Molecular mechanism of thermosensory function of human heat shock transcription factor Hsf1

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Abstract The heat shock response is a universal homeostatic cell autonomous reaction of organisms to cope with adverse environmental conditions. In mammalian cells, this response is mediated by the heat shock transcription factor Hsf1, which is monomeric in unstressed cells and upon activation trimerizes, and binds to promoters of heat shock genes. To understand the basic principle of Hsf1 activation we analyzed temperature-induced alterations in the conformational dynamics of Hsf1 by hydrogen exchange mass spectrometry. We found a temperature-dependent unfolding of Hsf1 in the regulatory region happening concomitant to tighter packing in the trimerization region. The transition to the active DNA binding-competent state occurred highly cooperative and was concentration dependent. Surprisingly, Hsp90, known to inhibit Hsf1 activation, lowered the midpoint temperature of trimerization and reduced cooperativity of the process thus widening the response window. Based on our data we propose a kinetic model of Hsf1 trimerization.

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Introduction To cope with changes in physical and chemical properties of the environment as well as with physiological and pathophysiological conditions which cause protein misfolding, organisms mount a homeostatic transcriptional program, the so-called heat shock response (Jolly and Morimoto, 2000). In all eukaryotic cells, heat shock transcription factor (HSF) 1 is the master regulator of this response and alters transcription of a large number of genes, some of which encode chaperones and proteases (Anckar and Sistonen, 2011). Although this response is essentially cell autonomous, systemic modulation of this response has been observed in metazoa (Morimoto, 2008; Prahlad et al., 2008; Prahlad and Morimoto, 2011).

Metazoan Hsf1 consists of a N-terminal winged helix-turn-helix DNA binding domain (Harrison et al., 1994; Vooister et al., 1994), a hydrophobic shorter heptad repeat regions (HR-A/B) proposed to function as a leucine zipper coiled-coil trimerization domain (Clos et al., 1990; Rabindran et al., 1993), a regulatory domain, a second heptad repeat (HR-C) and a C-terminal transcription activation domain (Figure 1A) (Anckar and Sistonen, 2011; Voellmy, 2004). In unstressed cells metazoan Hsf1 is monomeric and supposed to be in complex with molecular chaperones, including Hsp70, Hsp90 and TRiC/CCT (Shi et al., 1998; Zou et al., 1998; Neef et al., 2014). At physiological concentrations monomeric Hsf1 does not bind appreciably to heat shock elements (nGAAn). In the activated state Hsf1 forms trimers or higher order oligomers and binds to its response elements in heat shock gene promoters (Clos et al., 1990; Rabindran et al., 1993). Currently, two models are discussed for the heat-induced activation of Hsf1: (1) Based on the observation that deletion or mutational alteration of HR-C leads to constitutive trimerization Wu and co-workers proposed that Hsf1 is a thermosensor itself and kept monomeric by intramolecular leucine
eLife digest Cells cope with excessive heat, toxic compounds and other adverse environmental conditions by triggering an internal repair process called the heat shock response. In mammalian cells, a protein called Hsf1 is activated by stress and regulates the activity of a large set of target genes. These genes code for proteins that help the cell cope with the effects of stress, for example, by repairing or breaking down damaged proteins. Under normal conditions, Hsf1 exists as a single molecule, but when it is activated, three molecules come together to make a complex called a trimer that is able to bind to DNA and activate the target genes.

Proteins are made of long chains that then fold into specific three-dimensional shapes. It is not known how Hsf1 is kept in an inactive state in healthy, unstressed cells. One possibility is that the protein folds into a three-dimensional shape that prevents it from being activated. Alternatively, Hsf1 may be bound to other proteins called chaperones that move away when the cell is under stress because they are needed to help the damaged proteins refold into their own three-dimensional shapes.

Hentze et al. used a variety of biochemical techniques to study the human Hsf1 protein. The experiments showed that there are two regions of the Hsf1 protein that changed shape dramatically when the temperature increased. A region that regulates the activity of Hsf1 unfolded, while a region involved in making the trimer became more stable. Detailed analysis showed that once the regulatory region unfolded, the protein was able to interact with other Hsf1 units to make the trimer. Therefore, Hsf1 can directly sense and respond to changes in temperature without the aid of any chaperone proteins.

Further experiments showed that the formation of Hsf1 trimers and the ability of these trimers to bind to DNA depend upon both the temperature and the amount of Hsf1 present. In addition, a chaperone protein called Hsp90 – which is known to be able to interact with Hsf1 – influenced how Hsf1 responded to changes in temperature. Hentze et al. also present a model for the activation of Hsf1 that allows for flexibility in the response of Hsf1 to changes in temperature. Previous studies have shown that Hsf1 is chemically modified during stress and also while the cell recovers from stressful conditions. Therefore, the next challenge will be to find out how these modifications influence the way in which Hsf1 responds to stress.

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zipper formation (Rabindran et al., 1993). However, activation of human Hsf1 in human and insect cells and Xenopus oocytes occurs at different temperatures, arguing against an Hsf1 intrinsic mechanism of heat shock activation (Baler et al., 1993; Clos et al., 1993). (2) Owing to the fact that the large variety of Hsf1-inducing signals have in common to cause protein misfolding and in analogy to the regulation of the heat shock response in E. coli (Guisbert et al., 2008), chaperones were proposed to prevent Hsf1 activation and to be titrated away from Hsf1 under stress conditions, resulting in heat shock response induction (Morimoto, 1998). Consistent with this hypothesis is the observation that inhibition of Hsp70, Hsp90 or TRiC/CCT or knock-down of their expression leads to the induction of the heat shock response (Powers and Workman, 2007; Powers et al., 2008; Neef et al., 2014; Whitesell et al., 2003; Lee et al., 2013; Abravaya et al., 1992; Zou et al., 1998).

Further regulation of Hsf1 is provided by posttranslational modifications, including phosphorylation, acetylation, sumoylation and oxidation of cysteines to disulfide bridges (Hietakangas et al., 2003; 2006; Sarge et al., 1993; Westerheide et al., 2009; Brunet Simioni et al., 2009; Zhong et al., 1998; Lu et al., 2008). The contribution of these modifications to the primary activating mechanism are still unclear (Budzynski et al., 2015).

To resolve the molecular mechanism of the temperature-induced activation of Hsf1 we analyzed the conformational dynamics of purified monomeric human Hsf1 pretreated at different temperatures using hydrogen-1H/2H-exchange (HX) mass spectrometry (MS). We found temperature-dependent unfolding of HR-C and concomitant stabilization of HR-A/B, demonstrating that isolated Hsf1 acts as temperature sensor. At short incubation times the temperature response curve exhibits high cooperativity with a transition midpoint of 36°C. Using fluorescence anisotropy we demonstrate that
the acquisition of DNA-binding competency depends on temperature and concentration of Hsf1. Phosphomimetic Hsf1 variants corresponding to phosphorylation at two serine residues previously shown to negatively affect Hsf1 activation did not have an increased temperature transition midpoint. Hsp90 known to negatively regulate Hsf1-mediated transcription decreased the slope of the transition. DOI: 10.7554/eLife.11576.003
temperature response curve, thereby lowering the transition midpoint and widening the response window. Our data suggest a kinetic model of Hsf1 trimerization.

Results
Recombinant human Hsf1 was purified out of *E. coli* by affinity chromatography and size-exclusion chromatography, resulting in mostly monomeric species in the final fraction (*Figure 1B and C*). Upon incubation at 42°C, Hsf1 formed trimers and higher-order oligomers, as verified by blue native gel electrophoresis consistent with published data (*Clos et al., 1990*), and acquired DNA-binding competence as shown by electrophoretic mobility shift assays (*Figure 1C and D*). The conformational dynamics of Hsf1 was investigated by HX-MS as described previously (*Rist et al., 2006; Graf et al., 2009*). Monomeric Hsf1 was incubated for 30 s in D2O at 20°C, subsequently mixed with ice-cold, low-pH quench buffer to slow down back exchange, and analyzed on our HPLC-mass spectrometry setup including a column with immobilized pepsin for online digestion. As shown in *Figure 1E*, monomeric Hsf1 is highly dynamic with only few regions exhibiting significant protection from HX, including parts of the DNA binding domain and the trimerization domain (HR-A/B). Out of the C-terminal half of the protein, containing the regulatory region, HR-C and the transactivation domain, only the HR-C region showed significant protection at 20°C consistent with an earlier study showing the C-terminal half of Hsf1 largely unfolded (*Pattaramanon et al., 2007*). *Figure 1F* shows a heat map of the DNA binding domain and the trimerization domain of Hsf1, the only parts for which structural information is available.

**Hsf1 is a thermosensor**

To elucidate temperature-induced changes in conformational dynamics, we pre-incubated monomeric Hsf1 at different temperatures for different time intervals followed by incubation at constant temperature in D2O (*Figure 2A*). As control, we analyzed the pre-treated Hsf1 by blue native polyacrylamide gel electrophoresis (*Wittig et al., 2006*) and observed a temperature-dependent increase in trimeric Hsf1 species (*Figure 2B and Figure 2—figure supplement 1*). The 10 min-pre-incubation of Hsf1 dramatically changed conformational dynamics of two regions in Hsf1 (*Figure 2*): temperature-dependent increase in HX is observed in HR-C, indicating heat-induced unfolding, and a concomitant decrease in HX is observed in HR-A/B, consistent with heat-induced trimerization. Close inspection of the spectra of the peptic fragments exhibiting temperature-induced changes in HX revealed bimodal distributions of the isotope clusters indicative of the coexistence of two populations of molecules with different exchange properties (*Figure 2—figure supplement 2A and C, Figure 2—figure supplement 3*). An equation for two Gaussian curves was fitted to the intensity-versus-m/z plots of the data (*Figure 2D and F, and Figure 2—figure supplement 2B and D, Figure 2—figure supplement 3*) and the equation parameters used to back calculate the contribution of each population to the peak intensities (see *Figure 2—figure supplement 2*). For the HR-A/B region the relative frequency of high exchanging population decreases with pre-incubation at increasing temperatures resulting in a sigmoidal temperature response curve (*Figure 2E*). For the HR-C region the opposite is observed: the frequency of high exchanging species increased with increasing temperatures (*Figure 2G*). These data clearly demonstrate that Hsf1 has intrinsic thermo-sensory properties. The midpoint of transition Tm, the temperature at which 50% of the molecules are in the high exchanging conformation after 10 min, was identical for both regions equal to 36.15 ± 0.14°C.

We also determined the midpoint of transition for 30 min pre-incubation at different temperatures (*Figure 2, Figure 2—figure supplement 3*). Under these prolonged incubation conditions, the transition curves became more shallow, and the midpoint temperature for HR-C unfolding and HR-A protection (trimerization) was not identical anymore but decreased to 32.0 ± 0.4 and 34.7 ± 0.2°C, respectively. These data suggest that the temperature-induced conformational changes are not reversible under our conditions, otherwise the steepness of the curves, which is determined by the unfolding enthalpy, should remain the same as for the 10-min-pre-incubation. To investigate this in more detail we heat-shocked Hsf1 for 10 min at 42°C, then incubated the protein for different time intervals at 20°C, and analyzed the conformational state by HX-MS (*Figure 3*). As control, we incubated Hsf1 without prior heat shock for 30 min at 20°C before HX-MS analysis. We also tested whether dilution would lead to trimer dissociation in the time scale of our experiments (*Figure 3G*).
Figure 2. Human Hsf1 is a thermosensor. (A) Experimental design: monomeric human Hsf1 was pre-incubated at different temperature as indicated for 10 min or 30 min and then either analyzed by blue native polyacrylamide gel electrophoresis (BN) or diluted 20-fold into D_2O-buffer at 20°C and incubated for 30 s. The reaction was quenched and the samples analyzed by HPLC-MS. (B) Analysis of quaternary structure of Hsf1 after pre-incubation at different temperatures for 10 min as indicated. Hsf1 was detected by immunoblotting with an Hsf1 specific antiserum. M, monomer; D, dimer; T, trimer. (C) Difference plot of deuteron incorporation into monomeric Hsf1 pre-incubated at the indicated temperature minus deuteron incorporation of Hsf1 pre-incubated at 20°C for peptic peptides as indicated. Error bars are SEM of three independent experiments. Cartoons underneath the X-axis indicate the domains of Hsf1 corresponding to the respective peptic peptides and a homology model of the trimerized human HSF1 (kindly provided by A. Bracher [Neudegger et al., 2016]), colored according to HX as indicated. First and last amino acid of the model are indicated. (D, F) Intensity distributions of the isotope clusters...
Both assays clearly demonstrate that HR-C unfolding, HR-A protection and Hsf1 trimerization are not reversible under these conditions. Therefore, we cannot derive the unfolding enthalpy $\Delta H_U$ from our temperature response data but only use the fit to determine the temperature at which 50% of the transition occurred.

### Kinetics of temperature-dependent conformational changes

To resolve the kinetics of the conformational transitions, we pre-incubated monomeric Hsf1 for 10-1000 s at 35, 37, 39 or 42°C before diluting into D$_2$O and incubation for 30 s at 20°C (Figure 4). For the peptic fragments of HR-C (amino acids 378–395) the low-exchanging population decrease with a rate of 0.0028 ± 0.0002, 0.0038 ± 0.0002, 0.011 ± 0.001, and 0.018 ± 0.001 s$^{-1}$ at 35, 37, 39 and 42°C, respectively. For the peptic fragments of HR-A/B (amino acids 159–168 and 169–175) the low-exchanging population increased with slightly lower rates of 0.0018 ± 0.0004, 0.0033 ± 0.0002, 0.0094 ± 0.0009 and 0.016 ± 0.001 s$^{-1}$ (Figure 4G). The Arrhenius plot of the data yielded the activation energy for the temperature transition of 249 ± 47 kJ mol$^{-1}$ (Figure 4H). Taken together, our data demonstrate that Hsf1 is a thermosensor that directly senses increasing temperatures with conformational changes in HR-A/B and HR-C.

### HR-C interacts with HR-A

To uncouple the temperature-induced HR-C unfolding from trimerization, we replaced the hydrophobic heptad repeat residues in HR-A by serine, which should not engage in coiled-coil interactions. HR-A and HR-B are thought to be involved in trimerization, and HR-B had been previously implicated in negative regulation of trimerization, since deletion of HR-B lead to continuous active trimeric HSF1 (Zuo et al., 1994). HX-MS experiments with the mutant protein revealed that HR-A/B and HR-C are constitutively unfolded at all temperatures tested (Figure 5). These results demonstrate that the ability of HR-A to form a coiled-coil is essential for stabilization of HR-A, HR-B and HR-C, suggesting that HR-C also interacts with HR-A.

### Mechanism of Hsf1 activation

In our in vitro experiments, the midpoint of trimerization of Hsf1 was around 36°C, which seemed rather low given a body core temperature of 37°C, and DNA binding activity of Hsf1 in different human cells is rather low below 40°C and strongly increased above 42°C (Abravaya et al., 1991; Mosser et al., 1990; Baler et al., 1993). In contrast, in testis the major increase in DNA binding activity of Hsf1 is already observed at 38°C (Sarge, 1995; Sarge et al., 1995). Therefore, the above described unfolding of HR-C cannot be the sole determinant for the setpoint of Hsf1 trimerization and DNA binding activity. Trimerization as multi-molecular reaction would be intrinsically
Figure 3. Heat-induced trimerization of Hsf1 is not reversible. (A–F) Prolonged incubation at 20°C does not revert the heat shock induced changes in Hsf1 conformation. Hsf1 (5 μM) was incubated for 10 min at 42°C and then shifted to 20°C. Aliquots were diluted at different time points (0, 3, 10, 30, 100 min) for 30 s into D_2O buffer and subsequently analyzed by LC-MS. As control, Hsf1 was not heat-shocked and incubated for 30 min at 20°C before dilution into D_2O. Shown are the intensity-m/z data for the indicated peptides from HR-A region (A, aa 159–168, 578.29^{2+}; C, aa 169–175, 430.74^{2+}) and region HR-C (E, aa 389–395, 765.31^{1+}) with a global fit of an equation for two Gaussian peaks, as in Figure 2. The percentage of high exchanging species was calculated as described in Figure 2—figure supplement 2 (B, D, F). Data for two independent experiments are shown. (G) Dilution of heat-shocked Hsf1 does not lead to trimer dissociation. Hsf1 (5 μM) was heat-shocked at 42°C for 10 min, subsequently diluted as indicated, incubated at room temperature for 15 min, analyzed by blue-native gel electrophoresis, and detected by immune blotting with an human Hsf1 specific antiserum (lanes 1–7). Control samples were kept on ice before dilution and incubation at room temperature (lanes 8–14). M, monomer; T, trimer; HO, higher order oligomers (not detectable anymore upon dilution).

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Figure 4. Kinetics of heat-induced conformational transitions in human Hsf1. (A–D) Intensity distributions of the isotope clusters for peptide 678.32$^{3+}$ (aa 378–395) and 1155.58$^{1+}$ (aa 159–168) of Hsf1 incubated at 35°C (A and B) or 42°C (C and D) for 10 to 1000 s. Curves are fits of the sum of two Gaussian peak functions to the data. (E and F) Change in the fraction of low exchanging species for peptides 378–395 (E) and 159–168 (F) for 35, 37, 39 and 42°C, as indicated. Curves are fits of a single exponential equation to the data. (G) Transition rates determined by fits as in panels E and F for all four peptides (159–168, 169–175, 378–395, 389–395) evaluated. (H) Arrhenius plot of the data shown in G. Linear regression analysis yielded an activation energy of 258 ± 25, 273 ± 26, 239 ± 19, and 225 ± 22 kJ·mol$^{-1}$ for peptides 159–168, 169–175, 378–395, and 389–395, respectively. Error bars are SEM of three independent experiments.

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concentration dependent. In contrast, inhibition of trimerization by coiled-coil interaction of HR-C with HR-A/B, as originally proposed by Wu and colleagues (Rabindran et al., 1993), would be an intramolecular reaction and consequently independent of concentration. Thus, the concentration of Hsf1 could be an important parameter for controlling the setpoint of the temperature response curve. Consistent with this hypothesis we noticed spontaneous trimerization of Hsf1 even at 4°C upon concentrating the protein. To explore this hypothesis more quantitatively, we devised a fluorescence anisotropy assay to determine the fraction of Hsf1 capable of binding DNA after treatment at different temperatures. We treated different concentrations of Hsf1 at 30, 35, 37, 39, and 42°C for 10 min, serially diluted the samples, incubated them with fluorescent labeled double-stranded DNA containing heat shock elements and measured fluorescence anisotropy (Figure 6). None of the formed Hsf1 trimers dissociated upon dilution and incubation at room temperature (Figure 3G). If Hsf1 trimerization were not concentration dependent in the concentration range tested, all data points would fall onto the same titration curve. This was obviously not the case. The quadratic solution of the binding equilibrium, modified for fractional active protein was globally fitted to the data, assuming identical \( K_D \) values for all formed Hsf1 trimers and similar minimum and maximum anisotropy values for no binding and complete binding, respectively. This fit yielded the fraction of DNA binding competent Hsf1 trimers. For 100 nM Hsf1, no activation was observed up to 42°C (Figure 6F). At 300 nM, less than 20% of the theoretical possible Hsf1 trimers had formed at 42°C.
Figure 6. Temperature-induced acquisition of DNA-binding competence of Hsf1 is concentration dependent. (A–E) DNA-binding competence of monomeric Hsf1 after pre-incubation at the indicated temperature as measured by fluorescence anisotropy of 5′-Alexa 488-labeled HSE-DNA (5′ccccTTCGAAAtaTTccccc3′). Monomeric Hsf1 was pre-incubated at 30–42°C at different concentrations (100–5000 nM) as indicated, then twofold dilution series were prepared and labeled DNA added. Plotted is fluorescence anisotropy (relative values) versus theoretical concentration of Hsf1 trimer. Curves represent a global fit of the quadratic solution of the binding equilibrium modified for fractional activity of Hsf1 to all data together resulting in a $K_D$ of 1.10 ± 0.2 nM for fully active trimeric Hsf1 and a fraction of Hsf1 that formed DNA-binding competent trimers as shown in panel (F). Data of one representative experiment is shown. Error bars represent standard error of the mean of four technical replicates. (F) Fraction of Hsf1 that formed DNA-binding competent trimers at the given Hsf1 concentration and temperature as calculated from data in panels A to E. Mean and standard error of the mean of three independent sets of experiments are shown.

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within 10 min. In contrast, at 1 and 5 μM concentrations a substantial fraction of Hsf1 trimers had formed already at lower temperatures. At 42°C the apparent fraction of DNA-binding competent Hsf1 species may decrease due to the formation of high order oligomers as observed by blue native gel (Figure 1C).

Since these data indicated that the temperature transition of Hsf1 is concentration dependent, we wondered whether only trimerization is affected or HR-C unfolding as well. We therefore repeated the HX-MS experiments at lower concentration (2 μM). Interestingly, not only trimerization but also unfolding of HR-C was concentration dependent and the differences between the transition temperatures of HR-A/B and HR-C were not statistically significant (Figure 7B). However, the difference in transition temperature between Hsf1 at 5 μM and at 2 μM was highly statistically significant (p<0.0001). We could not test lower concentrations of Hsf1 due to lacking sensitivity in the mass spectrometric detection of the important peptides.

Taken together, these data clearly demonstrate that HR-C unfolding, trimerization and the fraction of DNA-competent Hsf1 trimers are a function of temperature and concentration.

**Influence of phosphorylation on the thermosensor function**

Human Hsf1 is heavily modified by posttranslational modifications (29 phosphorylation sites, 5 acetylation sites and 1 sumoylation site; www.phosphosite.org). Some of these modifications have been shown to influence Hsf1 activation (Holmberg et al., 2001; Guettouche et al., 2005; Xia et al., 1998; Xia and Voellmy, 1997; Chu et al., 1996; Wang et al., 2006; Soncin et al., 2003; Kim et al., 2005; Hietakangas et al., 2006; Westerheide et al., 2009; Raychaudhuri et al., 2014). Phosphorylation of Ser307 was proposed to negatively regulate the activation of Hsf1, because Hsf1-S307A was constitutively active in vivo (Chu et al., 1996; Xia et al., 1998). In contrast, phosphorylation of the close-by Ser303 had no influence on Hsf1 activation (Xia et al., 1998). Therefore, we constructed phosphomimetic variants of human Hsf1 (S307D and S303D as control) and determined their temperature response curves using HX-MS. The transition temperature for the phosphomimetic variants was slightly but statistically significantly lower than for Hsf1wt, indicating that phosphorylation at these sites does not prevent temperature-induced trimerization and might even aid it at physiological concentrations of Hsf1 (Figure 7B).

**Influence of Hsp90 on Hsf1 activation**

Several lines of evidence suggested that Hsp90 inhibits Hsf1 activation (Zou et al., 1998) and the current model assumes that Hsp90 binds Hsf1 in the monomeric state in unstressed cells (Anckar and Sistonen, 2011). We therefore studied the effect of human Hsp90Δ on the conformational dynamics of Hsf1, in particular the temperature response curve (Figure 8A–C). Surprisingly, in the presence of Hsp90 the midpoint of transition was lower than in its absence and the response curve was less steep, stretching the transition window from ~10°C in the absence of Hsp90 to ~20°C. Interestingly, the midpoint temperature of transition in the presence of Hsp90 was slightly lower for the peptide derived from HR-C than for the HR-A/B peptides (Figure 8G). A significant difference in midpoint temperature for the two regions was never observed in other experiments with 10-min pre-incubation time but with a 30-min-pre-incubation at the different temperatures (compare Figure 8G with Figure 7B).

To verify that the effect of Hsp90 on trimerization is not an artifact of HX-MS methodology we performed electrophoretic mobility shift assays under comparable conditions using a fluorescently labeled DNA probe containing heat shock elements (Figure 8D–F). Within the experimental error, the results of the DNA binding assay were identical to those of the HX-MS experiments (Figure 8G). Taken together, our results demonstrate that in an in vitro assay with purified components, Hsp90 neither inhibits Hsf1 trimerization nor its DNA binding but, on the contrary, lowers transition temperature and widens the activation window.

**Discussion**

In this study we demonstrate that human Hsf1 is a thermosensor. We are the first to show that the HR-C region of Hsf1 unfolds with temperature-dependent rates, resulting in a release of its repressive effect on Hsf1 trimerization and DNA binding. For relatively short heat shocks unfolding of HR-C and trimerization through intermolecular interactions of HR-A/B exhibit the same temperature...
Figure 7. Temperature response curve of Hsf1 is concentration dependent. (A) Fraction of high exchanging species of peptides 159–168 and 389–395 for 5 and 2 μM Hsf1 pre-incubated at the respective temperatures and analyzed by HX-MS as in Figure 2. Data points and fits of the unfolding equilibrium equation of one representative of three independent experiments are shown. (B) Calculated midpoint temperature for wild-type Hsf1 (2 and 5 μM) and two phosphomimetic Hsf1 variants (5 μM). For 5 μM wild-type Hsf1 (10-min and 30-min-incubations) each data point represents the average of the T_m values for the two peptides observed in the respective region, which were not significantly different from each other (HR-A: aa 159–168 and 169–175; HR-C: 389–395 and 378–395 or 380–388). For 2 μM wild-type and for 5 μM mutant proteins no statistically significant differences were observed between the T_m values for the different peptides (159–168, 169–175, 378–395, 389–395) within an experiment, each data point represents the average of the T_m values of all evaluated peptides for an independent experiment. In addition to the data points of three to six independent experiments, the mean and
response curves and follow identical kinetics, suggesting that these are coupled events. For longer heat shocks the cooperativity of the transition is reduced and HR-C unfolding and HR-A/B trimerization seem to uncouple. Most importantly, HR-C unfolding, temperature-induced trimerization and acquisition of DNA-binding competence depends on the concentration of Hsf1. Moreover, Hsp90 significantly modulated the temperature response of Hsf1, reducing midpoint and steepness of the response curve, thus widening the temperature window within which Hsf1 transits from monomer to trimer and low to maximal DNA binding competence. The response curves of Hsf1 in the presence of Hsp90 are similar to the response curves at prolonged incubation time at elevated temperatures. Thus, Hsp90 accelerates the response at intermediate heat shock conditions.

Our data with purified human Hsf1 are consistent with previous in vivo and in vitro work (Rabindran et al., 1993; Zhong et al., 1998; Baler et al., 1993; Zuo et al., 1994; 1995; Sarge et al., 1993; Zou et al., 1998), substantiating the hypothesis that Hsf1 trimerization and DNA binding is controlled by HR-C. Most previous work was performed in cellular systems or complex cell extracts and only few studies used purified components. Purified Drosophila Hsf1 was shown to exist in a trimer-monomer equilibrium that was influenced by temperature and oxidative stress (Zhong et al., 1998). However, human HSF1 does not seem to exist in such an equilibrium, as we did not observe dissociation of trimeric human Hsf1 upon dilution and heat-induced trimerization was irreversible in our hands (Figure 3). These data suggest that there are principle differences between Drosophila and human Hsf1.

For human Hsf1 we observed striking differences between the temperature response curves of 10 and 30-min-incubation at elevated temperatures. How can these differences be explained? Under our conditions temperature-induced Hsf1 transitions were irreversible. Therefore, at low temperatures even rare unfolding fluctuations of HR-C will eventually lead to trimerization, which will not be observed at short incubation times. Two effects could be responsible for the difference in Tm for HR-A/B and HR-C. At low temperature HR-C dissociation from HR-A/B and re-association might be fast as compared to trimerization. In the free state, HR-C could exist in an unfolding-refolding equilibrium, allowing exchange of protons for deuterons. If refolding is slow in comparison to the intrinsic chemical exchange rate, this would be visible as unfolded species, but HR-C would still refold and reassociate with HR-A/B, repressing trimerization. With increasing temperature unfolding rates would increase, shifting the equilibrium to the completely unfolded state, then allowing HR-A/B trimerization. Alternatively, additional temperature-induced conformational changes in HR-A/B are necessary to allow trimerization and these changes are slow as compared to HR-C unfolding at low temperatures.

Phosphorylation of Ser307 was suggested to repress human Hsf1 activation because the Ser307 to Ala replacement caused constitutively active Hsf1 in vivo (Xia et al., 1998). The temperature response curve of the phosphomimetic Hsf1-S307D variant measured by HX-MS showed a slightly reduced midpoint of transition as compared to wild-type Hsf1, suggesting that phosphorylation at this site does not inhibit heat-induced trimerization but might rather favor it and the repressive effect must be at a different level. Our observations are consistent with more recent data on non-phosphorylatable Hsf1 variants which were not constitutively active, casting a doubt on the repressive effect of phosphorylation at this site (Budzynski et al., 2015).

Most surprising was our finding that Hsp90 does not prevent trimerization and DNA binding of Hsf1 but in the contrary reduces midpoint and steepness of the temperature response curve. This observation seems to be at odds with the known repressive function of Hsp90 on the heat shock response (Zou et al., 1998; Whitesell et al., 2003; 2014; Sittler et al., 2001; Ali et al., 1998). This discrepancy may have different reasons. First, Hsp90 also inhibits the heat shock response in yeast, although yeast Hsf1 is constitutively trimeric and bound to DNA, suggesting that Hsp90 could exert its inhibiting function on human Hsf1 after trimerization and DNA binding as well, consistent with observations for human Hsf1 (Duina et al., 1998; Sorger et al., 1987; Guo et al., 2001). Second, in our experiments we only used one isoform of Hsp90, Hsp90β, and did not add any of the some 30
Figure 8. Hsp90 modulates midpoint and steepness of the temperature response curves of human Hsf1. (A) Difference plot of deuteron incorporation of human Hsf1 in the presence of Hsp90β minus deuteron incorporation into Hsf1 at the indicated temperatures. (B and C) Fraction of high-exchanging species of peptic peptides 169–175 (B) and 389–395 (C) of Hsf1 (5 μM) pre-incubated in the absence (blue) and presence of human Hsp90β (15 μM, red) at the indicated temperature before HX at 20°C for 30 s, quenching with low-pH buffer, peptic digestion, and MS analysis. Data points and fits of the unfolding equilibrium equation for three independent experiments are shown. (D and E) Electrophoretic mobility shift assay (EMSA). HSE-DNA binding of monomeric Hsf1 pre-incubated at the indicated temperature in the absence (D) or presence of Hsp90 (E). (F) Quantification of data from panels D and E. Fraction of DNA bound by Hsf1 versus temperature is plotted. Data points and fits of the unfolding equilibrium equation to the data of two independent experiments are shown. (G) Transition midpoints calculated from HX-MS and EMSA data. Data points and fits of the unfolding equilibrium equation to the data of two independent experiments are shown.
co-chaperones known to assist chaperoning by Hsp90. Further experiments with Hsp90α and different combinations of co-chaperones will be necessary to elucidate whether there exist isoform specificity in Hsf1 regulation or whether Hsp90-co-chaperone complexes have a different effect on Hsf1 trimerization than Hsp90β alone. Third, effects of Hsp90 down-regulation or inhibition could also be indirect, especially because interaction of Hsp90 with Hsf1 seems only to be observed after cross-linking (Neef et al., 2014; Zou et al., 1998), suggesting a very transient interaction. Hsp90 chaperones many kinases, and inhibition of Hsp90 leads to inactivation and degradation of these client proteins. Loss of such kinases could reduce inhibitory effects of phosphorylation or phosphorylation-dependent sumoylation of Hsf1 (Soncin et al., 2003; Wang et al., 2006; Hietakangas et al., 2003; 2006).

Originally, it was proposed that HR-C forms a coiled-coil with HR-A or HR-B to prevent trimerization in unstressed HSF1 (Rabindran et al., 1993; Zuo et al., 1994). Together with the observation that Hsf1 spontaneously trimerizes at high concentrations in the absence of a heat shock (Zhong et al., 1998) and our own observations) and with our HX-MS data, showing temperature-induced unfolding of HR-C, the model shown in Figure 9A can be derived. Under non-stress conditions, HSF1 is in a conformational equilibrium between a closed conformation with HR-C interacting with HR-A/B and an open conformation, in which the two heptad repeat regions are dissociated. Since association of HR-C with HR-A/B is an intramolecular interaction, limiting the diffusional freedom of the interaction partners, association rates would be very high due to the apparent high local concentration. In addition, the net charge of HR-A/B and HR-C are +6 and -7, respectively, favoring association of the two regions by electrostatic attraction. Only at high concentrations of HSF1 association of HR-A/B regions of several HSF1 molecules, which are in the open conformation, would be able to compete with the intramolecular reaction, forming the thermodynamically more stable trimers. This would explain the spontaneous Hsf1 trimerization at high concentrations even at low temperatures (Zhong et al., 1998). Temperature-induced unfolding of HR-C in the closed conformation favors HR-C dissociation. Alternatively or in addition, HR-C unfolding in the open conformation prevents coiled-coil interaction with HR-A/B and thus reduces the back-reaction to the closed conformation. As a consequence, trimer association is favored even at lower HSF1 concentrations.

For two reasons we do not consider this model as very likely: First, single helices free in solutions are usually not stable but are in a rapid equilibrium with the unfolded state due to the low energy difference between helix-internal hydrogen bonds and hydrogen bonds with water. In such a state, we would not expect to see much protection in HX-MS experiments, which is in contrast to our observation (Figure 1E). The amount of heat necessary to unfold a single helix seems too small to account for the temperature control, the steepness, the kinetics and the substantial activation energy of 249 kJ mol⁻¹ for the unfolding/trimerization transition, observed in our experiments. Second, HR-C unfolding would be independent of concentration in this model, also inconsistent with our data (Figure 7).

Based on our data we propose a novel ‘dimer activation model’ (Figure 9B). In this model HR-C remains bound to HR-A/B in the unstressed Hsf1 monomer. However, unstressed Hsf1 could transiently dimerize due to the larger size of the HR-A/B region (75 amino acids) as compared to HR-C (42 amino acids). Such dimers would be destabilized by the interaction of HR-C with HR-A/B, resulting in high dissociation rates and a high fraction of Hsf1 monomers in unstressed cells. However, HR-A/B-HR-A/B interaction could also destabilize the interaction of HR-C with HR-A/B. At high Hsf1 concentrations, association of a third Hsf1 monomer to the transient Hsf1 dimer could occur, displacing the then more-weakly bound HR-C and leading to a stable Hsf1 trimer. Heat-induced unfolding of HR-C and possibly also its binding partner within HR-A/B (in our experiments obscured by subsequent trimerization) would lead to HR-C undocking. This would reduce Hsf1 dimer dissociation rates and favor Hsf1 trimerization even at low Hsf1 concentrations.

In this model, HR-C unfolding and trimerization are kinetically coupled processes, at least for short heat shocks (up to 15 min), explaining why they occur at identical rate constants (Figure 4).
Figure 9. Kinetic models of the thermosensor function of Hsf1. (A) Monomer activation model, based on the originally proposed mechanism modified with our HX-MS data. In unstressed cells monomeric Hsf1 is in equilibrium between a closed, HR-C docked to HR-A/B, and open conformation, with HR-C dissociated from HR-A/B. Owing to high local concentration and electrostatic attraction the intramolecular association rate \( k_{on,i} \) of the HR-C–HR-A/B interaction are very high as compared to the dissociation rate \( k_{off,i} \). Since only uncomplexed HR-A/B can trimerize and Hsf1 trimerization therefore depends on the concentration of the open conformation, at low temperatures, trimerization only occurs at high Hsf1 concentrations. Temperature-induced unfolding of HR-C in the docked or undocked state reduces the intramolecular association rates and/or increases the dissociation rate of the intramolecular HR-C–HR-A/B complex, thereby increasing the concentration of Hsf1 in the open conformation and allowing trimerization at low Hsf1 concentrations. (B) Dimer activation model. At low temperatures, HR-C is constitutively docked onto HR-A/B and monomeric Hsf1 transiently dimerizes through the free part of HR-A/B. Such transient dimerization may partially destabilize the HR-C–HR-A/B interaction. At high Hsf1 concentrations a third Hsf1 monomer could interact with a transient Hsf1 dimer to form a thermodynamically stable Hsf1 trimer with completely released HR-C even at low temperatures. Increasing temperatures lead to unfolding of HR-C in the dimeric Hsf1 species leading to stabilization of the Hsf1 dimer and increased probability of trimerization. Hsp90 might modulate the temperature response by stabilizing the dimeric Hsf1 species. (C) Estimation of the concentration dependence of the transition temperature of Hsf1. Data points are all the \( T_m \) values determined for 10 min incubation at elevated temperatures for Hsf1 wild type in the absence (black) or presence (green) of Hsp90 by HX-MS and by anisotropy. Black curve is a fit of the quadratic solution of the law of mass action of the monomer-dimer equilibrium, assuming that the fraction of dimer determines the \( T_m \). This fit results in a \( T_{m_M} \) for the monomer of Hsf1 (extrapolation to 0 nM) of 53°C, the \( T_{m_D} \) for the dimer of 33°C, and a \( K_D \) of the monomer-dimer equilibrium of 330 nM. Due to the sensitivity of the fit to data points at low Hsf1 concentrations, these are only rough estimates. The blue and red dotted lines are simulations using a lower value for \( K_D \) (100 nM, blue) or \( K_D \) (200 nM) and \( T_{m_D} \) (29°C, red) to simulate the effect of Hsp90. (D) Tentative model of the dimeric Hsf1 based on the recent crystal structure of the trimerization domain of C. thermophilum Skn7, which formed tetramers in two different crystal forms (PDB ID SDSY and SDSZ, [Neudegger et al., 2016]). HR-A, HR-B and HR-C were homology modeled on the tetrameric Skn7 using I-TASSER (Roy et al., 2010; Zhang, 2008; Yang and Zhang, 2015; Yang et al., 2015). HR-C was positioned to accommodate interactions with HR-A and HR-B. The homology model is colored according to HX-MS data (Figure 1E). Residues of the heptad repeat involved in the tetramer interface are shown as sticks. DOI: 10.7554/eLife.11576.015
Furthermore, the average energy to unfold a protein which does not contain a co-factor is 1.4 kJ·mol⁻¹ (Privalov and Gill, 1988; Alexander et al., 1992). Dividing the activation energy for HR-C unfolding determined by us, 249 kJ·mol⁻¹, by 1.4 results in 175, suggesting that 175 residues are involved in this unfolding process. This is close to 168, the number of residues corresponding to a dimer of the coiled-coil between HR-C (42 residues) with a similar sized region in HR-A/B.

Our dimer activation model assumes that the proposed Hsf1 dimer has a lower Tₘ (Tₘ,D) than the Hsf1 monomer and that the measured Tₘ depends on the fraction of Hsf1 dimer present in the assay. We therefore plotted all of our Tₘ values derived for wild-type Hsf1 and incubation times of 10 min versus Hsf1 concentration and fitted the quadratic solution of the law of mass action for the monomer-dimer equilibrium to the data (Figure 9C). This fit results in a Kᵦ for the monomer-dimer equilibrium, the Tₘ,M for the monomer and the Tₘ,D for the dimer. However, the derived values are only very rough estimates due to the sensitivity of the curve to values of very low concentrations of Hsf1, which, for technical reasons, we could not determine so far. However, the steepness of the curve demonstrates that already small changes in concentration can dramatically change the transition temperature for Hsf1 activation.

Although there are no concentration determinations for Hsf1 in different tissues available to our knowledge, using the relative quantification data determined by mass spectrometry for 11 different cancer cell lines (Geiger et al., 2012), and assuming that these cells have a total protein concentration of about 150 mg/ml as determined for HEK293 cells (Gillen and Forbush, 1999), results in Hsf1 concentrations between 10 and 130 nM. The resulting Tₘ values would be between 47 and 53°C. However, Hsf1 is not equally distributed throughout the cytosol but shuttles in and out of the nucleus with various stress conditions preventing Hsf1 export out of the nucleus (Vujanac et al., 2005), resulting in a locally increased Hsf1 concentration between 4- and 17-fold (Fujikaka et al., 2006) and a local concentration between 40 and 2210 nM. This is well within the range that would lead to a functional heat shock response according to our model.

Interestingly, the recent crystal structure of the trimerization domain of the Hsf1 homolog Skn7 of Chaetomium thermophilum contains trimers but also tetramers (PDB ID 5D5Z and 5D5Y, [Neudegger et al., 2016]). This tetramer might be a proxy for the dimer of HR-C-HR-A/B coiled-coils. To visualize how such an Hsf1 dimer might look like, we modeled the structure of HR-A, HR-B and HR-C using I-TASSER (Roy et al., 2010; Zhang, 2008; Yang and Zhang, 2015; Yang et al., 2015) and the tetramer structure of the trimerization domain of Skn7 as template (Figure 9D).

Hsp90 could modulate the monomer-trimer transition by stabilizing HR-A-HR-A interactions and/or destabilize HR-A/B-HR-C interactions, resulting in a reduced dimer dissociation rate and an increased rate of trimerization at lower temperatures. Stabilization of the HR-A-HR-A dimer and concomitant destabilization of HR-A/B-HR-C interaction would automatically destabilize HR-C, since single helices are not stable in solution and only stabilized by interaction with other structural elements, leading to a reduced unfolding transition temperature. To distinguish between stabilization of HR-A-HR-A interaction and destabilization of HR-A/B-HR-C interactions we simulated the effect of Hsp90 on the Tₘ by varying the apparent Kᵦ for dimerization and/or the Tₘ,D of the dimer (see Figure 9C, dotted lines). With the current data available only changing the Kᵦ does not reduce the Tₘ sufficiently to fit the measured values. However, small changes in Kᵦ and Tₘ,D would give satisfying results. This would also explain the observation that Hsp90 reduced the Tₘ for HR-C unfolding significantly more than the Tₘ for trimerization. In this respect Hsp90, curiously, had a similar effect as the prolonged incubation at elevated temperatures (compare transition curves in Figure 2E and G and Figure 8B, C, and F). Thus, Hsp90 accelerates temperature-induced changes in conformation of Hsf1. It is not surprising that the chaperone Hsp90 destabilizes the Hsf1 conformation. Chaperones have been shown to locally unfold native proteins (Rodriguez et al., 2008; Sharma et al., 2010; Kirschke et al., 2014) and Hsp90 is believed to destabilize an α-helix in steroid hormone receptors to allow hormone binding.

How could Hsf1 be active at non-heat stress conditions, for example during development (Xiao et al., 1999), and how could it be activated by salicylate, low pH, Ca²⁺ ions, hypoxia, or proteotoxic stress other than heat shock as demonstrated previously (Mosser et al., 1990; Jurivich et al., 1992; Huang et al., 1995; Liu et al., 1996; Zhong et al., 1999; Ahn and Thele, 2003)? According to our model Hsf1 exists in a monomer-dimer equilibrium, and trimerization of Hsf1 with subsequent DNA binding may occur continuously at low levels, promoted by Hsp90, as shown by us, and inhibited by TriC/CCT (Neef et al., 2014). This may ensure basal Hsf1
transcriptional activity under non-stress conditions. At the same time inhibition of Hsf1 activity by Hsp90, Hsp70-mediated attenuation and continuous Hsf1 monomerization would keep heat shock gene transcription at a low level. Any condition that would favor Hsf1 dimerization and thus trimerization or inhibit chaperone-mediated inhibition or attenuation would, as a consequence, increase heat shock gene transcription. All of the conditions mentioned above, including development, have been associated with an imbalance in proteostasis affecting Hsf1 through titrating away chaperones. However, Hsf1 dimerization could also be affected directly by changing its local concentration, as through transport of Hsf1 into the nucleus (Dai et al., 2003) or preventing its export (Vujanac et al., 2005), and by posttranslational modifications, including glutathionylation of the cysteine in HR-A in response to oxidative stress or alkylation agents (Liu et al., 1996), and phosphorylation of Thr142 in HR-A (Soncin et al., 2003), both of which would reduce the positive net charge of HR-A and thus the electrostatic repulsion.

Our model could explain why the temperature setpoint of activation was lower when human Hsf1 was expressed in Drosophila cells or in Xenopus oocytes (Baler et al., 1993; Clos et al., 1993). In transient or stable transfection experiments, usually a strong promoter is used to express the transfected gene. Thus, the concentration of Hsf1 might have been much higher in the transfected cell than is naturally the case in human cells. Similarly, in Xenopus oocyte-injection experiments the amount of injected mRNA determines the final concentration of HSF1 and might have been so high that the resulting Hsf1 concentration might have allowed activation already at 37°C. Finally, our kinetic Hsf1 activation model would allow each cell to adjust its setpoint of activation by changing the concentration of Hsf1 by producing more Hsf1 or by concentrating it in a smaller compartment, for example by transport from the cytoplasm into the nucleus. This would easily explain the different setpoints in testis (Sarge et al., 1995; Sarge, 1995), mouse T-lymphocytes (Gothard et al., 2003) and mouse motor neurons (Batulan et al., 2003). This might be particularly important for cancer cells for which it was shown that Hsf1 is a driver of malignancy (Dai et al., 2007).

Materials and methods

Protein purification

A culture of BL21 Rosetta, freshly transformed with a plasmid encoding the 6xHis-SUMO-Hsf1 wild-type or Hsf1-HR-A-S11 mutant sequence (Hsf1-I130S,V137S,L140S,V144S,M147S,M154S, L158S, M161S,L168S,V172S,L175S), was grown at 37°C to an OD_600 of 0.6 and then shifted to 20°C. Expression was induced by addition of IPTG to a final concentration of 0.1 mM, the culture grown for 2 hr at 20°C and cells were subsequently harvested by centrifugation (4500 × g for 15 min). All following steps need to be carried out at 4°C. Cell pellets were resuspended in lysis buffer (25 mM Hepes pH 7.4, 150 mM NaCl and 10% glycerol, 3 mM β-mercaptoethanol) containing protease inhibitors (10 μg/ml aprotinin, 5 μg/ml leupeptin, 8 μg/ml pepstatin, one cOmplete Protease Inhibitor Cocktail tablet [Roche Diagnostics, Mannheim, Germany]). Cells were disrupted by subjecting the suspension two times to a chilled microfluidizer at a pressure of 1000 bar. The resulting lysate was immediately centrifuged (16000 × g for 45 min) to remove cell debris.

The supernatant fraction containing 6xHis-tagged HSF1 was incubated for 20 min at 4°C with 1 g of Protino Ni^{2+}-IDA resin (Macherey-Nagel, Düren, Germany) in a rotation shaker. The resin was transferred to an empty gravity-flow column and the flow-through was collected. In a first step the resin was washed with 10 column volumes (CV) of wash buffer and 10 CV of high salt buffer (25 mM Hepes pH 7.4, 1 M NaCl, 10% glycerol, 3 mM β-mercaptoethanol) containing protease inhibitors (10 μg/ml aprotinin, 5 μg/ml leupeptin, 8 μg/ml pepstatin, one cOmplete Protease Inhibitor Cocktail tablet [Roche Diagnostics, Mannheim, Germany]). Cells were disrupted by subjecting the suspension two times to a chilled microfluidizer at a pressure of 1000 bar. The resulting lysate was immediately centrifuged (16000 × g for 45 min) to remove cell debris.

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The supernatant fraction containing 6xHis-tagged HSF1 was incubated for 20 min at 4°C with 1 g of ProteoNi^{2+}-IDA resin (Macherey-Nagel, Düren, Germany) in a rotation shaker. The resin was transferred to an empty gravity-flow column and the flow-through was collected. In a first step the resin was washed with 10 column volumes (CV) of wash buffer and 10 CV of high salt buffer (25 mM Hepes pH 7.4, 1 M NaCl, 10% glycerol, 3 mM β-mercaptoethanol). After a final washing step with another 10 CV of wash buffer the protein was eluted by addition of 1.5 CV elution buffer (25 mM Hepes pH 7.4, 1 M NaCl, 10% glycerol, 3 mM β-mercaptoethanol, 250 mM imidazole) to the column. The SUMO-Tag was cleaved off by incubation with Ulp1 SUMO-protease for 2 hr at 4°C.

The cleaved Hsf1 was further separated by size-exclusion on a S200 HiLoad 16/60 column (GE Healthcare Europe, Freiburg, Germany), equilibrated with Hsf1-buffer (25 mM Hepes pH 7.4, 150 mM NaCl, 10% glycerol, 2 mM DTT). The fractions containing monomeric Hsf1 were adjusted to a concentration of 10 μM, flash-frozen in liquid nitrogen and stored at -80°C.
EMSA for trimerization of HSF1

300 nM Hsf1 premixed with 200 nM Cy3–labeled HSE-oligonucleotide were incubated for 10 min either on ice or at 42°C. As a positive control purified trimeric Hsf1 was kept on ice for 10 min. After incubation the samples were kept at room temperature for additional 30 min, mixed with glycerol and loaded onto a pre-chilled 1% agarose gel (TBE) at 4°C. The agarose gel was run for 30 min at 150 V in the cold room. Labeled HSE-DNA was detected on a FUJI LAS-4000 fluorescence imager (Fuji Photo Film, Düsseldorf, Germany).

For Hsp90 experiments, 5 µM Hsf1 was premixed with 2.5 µM Cy3–labeled HSE-oligonucleotide and 20 µM Hsp90β in buffer containing 10 mM ATP/20 mM MgCl2.

Samples were incubated for 10 min at different temperatures (20°C–42°C), diluted 1:6 in Hsf1-buffer and incubated for 30 min at room temperature. An amount of sample containing 850 nM Hsf1, 3.4 µM Hsp90β and 425 nM HSE-DNA was loaded onto a 1% agarose gel and processed as described above.

HDX for temperature response curve

For the determination of the temperature response curve of Hsf1 activation, 5 µM Hsf1 or mutants (S303D or S307D) in Hsf1-buffer (25 mM Hepes pH 7.4, 150 mM NaCl, 10% glycerol, 2 mM DTT) were heat shocked for 10 or 30 min at different temperatures (20°C–42°C). The samples were then diluted 1:20 in D2O buffer and incubated for 30 s at 20°C. Deuterated samples were quenched 1:1 with ice-cold quench buffer (400 mM sodium phosphate pH 2.2), quickly injected into the injection valve and subjected to LC-MS using an Agilent UPLC and a MaXis mass spectrometer (Bruker, Bremen, Germany). For each experiment at least one unexchanged sample and one fully deuterated control was measured. To determine the Hsf1 activation temperature at lower concentrations, Hsf1 was diluted to a concentration of 2 µM before the experiment.

For HX-MS experiments in the presence of Hsp90β, 10 µM Hsf1 were mixed 1:1 with 40 µM human Hsp90β in reaction buffer (25 mM Hepes, 150 mM NaCl, 10% Glycerol, 20 mM MgCl2, 10 mM ATP and 2 mM DTT) and incubated for 10 min at 20°C. Equilibrated samples were then transferred to a thermomixer for a 10 min-incubation at seven different temperatures (20°C–42°C). Dilution in D2O buffer and subsequent steps were performed as described above.

The unexchanged protein sample was diluted 1:20 in H2O buffer and then mixed 1:1 with quench buffer. The fully deuterated sample (protein in Hsf1-buffer containing 6 M guanidine hydrochloride, lyophilised and redissolved in pure D2O at least three times) was treated equally to normal samples.

Data analysis was performed manually (Data Analysis 4.1, Bruker).

HDX for kinetic studies

5 µM HSF1 in Hsf1-buffer were heat-shocked for different amounts of time (10 s, 30 s, 60 s, 100 s, 300 s, 600 s, 1000 s) at four different temperatures (35°C, 37°C, 39°C, 42°C). The samples were then diluted 1:20 in D2O buffer and incubated for 30 s at 20°C. Deuterated samples were quenched 1:1 with ice-cold quench buffer (400 mM sodium phosphate pH 2.2) and quickly injected into the injection valve and subjected to LC-MS. For each experiment at least one unexchanged sample and one fully deuterated control was measured.

Data analysis was performed manually (Data Analysis 4.1, Bruker).

Data evaluation

Evaluation of bimodal isotope peak distribution: the intensity versus m/z plots of the isotope peaks were fitted with an equation for two Gaussian peaks (see Figures 2,4 or Figure 2—figure supplement 3):

\[ I = \frac{A_1}{\sigma \sqrt{2\pi}} e^{-\frac{(m - \mu_1)^2}{2\sigma^2}} + \frac{A_2}{\sigma \sqrt{2\pi}} e^{-\frac{(m - \mu_2)^2}{2\sigma^2}} \]

with \(A_{1/2}\) being the area of the two peaks; \(\mu\), the m/z values; \(\mu_{1/2}\), the means of the Gaussian peaks, representing the centroid of each of the two subpopulations; and \(\sigma\), the standard deviation of the Gaussian peaks, representing the width of the isotope peak distribution (see Figure 2—figure supplement 2B and D for individual Gaussian curves, the sum of which results in the fit curves of Figure 2). For each peptide showing a bimodal distribution all intensity values belonging to one
temperature (Figures 2, 3, 5, 7, 8) or time (Figure 4) series was globally fitted assuming that \( \sigma, \mu_1 \) and \( \mu_2 \) are constant within this series. Independent experiments were treated independently. Then the parameters of the fit results, \( A_{1/2}, \mu_{1/2} \) and \( \sigma \), were used to calculate for each individual isotope peak which part of the intensity belongs to the low exchanging subpopulation and which part belongs to the high exchanging subpopulation (see Figure 2—figure supplement 2A and C panels: calculated intensity values for low [blue] and high [red] exchanging subpopulations for each isotope peak stacked on top of each other for comparison with original spectra). For all isotope peaks the intensities belonging to one subpopulation (low or high) was summed up to calculate the fraction of this subpopulation within the sample.

To calculate the temperature midpoint of the transition we used the thermal unfolding equation:

\[
F = f_0 + (f_{\text{max}} - f_0) \frac{e^{\left(\frac{T}{T_{\text{m}}} - 1\right) / \Delta H}}{1 + e^{\left(\frac{T}{T_{\text{m}}} - 1\right) / \Delta H}}
\]

with \( f_0 \) and \( f_{\text{max}} \) being the fraction of high exchanging subpopulation at low and high temperatures, respectively; \( T \), absolute temperature in K; \( T_{\text{m}} \) temperature at midpoint of activation; \( R \), gas constant; \( \Delta H \), unfolding enthalpy.

**Blue native gel**

Hsf1 (10 \( \mu \)M) in Hsf1-buffer were incubated for 30 min at 0 \(^\circ\)C (control) or 42 \(^\circ\)C (heat shock). Natively purified dimer and trimer of Hsf1 were added as additional controls. After incubation 7 \( \mu \)g of Hsf1 were loaded on a 7% native gel or a 4–16% native gradient gel and separated by blue native polyacrylamide gel electrophoresis as described in (Wittig et al., 2006) except that Coomassie Brilliant Blue G250 was only present in the sample buffer (0.2%) not in the running buffer. For western blot analysis, an anti-Hsf1 antibody was used (Santa Cruz Biotech, HSF1 H-311).

**Fluorescence anisotropy**

Aliquots of Hsf1 were thawed and immediately centrifuged (4 \(^\circ\)C, 15 min, 15000 rpm). In order to capture any occurring trimeric Hsf1, the supernatant was incubated for 20 min on ice with DNA containing three HSEs coupled to magnetic beads (5'-CCCCTTCCCGAATATTCCCCC-3', 0.5 mg per aliquot, Dynabeads M-280 by Invitrogen). The supernatant concentration was determined by absorbance at 280 nm. Subsequently, four discrete concentrations of Hsf1 (5 \( \mu \)M, 1 \( \mu \)M, 300 nM, 100 nM) were prepared with Hsf1-buffer and heat-shocked for 10 min at five different temperatures (30 \(^\circ\)C, 35 \(^\circ\)C, 37 \(^\circ\)C, 39 \(^\circ\)C and 42 \(^\circ\)C) using a temperature-controlled water bath. Additionally, 300 nM Hsf1 was kept on ice for the same period of time as a control. Fluorescence anisotropy measurements were performed with a CLARIOstar microplate reader (BMG Labtech) and 384-well black flat-bottom microplates (Corning) in a final sample volume of 30 \( \mu \)L. Samples were serial diluted 1:2 until concentrations were below 1 nM. 10 nM of Alexa Fluor 488-labelled DNA containing three HSEs (5'-[A488]CCCCTTCCCGAATATTCCCCC-3' (Sigma-Aldrich) was added by the injection system of the plate reader to start the measurement.

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