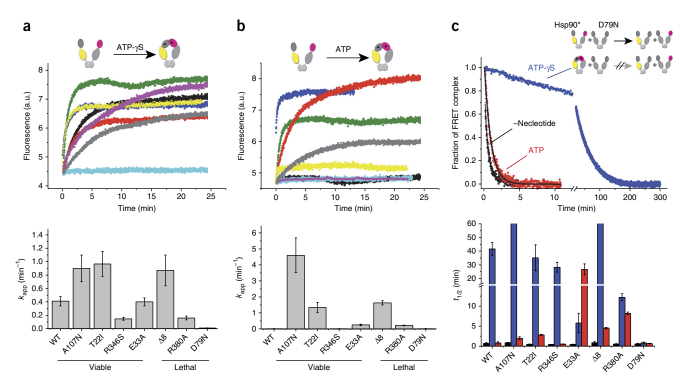


Figure 4 N-terminal closing of the different Hsp90 variants. Monitoring of nucleotide-induced closing kinetics via FRET. ($\bf a,b$) Changes in acceptor fluorescence of the different Hsp90 variants in response to the binding of 2 mM ATP-γS ($\bf a$) or 2 mM ATP ($\bf b$) (top). Black, WT; yellow, A107N; blue, Δ8; green, T22I; purple, R346S; gray, R380A; red, E33A; cyan, D79N. The increase in signal was fitted to a monoexponential function to obtain the apparent rate constants $k_{\rm app}$ (min⁻¹). Data are shown as mean \pm s.d. (n = 3 technical replicates) (bottom). ($\bf c$) N-terminal dimerization stability of the Hsp90 variants, deduced by FRET chase experiments. The chase was induced by addition of an excess of unlabeled Hsp90 D79N variant to 400 nM preformed Hsp90 FRET complexes in the absence of nucleotide (black) or the presence of 2 mM ATP-γS (blue) or 2 mM ATP (red). Top, chase experiments for WT Hsp90. Bottom, apparent half-lives ($t_{1/2}$) derived from a nonlinear fit of the acceptor signal changes. Data are shown as mean \pm s.d. (n = 3 technical replicates).

To differentiate between the closed states, we used the cochaperone p23, whose binding is restricted to the closed-2 state^{18,34}. We performed analytical ultracentrifugation (AUC) to monitor the interaction (**Fig. 5**).

Whereas all viable Hsp90 variants bound p23 in the presence of ATP- γ S, similarly to the WT, the two lethal variants $\Delta 8$ and R380A bound more and less p23, respectively, and D79N did not bind p23 at all (Fig. 5a and Supplementary Fig. 8a). For R380A, this result indicated a destabilization of the closed-2 state; moreover, the lower p23 binding of this mutant also occurred in the presence of ATP, and the closed-1 state appeared to accumulate under cycling conditions (Fig. 5b and Supplementary Fig. 8b). In contrast, the Δ8-variant exhibited an increased p23 binding consistent with a population of the closed-2 state in the presence of ATP (Fig. 5b and Supplementary Fig. 8b). However, without ATP, $\Delta 8$ did not bind to p23 (Supplementary Fig. 8c). Furthermore, in the presence of ATP, p23 also bound to the variants T22I, E33A and A107N (Fig. 5b) and Supplementary Fig. 8b). For E33A, the closed-2 state was the final conformation reached in the presence of nucleotide, because hydrolysis could not occur.

When we preincubated FRET complexes with ATP and p23, the chase with Hsp90 resulted in complex disruption for the D79N and R346S, a result similar to that observed for the WT (**Fig. 5c** and **Supplementary Table 4**). For the variants E33A and Δ 8, we observed an increase in the half-lives of the FRET complexes (**Fig. 5c** and **Supplementary Table 4**). We found a less pronounced effect for A107N, T22I and R380A compared with Hsp90 WT (**Fig. 5c** and **Supplementary Table 4**). However, the closed states of the variants

A107N, T22I and R380A were not further stabilized by p23, because the half-lives were similar for those variants without p23 (**Supplementary Table 4**). These findings suggested that the mutations modulated the population of the closed-1 and closed-2 states.

Population of closed states decreases Hsp90 chaperone function

The mutations affected yeast growth to different extents. Some of the viable variants showed growth defects (**Fig. 1** and **Fig. 6a**), such as T22I which grew slowly, thus confirming its temperature-sensitive phenotype²⁴. We detected normal growth under physiological temperatures and sensitivity under increased temperatures for E33A. Only the R346S and A107N mutants behaved like the WT protein.

To test for specific functions of Hsp90, we induced DNA damage by UV light^{35,36}, which activates the Hsp90-dependent nucleotide excision repair machinery^{35,36}. Yeast cells expressing the T22I or E33A variants displayed increased sensitivity to UV damage, whereas those expressing A107N and R346S behaved similarly to the WT control (**Fig. 6b**).

Furthermore, we determined the ability of the variants to activate the two well-established Hsp90 clients viral Src kinase (v-Src) and glucocorticoid receptor (GR) in yeast. Expression of active v-Src is toxic 37,38 and inhibits yeast growth. The variants T22I and E33A showed reduced toxicity indicating impaired v-Src activation (**Fig. 6c**), whereas the other Hsp90 variants tested (A107N and R346S) exhibited WT-like v-Src maturation. To assess GR activation in yeast, we used a β -galactosidase-based reporter assay 24 . GR processing was slightly reduced in cells expressing the T22I or E33A variants (**Fig. 6d**). For T22I, this result was consistent with published data 24 . A107N

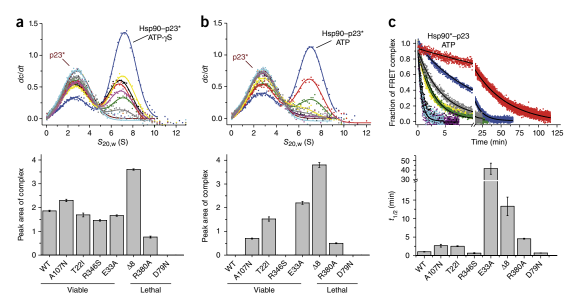


Figure 5 Binding of the cochaperone p23 (Sba1) to the Hsp90 variants. (a,b) Binding of fluorescein-labeled p23 to Hsp90 variants, as analyzed by analytical ultracentrifugation with fluorescence detection in the presence of 2 mM ATP-γS (a) or 2 mM ATP (b). Left, p23* alone (brown; asterisk indicates labeled protein) or p23* in complex with WT (black), A107N (yellow), $\Delta 8$ (blue), T22I (green), R346S (purple), R380A (gray), E33A (red) and D79N (cyan) are shown. Bottom, areas of the complex peaks, obtained by fitting. Error bars, standard error of the fit. $s_{20,w}$, standardized sedimentation coefficient; dc/dt, concentration change over time. (c) N-terminal dimerization stability of the Hsp90 variants in the presence of ATP and p23, obtained by FRET chase experiments. Apparent half-lives ($t_{1/2}$) were derived from a nonlinear fit of the acceptor signal changes. Data are shown as mean \pm s.d. (n = 3 technical replicates).

and R346S supported WT-like maturation of GR. Thus, of the functional mutations, the variants E33A and T22I in particular appeared to broadly affect client protein processing.

To determine the reasons for these observations, we analyzed the interaction with GR by AUC. GR preferentially interacts with ATP-bound Hsp90 (ref. 31). Accordingly, D79N showed the weakest

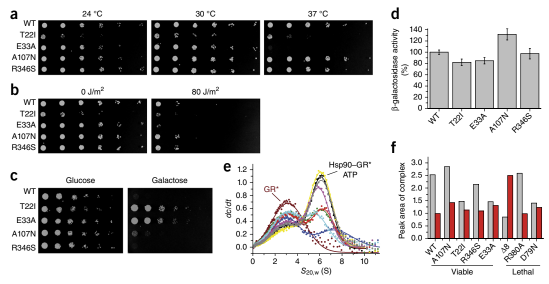


Figure 6 Influence of Hsp90 variants *in vivo*. (a) Temperature sensitivity of yeast cells expressing Hsp90 WT and variants as the sole Hsp90 source at 24 °C, 30 °C or 37 °C. The assay was carried out at least three times, and data represent technical replicates from independent single yeast colonies. (b) Influence of the Hsp90 variants on DNA-repair activity in yeast cells exposed to UV light (80 J/m²) and incubated at 30 °C. As a control, one plate was not treated with UV radiation (0 J/m²). The assay was carried out at least three times, and data represent technical replicates from independent single yeast colonies. (c) Influence of Hsp90 variants on v-Src activation. Yeast cells expressing Hsp90 variants as the sole Hsp90 source and v-Src under the control of a galactose promoter were spotted on glucose (control, no v-Src expression)- and galactose-containing dropout plates (v-Src expression) and incubated at 30 °C. The assay was carried out at least three times, and data represent technical replicates started from independent single yeast colonies. (d) Influence of Hsp90 mutations on GR processing in yeast cells. GR activation, measured with a β-galactosidase-coupled assay, in yeast cells expressing different Hsp90 variants. Experiments were carried out at least three times, and data represent technical replicates (n = 3) started from independent single yeast colonies. Error bars, s.d. (e) Binding of the GR to different Hsp90 mutants, monitored by analytical ultracentrifugation with fluorescence detection and derived from dc/dt plots. GR* alone (brown; asterisk indicates labeled protein) or GR* in complex with WT (black), A107N (yellow), Δ8 (blue), T22I (green), R346S (purple), R380A (gray), E33A (red) and D79N (cyan) are shown. (f) Peak areas of the GR*-Hsp90 complex in the presence (gray bars) or absence of ATP (red bars).

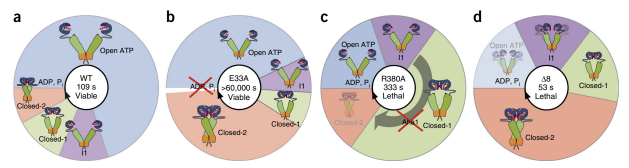


Figure 7 Importance of cycle timing in the function of Hsp90. The scheme illustrates the differences in dwell times in different states (blue, open; violet, I1; green, closed-1; orange, closed-2) for WT Hsp90 (ref. 19) and mutants analyzed in this study. The sizes of the sectors indicate the relative time spent by the mutated protein in the different states during one round of hydrolysis. The average cycle time is indicated by the numbers in the centers of the circles. Mutants showing less intense effects, such as R364A, A107N and T22I, or those that did not bind nucleotide, such as D79N, are not depicted. (a) Hsp90 WT. (b) E33A. The white sector in the scheme for E33A indicates that hydrolysis did not detectably occur, and therefore the mutant does not appear to pass beyond this state. Release of ATP appears to rest the cycle. The variant shifts primarily between the open and the closed-2 state. (c) R380A. The conformational cycle of this mutant is dominated by closed states, especially closed-1. The defect cannot be corrected by Aha1. (d) Δ8. This mutant adopts a compact state even in the absence of nucleotide. It hydrolyzes ATP efficiently but populates primarily the closed-2 state.

affinity for GR, as expected from its deficiency in binding nucleotide (**Fig. 6e,f**). Surprisingly, the $\Delta 8$ variant also exhibited strongly reduced GR binding. Furthermore, for E33A, and to a lesser extent for T22I, GR binding was compromised in the presence of ATP, thus suggesting that mutants for which the closed-2 state accumulates in the presence of ATP show weaker client interactions. Without nucleotide, all mutants interacted similarly with GR (**Fig. 6f** and **Supplementary Fig. 8d**). Thus, the binding site was not affected by the mutations. Only $\Delta 8$ -Hsp90 bound strongly to the client protein, thereby indicating that the GR-interacting state had already accumulated.

DISCUSSION

In this study, we used a set of Hsp90 mutants exhibiting changes in ATP hydrolysis, cochaperone interaction, biological function and client binding $^{26,39-41}$ to probe the conformational cycle of Hsp90. These mutants allowed us to elucidate the mechanism of this molecular machine, which is characterized by large and slow conformational transitions (**Fig. 7a**). In this respect, the association of the N-terminal domains to form the closed-1 state and the subsequent interaction of the N and M domains to form the closed-2 state is important, because these processes are the rate-limiting steps of the ATPase cycle. The mutations are located in the N-terminal loop (which closes over the ATP-binding site), the N-terminal helix α 1 and the N-M interface (which includes the catalytic loop (Pro375–Ile388)). Helix α 1 is part of the N-terminal dimerization interface¹⁸. In the open state, helix α 1 and the lid are in spatial proximity⁸. Closure of the lid and N-terminal dimerization are both coupled to ATP binding 12.

Analysis of Hsp90 mutants has suggested that for viability of yeast cells, ATP hydrolysis is essential but that a range of ATPase activities of Hsp90 is tolerated 26,40,42 . Our study showed that, in agreement with previous reports, ATP hydrolysis is not required for viability, and even mutants with high ATPase rates 23 can be lethal 42,43 . The most striking examples were $\Delta 8$ and E33A: $\Delta 8$ Hsp90 exhibited a two-fold increase in ATP turnover rate relative to that of the WT protein and was therefore expected to be a good substitute for WT Hsp90. However, this mutant was completely defective *in vivo*. In contrast, E33A bound ATP but did not hydrolyze it and therefore was expected to be catalytically dead. However, we found that yeast cells expressing only E33A Hsp90 were viable. These results cannot be explained on the basis of the current understanding of the Hsp90 cycle. Our mechanistic analysis revealed that the timing of the steps

of the Hsp90 cycle is important. In this context, FRET experiments, SAXS and NMR analysis together with the effect of conformation-sensitive cochaperones provided direct insight into the presence of specific conformational states.

 $\Delta 8$ Hsp90 was especially interesting in this context, because this mutant did not populate the open conformation as effectively as the WT protein. Instead, it exhibited a compact, N-terminally associated form. Thus, even in the absence of nucleotides, $\Delta 8$ appeared to be in the closed-1 state, whereas the addition of ATP induced the closed-2 state. This model is consistent with the differences between the mutant $\Delta 8$ and the WT protein seen in NMR chemical shifts that spread over the entire N-terminal domain in the absence of ATP, thus suggesting a different conformation that allows N-terminal dimerization (**Fig. 3b**).

In contrast, the E33A variant unexpectedly did not hydrolyze ATP at detectable rates but supported growth. Our analysis showed that this mutant still responded to ATP binding with defined conformational changes. Ultimately, ATP binding led to the formation of the closed-2 state, because p23 bound *in vitro* and *in vivo*³⁹. The D79N variant, in contrast, was not able to bind ATP (**Supplementary Fig. 4**), could not undergo conformational changes and therefore was not able to support viability.

These findings suggest that the speed of the cycle plays a subordinate role in the function of Hsp90. Instead the conformations that are adopted along the way and, importantly, the time spent by Hsp90 in a specific conformational state are critical. Thus, the cycling between conformational states has tremendous consequences for the biological effects of Hsp90 (**Fig. 7a**).

Together, our results revealed several characteristics of the conformational cycle: (i) Dramatic changes in ATP hydrolysis are tolerated if they allow conformational changes to occur, as observed for E33A and T22I (**Fig. 7b**). These mutations apparently do not affect functionality to an extent that precludes viability, but they render yeast sensitive to stress. (ii) The deceleration of the transition between the closed-1 and closed-2 states, as is the case of R380A, and the accumulation of the closed-1 state under cycling conditions, affects Hsp90 function (**Fig. 7c**). (iii) Mutations that alter initial conformational changes of the chaperone can still support growth (A107N and R346S). (iv) The accumulation of closed states at the expense of open states is deleterious, and the respective mutants are dysfunctional even if they show ATPase activity, as is the case for Δ8 (**Fig. 7d**).

We found that mutations affecting the timing of the conformational transitions strongly impaired the biological function of Hsp90 regardless of their position. In this context, several mutants capable of hydrolyzing ATP (R380A, Δ8, T22I and A107N) exhibited accumulation of N-terminally dimerized intermediate states under continuous cycling conditions and ongoing hydrolysis. This finding was in line with the results of our SAXS experiments, in which $\Delta 8$, E33A, T22I and A107N, in contrast to WT Hsp90, tended to populate closed states in the presence of ATP. These results were consistent with NMR data for T22I and A107N, which showed similar chemical shift-changes that affected the N-terminal dimerization interface of Hsp90. The FRET experiments and cochaperone binding further demonstrated that these mutants exhibited accumulation of the closed-2 state, in contrast to the normal predominance of the open conformation in the presence of ATP 19,22 . E33A, T22I and $\Delta 8$ bound less GR in the presence of ATP, thus suggesting that this conformation is not competent for efficient client binding.

Cochaperones can be used as conformational sensors to pinpoint defects in cycling. In the case of Aha1, we observed effects exclusively for mutants with defects in the transition from the open to the N-terminally closed state; these results were consistent with the function of Aha1 in promoting N-terminal closure²⁸. Comparison of the mutants R380A and R346S provided further clues about important features of the conformational cycle. R380 is located in the so-called catalytic loop of the M domain, a structural element that contacts the ATP-binding site in the N domain, and R346 is part of the N-M interface formed after nucleotide binding. Both mutants led to an almost identical decrease in ATPase activity, to approximately one-third that of the WT. If ATP turnover were the critical determinant of Hsp90 function, then these mutants should behave similarly. However, the mutants showed opposite in vivo effects. Whereas the R380A substitution was lethal, R346S supported viability. What is the critical difference between the two mutants? Strikingly, Aha1 doubled the ATP hydrolysis rate of only R380A. In contrast, R346S was stimulated almost 80-fold, reaching a maximum velocity similar to that of the WT protein. Thus, R346S had difficulties in the transition from open to closed-1. In contrast, for R380A, this step was unaffected, but the subsequent formation of the closed-2 state was compromised and consequently led to the accumulation of the closed-1 state. Thus, although both N-M-interface mutants exhibited a slowed closing reaction, the precise point at which the defect occurred played a decisive role in the reaction.

The picture that emerges is that Hsp90 must populate the open and closed-1 conformations for a certain time for efficient client binding. This model is in accordance with results obtained for the interaction of the GR ligand-binding domain (LBD) with Hsp90, in which the use of nonhydrolyzable nucleotides has been found to reduce GR binding³¹. Thus, even if ATP turnover is possible, the functionality of Hsp90 can be compromised to an extent that excludes *in vivo* function, depending on how much time Hsp90 spends in certain conformations under cycling conditions. It is conceivable that some post-translational modifications may regulate Hsp90 by affecting this balance^{44,45}. In conclusion, the timing of the cycle, i.e., the time that Hsp90 spends in a certain conformation, determines the interaction with client proteins and thus Hsp90's *in vivo* function, regardless of the overall cycle speed.

METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

B.K.Z. and F.T. performed experiments and data analysis. M.R. recorded and analyzed NMR data. T.M. performed and analyzed SAXS measurements. F.H.S. generated Hsp90-knockout strains and performed yeast tetrad analysis. K.R. performed the AUC experiments and analyzed data, J.B. designed experiments. B.K.Z., K.R., M.S. and J.B. wrote the manuscript. D.A.R. provided ATTO488-labeled GR and performed AUC runs.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Cloning and protein purification. All Hsp90 point mutants were generated with QuikChange (Stratagene) site-directed mutagenesis with pET28a containing WT yeast Hsp82 as the template vector. The proteins were expressed in the *Escherichia coli* strain BL21(DE3) RIL (Stratagene) and were purified as described previously⁴⁴ and stored in 40 mM HEPES, pH 7.5, 150 mM KCl, 5 mM MgCl₂ (standard buffer) at -80 °C until usage. Hsp90 cochaperones and the GR LBD mutant were purified as described previously^{31,46}.

Protein labeling. Labeling of the Hsp90 cysteine variants was performed with ATTO-488 maleimide and ATTO-550 maleimide (ATTO-TEC), as previously described¹⁹. p23 was labeled via an engineered cysteine at position 2 with 5-iodacetamidofluorescein (Invitrogen) according to the manufacturer's protocol. Aha1 was labeled with 5(6)-FAM via a reactive succimidyl ester group bound to lysine (Invitrogen), according to the manufacturer's protocol. The GR LBD mutant was randomly labeled with ATTO-488 malemide (ATTO-TEC) on cysteine residues. Free label was removed via Superdex 75 FPLC (Pharmacia Biotech) as previously described³².

ATPase activity. ATPase assays were performed as previously described, with an ATP-regenerating system 47 . Assays were measured with an Hsp90 concentration of 3 $\mu\rm M$ in standard buffer supplemented with 2 mM ATP (Roche) at 30 °C. The Hsp90-specific ATPase activity was inhibited by addition of 50 $\mu\rm M$ of the inhibitor radicicol (Sigma). The remaining activity was subtracted as background. The assays were evaluated with Origin software (OriginLab). The $K_{\rm m}$ values for ATP of each Hsp90 variant were derived from the ATPase activities with following equation:

$$V = V_{\text{max}} \times c/(c + K_{\text{m}})$$

The stimulation of ATPase activity was measured under comparable conditions, except with 1 μM of Hsp90 in the presence of varying concentrations of Aha1 in 40 mM HEPES, pH 7.5, 50 mM KCl and 5 mM MgCl2. The stimulation factor was determined according to the following equation, with V_{max} in the presence of Aha1 and k_{cat} in the absence of Aha1:

 $stimulation(x fold) = V_{max} / k_{cat}$ (without cochaperone)

$$V = V_{\text{max}} \times c/(c + K_{\text{m}})$$

NMR spectroscopy. All spectra were recorded on a Bruker AVIII 600 MHz spectrometer with a cryogenic triple-resonance gradient probe. A construct comprising the first 210 residues of yeast Hsp90 was used to record the $^1H,^{15}N$ -HSQC spectra of the WT protein and the mutants $\Delta 8,\, T22I,\, E33A,\, D79N$ and A107N. Protein concentrations ranged from 110 to 465 μM . In the case of the E33A and D79N mutants, ^{15}N -edited NOESY spectra were recorded to confirm assignments of the mutants. All spectra were acquired in a 20 mM sodium phosphate buffer, pH 6.5, containing 100 mM sodium chloride, 2 mM EDTA, 1 mM DTT and 5% D2O. For the Hsp90 WT N domain and the variants $\Delta 8,\, T22I,\, E33A$ and A107N, spectra with a protein concentration of 300 μM in the presence of an excess of ATP (4 mM) and of 6 mM magnesium sulfate were measured. The mutants E33A and D79N were also measured in complex with a 2:1 excess of AMP-PNP. All spectra were processed with NMRPipe/Draw 48 and analyzed with CcpNmr analysis 49 . The chemical-shift perturbations (CSPs) were calculated according to the following formula, where δ is the proton (H) or nitrogen (N) chemical shift of the WT or mutant (mut):

$$CSP = \sqrt{(\delta_{Hwt} - \delta_{Hmut})^2 + \left(\frac{(\delta_{Nwt} - \delta_{Nmut})}{10}\right)^2}$$

Small-angle X-ray scattering. SAXS data for solutions of the nucleotide-free and ATP- and ATP-γS-bound forms of WT yeast Hsp90 and Hsp90 mutants were recorded on an in-house SAXS instrument (SAXSess mc2, Anton Paar) equipped with a Kratky camera, a sealed X-ray tube source and a two-dimensional Princeton Instruments PI-SCX:4300 (Roper Scientific) CCD detector. The scattering patterns were measured with a 60-min exposure time

(360 frames, each 10 s) for several solute concentrations ranging from 0.8 to 3.3 mg/ml (**Fig. 4**). Radiation damage was excluded on the basis of a comparison of individual frames of the 60-min exposures, wherein no changes were detected. A range of momentum transfer of $0.012 < s < 0.63 \ {\rm \AA}^{-1}$ was covered

$$s = \frac{4\pi \times \sin(\theta)}{\lambda}$$

where 2θ is the scattering angle, and λ is the X-ray wavelength, in this case 1.5 Å.

All SAXS data were analyzed with the ATSAS package (version 2.5). The data were processed with SAXSQuant (version 3.9) and desmeared with GNOM 50 . The forward scattering (I(0)), the radius of gyration, (Rg), the maximum dimension (Dmax) and the interatomic distance distribution function ((P(r)) were computed with GNOM 50 . The masses of the solutes were evaluated by comparison of the forward scattering intensity with that of a human serum albumin reference solution (molecular mass 69 kDa).

Fluorescence measurements. Fluorescence spectra and subunit exchange were carried out as previously described¹⁹.

Nucleotide binding kinetics. To analyze the conformational rearrangements after nucleotide binding, Hsp90 heterodimers (400 nM) were formed by mixture of an equal amount of donor-labeled and acceptor-labeled Hsp90 in standard buffer. The experiment was started by addition of 2 mM nucleotide (ATP, ATP- γ S, all from Roche), and the increase in fluorescence intensity was recorded with a Fluoromax 3 or Fluoromax 2 fluorescence spectrophotometer (Horiba Jobin Yvon) at 575 nm after excitement at 490 nm at 30 °C. The apparent rate constants of the conformational changes were determined by fitting the data to a monoexponential function with Origin software (OriginLab).

FRET chase experiments. After heterodimer formation, the subunit exchange was recorded by addition of a ten-fold excess of unlabeled Hsp90 D79N, and the decay of fluorescence intensity was recorded with a Fluoromax 3 or Fluoromax 2 fluorescence spectrophotometer (Horiba Jobin Yvon) at 575 nm after excitement at 490 nm at 30 °C. For experiments in the presence of different nucleotides (ATP and ATP- γ S, all from Roche) and/or cochaperones (4 μ M Aha1 or 4 μ M p23), Hsp90 variants were preequilibrated 30 min in the presence of 2 mM of the respective nucleotide to allow the formation of the closed state. The apparent half-life of the reaction was determined by fitting the data with the function for exponential decay with Origin software (OriginLab).

Analytical ultracentrifugation. To analyze the binding of fluorescein-Sba1, 5(6)-FAM-Aha1 and ATTO 488-GR to Hsp90, AUC was performed in a Beckman ProteomeLab XL-A (Beckman) equipped with a fluorescence detection system (Aviv Biomedica). Sedimentation-velocity experiments were performed with labeled protein supplemented with various combinations of unlabeled proteins in standard buffer at 42,000 r.p.m. A Ti-50 rotor (Beckman) was used at 20 °C. To determine the sizes of complexes, the raw data were converted to *dc/dt* profiles by subtraction of nearby scans and conversion of the difference into *dc/dt* plots, as previously described⁵¹. The plots generally correlated with those from the SEDVIEW *dc/dt* program⁵². *dc/dt* profiles were analyzed to determine the *s* values and the areas of the corresponding peaks.

In vivo function of Hsp90 variants. The *in vivo* function of the different Hsp90 variants was tested with a plasmid-shuffling approach based on a system described previously²⁴. We used the ΔPCLDα *Saccharomyces cerevisiae* strain (a derivative of W303) obtained from S. Lindquist's laboratory²⁴. This yeast strain is deficient in genomic hsp82 and hsc82 and contains a plasmid encoding hsp82 to rescue lethality. This plasmid, pKAT6, carries a URA selection marker, which enables selection for cells that had lost the WT Hsp82 plasmid in the medium supplemented with 5-FOA. Hsp90 WT and the mutant variants were constitutively expressed from a 2ì high-copy-number plasmid under the control of a constitutive glyceraldehyde-3-phosphate dehydrogenase gene (GPD) promoter (p423GPD vector). The cells surviving the shuffling were tested for loss of the URA plasmid by plating on selection medium lacking URA. To confirm the growth support by the Hsp90 E33A variant, several yeast shuffling strains were used ^{14,15,24,39}.

Tetrad dissection analysis. A diploid strain (*Mat a/o; his3*Δ1/*his3*Δ1; leu2Δ0/ leu2Δ0; lys2Δ0/LYS2; MET15/met15Δ0; ura3Δ0/ura3Δ0; HSP82::kanMX4/HSP82; HSC82::kanMX4/HSC82); was constructed by mating of BY4741 HSC82::kanMX4 and BY4742 HSP82::kanMX4 and was transformed with p415-GPD-HSP82^{E33A} or the empty vector as control. After sporulation, tetrad dissection was performed, and cells were grown on leucine-depleted medium for 3 d. Finally, cells were replica-plated on YPD medium containing 200 μg/ml G418 to test for the distribution of the kanMX4-cassette.

Yeast cell lysis and western blotting. After 5-FOA shuffling, single clones were grown at 30 °C in minimal medium (5 ml) to stationary phase. After an OD_{600} of 5 was reached, cells were harvested (4,500 r.p.m., 5 min). The cell pellets were washed with $\rm H_2O$, and the $\rm OD_{600}$ was measured again. Cells were spun down (4,500 r.p.m., 5 min) and resuspended in 200 μl $\rm H_2O$; then 200 μl of 0.2 M NaOH was added, and cells were incubated for 3 min at RT. Cells were harvested again (13,000 r.p.m., 1 min), and supernatant was discarded. The pellet was resuspended in 100 μl (OD $_{600}$ of 5) sample buffer and boiled for 5 min at 95 °C. Proteins were separated by SDS–PAGE and blotted on a PVDF membrane for 1 h.

The following primary antibodies were used to probe the membrane for 1 h at RT: anti-yeast Hsp90 (Pineda Antibody Service)³³, polyclonal, produced in rabbit (1:40,000 dilution in PBS 0.1% Tween 1% milk) and anti-PGK1 (Invitrogen, cat. no. A6457), monoclonal, produced in mouse (1:20,000 in PBS 0.1% Tween 1% milk). The secondary antibodies were anti-mouse and anti-rabbit IgG-peroxidase antibodies (Sigma-Aldrich, cat. nos. A9044 and A0545, respectively), diluted 1:20,000 in PBS 0.1% Tween 1% milk, were used to detect the respective first antibody for 45 min at RT. For all antibodies, validation is provided on the manufacturers' websites. Detection was performed with Western Bright ECL Spray (Advansta) and ImageQuant LAS4000 (GE Healthcare). Uncropped blots are shown in **Supplementary Data Set 1**.

Growth assays: temperature sensitivity and nucleotide excision repair assay.Drop dilution assays were performed to study the influences of various Hsp90

mutants on the growth of *S. cerevisiae* cells. After 5-FOA, several colonies were plated onto histidine and leucine (–His/–Leu)-depleted medium. After a 2-d incubation time at 30 °C, liquid –His/–Leu medium was inoculated and incubated again (30 °C; 18 h). The OD₆₀₀ of every culture was measured and set to 0.5 by dilution with –His/–Leu medium. The first dilution sample had a total volume of 1 mL. On the basis of this first dilution sample, serial dilution consisting of five more 1:5 dilutions with a sample volume of 200 μ L each was carried out. The serial dilution was finally plated on –His/–Leu medium, with 5 μ L of sample per drop. Dilution steps 1–6 for each mutant were plated in studies of growth at different temperatures (24 °C, 30 °C and 37 °C). To study nucleotide excision repair, the plates were exposed (without the lid of the petri dish) to UV-light radiation (80 × 100 μ J/cm², 10 s) with a CL-100 ultraviolet cross-linker (UVP). As control, one plate was not exposed to UV light. The growth of each Hsp90 mutant was always compared to WT Hsp90, which was present on every plate. Yeast cell growth was documented after 48 h.

v-Src and GR maturation assay. The influence on client maturation was investigated as previously described⁴⁴.

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