The quaternary organization and dynamics of the molecular chaperone HSP26 are thermally regulated

J. L. Benesch
*Physical and Theoretical Chemistry Laboratory*

Andrew Aquilina
*University of Wollongong*

Andrew Baldwin
*University of Toronto*

Agata Rekas
*Australian Nuclear Science and Technology Organisation*

Florian Stengel
*Physical and Theoretical Chemistry Laboratory*

Follow this and additional works at: [https://ro.uow.edu.au/scipapers](https://ro.uow.edu.au/scipapers)

**Recommended Citation**

Benesch, J. L.; Aquilina, Andrew; Baldwin, Andrew; Rekas, Agata; Stengel, Florian; Lindner, Robyn; Basha, Eman; Devlin, G.; Horwitz, J.; Vierling, Elizabeth; Carver, John; and Robinson, C. V.: The quaternary organization and dynamics of the molecular chaperone HSP26 are thermally regulated 2010. [https://ro.uow.edu.au/scipapers/5134](https://ro.uow.edu.au/scipapers/5134)
The quaternary organization and dynamics of the molecular chaperone HSP26 are thermally regulated

Abstract
The function of ScHSP26 is thermally controlled: the heat shock that causes the destabilization of target proteins leads to its activation as a molecular chaperone. We investigate the structural and dynamical properties of ScHSP26 oligomers through a combination of multiangle light scattering, fluorescence spectroscopy, NMR spectroscopy, and mass spectrometry. We show that ScHSP26 exists as a heterogeneous oligomeric ensemble at room temperature. At heat-shock temperatures, two shifts in equilibria are observed: toward dissociation and to larger oligomers. We examine the quaternary dynamics of these oligomers by investigating the rate of exchange of subunits between them and find that this not only increases with temperature but proceeds via two separate processes. This is consistent with a conformational change of the oligomers at elevated temperatures which regulates the disassembly rates of this thermally activated protein.

Keywords
CMMB

Disciplines
Life Sciences | Physical Sciences and Mathematics | Social and Behavioral Sciences

Publication Details

Authors
J. L. Benesch, Andrew Aquilina, Andrew Baldwin, Agata Rekas, Florian Stengel, Robyn Lindner, Eman Basha, G. Devlin, J. Horwitz, Elizabeth Vierling, John Carver, and C. V. Robinson

This journal article is available at Research Online: https://ro.uow.edu.au/scipapers/5134
THE QUATERNARY ORGANIZATION AND DYNAMICS OF THE MOLECULAR CHAPERONE HSP26 ARE THERMALLY REGULATED

Justin LP Benesch†, J Andrew Aquilina‡, Andrew J Baldwin§, Agata Rekas¶, Florian Stengel†, Robyn A Lindner¶, Eman Basha‡, Glyn L Devlin¶, Joseph Horwitz§, Elizabeth Vierling¶, John A Carver¶*, & Carol V Robinson†**

† These authors contributed equally

* Correspondence: JAC, Tel +61 8 83033110, Fax: +61 8 83034380, john.carver@adelaide.edu.au
** CVR, Tel: +44 1865 275473, Fax: +44 1865 275410, carol.robinson@chem.ox.ac.uk

Running Title: Quaternary Organization and Dynamics of ScHSP26
HIGHLIGHTS

- Combining biophysical approaches allows characterization of a molecular chaperone
- \textit{Sc}HSP26 populates a heterogeneous ensemble of oligomers
- Upon thermal activation \textit{Sc}HSP26 oligomers dissociate and shift to larger species
- Quaternary dynamics of \textit{Sc}HSP26 are underpinned by two separate processes

GRAPHICAL ABSTRACT
SUMMARY

The function of ScHSP26 is thermally controlled: the heat shock which causes the destabilization of target proteins leads to its activation as a molecular chaperone. We investigate the structural and dynamical properties of ScHSP26 oligomers through a combination of multi-angle light scattering, fluorescence spectroscopy, NMR spectroscopy, and mass spectrometry. We show that ScHSP26 exists as a heterogeneous oligomeric ensemble at room temperature. At heat-shock temperatures two shifts in equilibrium are observed: towards dissociation, and to larger oligomers. We examine the quaternary dynamics of these oligomers by investigating the rate of exchange of subunits between them, and find that this not only increases with temperature but proceeds via two separate processes. This is consistent with a conformational change of the oligomers at elevated temperatures which regulates the disassembly rates of this thermally activated protein.
INTRODUCTION

The small heat-shock proteins (sHSPs) are an almost ubiquitous family of intracellular molecular chaperones, characterized by the presence of a highly conserved ‘α-crystallin’ domain [1-5]. This domain is named after the eponymous vertebrate sHSP which is found primarily in the eye lens [6], and whose chaperone activity [7] is crucial to maintaining protein solubility [8]. α-Crystallin is composed of two closely related subunits, A and B. In mammals, mutations to these proteins have been found to lead to cataract [9], and, with αB-crystallin being systemically expressed [6], a number of devastating skeletal- and cardio-myopathies [10].

The mechanism of action of the sHSPs is distinct from many molecular chaperones in that they do not actively refold target proteins. Instead they bind partially unfolded targets and hold them in a state suitable for subsequent refolding by ATP-dependent chaperones [1-5]. This role of the sHSPs as the ‘paramedics’ of the chaperone ‘medical service’ is particularly apparent for ScHSP26, one of two sHSPs of *Saccharomyces cerevisiae* cytosol [11]. ScHSP26 is activated rapidly by an increase in temperature [12], and stabilizes non-native target proteins [13] until the energy dependent HSP104/Ssa1/Ydj1 refolding machinery is mobilized [14, 15]. Consequently, these target proteins are recovered as folded functional entities [14, 15].

Structurally the sHSPs remain poorly understood because, in the main, members of this protein family are extremely dynamic and heterogenous [2, 9]. What is clear however is that, while the monomers are small, in general the sHSPs associate into large oligomers with dimeric units representing a common building block [1-5]. Presently, the highest resolution structural data for ScHSP26 has been obtained by means of cryo-electron microscopy: though the protein was heterogenous in structure, via the
development of novel image processing strategies [16], two coexisting structures, each comprised of 24 subunits, were obtained [17].

Here we examine ScHSP26 principally by means of mass spectrometry (MS). With technological advances over the last decade enabling the transferral and analysis of intact protein complexes from solution into the gas phase [18], MS has become a bona fide structural biology approach [19]. MS has been employed to determine the stoichiometries of a number of monodisperse sHSPs [20, 21], and to deconvolute the populations of individual oligomers within heterogeneous sHSP ensembles [22-24]. Furthermore, real-time monitoring of the sHSPs has enabled us to describe and measure their dynamic quaternary structure through the process of subunit exchange [25, 26].

While this capacity of sHSPs to exchange subunits has been known for some time [27], the cellular implications, including the role in chaperone action, remain unclear. It has been proposed that this phenomenon occurs to establish a dynamic and plastic chaperone ensemble, able to cater for the wide variety of target proteins destabilized under different stress conditions [28]. Additionally, the rate-determining dissociation of the oligomers [29] may amount to an activation of the sHSPs [5, 30]. However, it has since been shown for αA-crystallin that subunit exchange does not necessarily correlate with chaperone efficiency [31], and for ScHSP26 that oligomer dissociation is not a prerequisite for activation [32]. In the latter case activation has been proposed to be controlled by temperature-induced changes in conformation of the ‘middle domain’, a region of sequence just N-terminal to the α-crystallin domain unique to ScHSP26 [12].

In the present work we investigate the consequences of the transition to heat-shock temperatures on the quaternary structure and dynamics of ScHSP26. We demonstrate that though the most abundant stoichiometry of this protein is a 24mer at room temperature, a range of oligomeric states are
populated. At higher temperatures we notice a shift in equilibrium to favour monomers and dimers, but also that the predominant higher-order oligomeric species is a 40mer. We examine the properties of these large oligomeric forms of ScHSP26 and show that not only are they dynamic, but also that the mechanism of its subunit exchange must involve more than one concurrent process.
RESULTS

**ScHSP26 populates a polydisperse ensemble which exhibits chaperone activity**

To investigate the mass distribution of the ScHSP26 oligomer at room temperature, we first used size-exclusion chromatography (SEC) coupled to a multi-angle light-scattering (MALS) detector (Fig. 1A). The protein eluted from the column as a single broad, but slightly asymmetric, peak centered on approximately 10.5 min. The MALS data shows that at the front end of the peak the average mass was approximately 950 kDa, at the tail end 580 kDa, and at the maximum 810 kDa. This corresponds to an approximate average-oligomer distribution range of 24-40 subunits.

The ability of ScHSP26 to act as a molecular chaperone was assessed via its inhibition of the aggregation and precipitation of reduced insulin B chain. Under reduction stress in this *in vitro* assay at 25 ºC ScHSP26 was an efficient molecular chaperone, with a 1.0:1.0 ratio of ScHSP26:insulin providing complete suppression of aggregation (Fig. 1B). The efficiency of chaperone action is comparable to that observed previously for ScHSP26 [30] and in similar studies of other sHSPs [33].

**ScHSP26 undergoes temperature- and concentration-dependent structural changes leading to C-terminal exposure**

The dependence of the structure of ScHSP26 on temperature was investigated by observing changes in intrinsic tryptophan fluorescence. There are two tryptophan residues in ScHSP26: Trp-72 which lies in the middle domain [12], and Trp-211 which resides in the C-terminal region [34]. Spectra acquired at 20 ºC, 45 ºC and 90 ºC exhibit a substantial red shift with temperature (figure 2A, left). Monitoring the intensity at 330 nm demonstrates a relatively uncooperative transition between 20 ºC and 45 ºC with a midpoint of ~35 ºC (Fig. 2A, right). Upon cooling of the solution back to 20 ºC, almost full recovery of
a native-like spectrum could be observed within 12 hours. Together, these data indicate that the environment of one or both of the tryptophan residues in ScHSP26 is becoming on average more solvent exposed with increasing temperature.

Sequence comparison suggests that the last few amino acids of ScHSP26, including Trp-211, are aligned with the C-terminal extension of the α-crystallins and other mammalian sHSPs (Fig. 2B). This region is typically highly mobile and unstructured [35, 36] and is downstream of the highly conserved C-terminal I/VXt/V motif, which serves to attach dimers together within sHSP structures [2, 3, 5]. Flexible regions in proteins can be observed in NMR experiments, despite belonging to species far beyond the accepted molecular mass limit of the technique, as they tumble essentially independently and hence more rapidly than the bulk oligomers [37]. NMR spectra ($^1$H-$^1$H TOCSY and $^1$H-$^{15}$N HSQC) were acquired at various temperatures and protein concentrations in order to ascertain whether similar C-terminal flexibility is present in ScHSP26 (Fig 2B). At 25 °C and 1.66 mM, cross-peaks are only observed for Asn-213 (Fig 2B I). As the temperature was raised to 45 °C a peak for Gly-212 was observed (Fig 2B II), and when the concentration was lowered to 0.33 mM at 45 °C Trp-211 became visible (Fig 2B III). At 0.17 mM and 45 °C additional cross-peaks were observed at random-coil chemical shift values (Fig 2B IV). Thus, unlike the mammalian sHSPs [37], ScHSP26 lacks a highly mobile C-terminal extension at 25 °C. However, the appearance of additional cross peaks with elevated temperature and reduced concentration demonstrate enhanced C-terminal flexibility of ScHSP26 under these conditions. Our data therefore combine to suggest that changes in intrinsic fluorescence with elevated temperature are due primarily to an alteration in structure which affects the solvent accessibility of Trp-211.
ScHSP26 undergoes quaternary rearrangement at elevated temperature

The temperature-dependent changes observed in our fluorescence and NMR data could feasibly arise from either changes in oligomerization or an unfolding of the protein chains. To address this we examined ScHSP26 organization at a range of temperatures by means of nanoelectrospray MS. While our SEC-MALS data show that at 25 °C under our conditions ScHSP26 exists as a heterogeneous ensemble (Fig 1A), they only provide an average mass of the oligomers in the volume being sampled. In contrast the high resolution afforded by MS renders it well suited to the analysis of complex protein assemblies [38]. Figure 3A shows a mass spectrum of ScHSP26 obtained at 25 °C. A large number of peaks were observed between 9000 and 14000 m/z in a bimodal distribution, with the first broad concentration of signal between approximately 9000 and 11000 m/z, and the second between 11000 and 14000 m/z. The presence of this multitude of peaks suggests the presence of many differently sized oligomers, and is in agreement with the SEC-MALS data. In the low m/z region of the spectrum, some minor peaks were observed corresponding to the presence of a small population of monomers and dimers.

To determine whether these monomers and dimers were in equilibrium with the oligomers we obtained mass spectra at a range of different temperatures using a device which enables thermo-control of the sample solution during MS analysis [39] (Fig. 3B and 3C). As the temperature was raised, an increase in the amount of signal corresponding to monomer and dimer was observed. At 60 °C, the highest temperature investigated, these dissociated species dominated the spectrum. The remaining intact oligomers were observed in the range of 11000-14000 m/z, corresponding to the higher m/z range of the two distributions observed at room temperature (Fig. 3A). These results imply that a significant proportion of the monomers and dimers arose from dissociation of the smaller oligomers, i.e. those which appeared between 9000 and 11000 m/z at room temperature, as the temperature was increased.
Polydisperse ScHSP26 is composed of dimeric building blocks and has variable quaternary organization

Because of the polydispersity of this system, the individual oligomeric species could not be completely identified. In the main this is due to the poor level of desolvation afforded by the ‘gentle’ MS conditions used in the heating experiments. In an attempt to improve desolvation, and consequently obtain narrower peak-widths and improved separation between peaks, the voltages accelerating the ions into the collision cell were increased. The resultant collisional activation results in the removal of adducted buffer and solvent, and hence peaks of higher effective resolution [40]. Figure 4A shows such spectra of ScHSP26 which had been equilibrated at 43°C for 30 mins (lower panel), or had been left at room temperature (upper). The heat-treated protein displays peaks between 11000 and 14000 m/z. Two components, a 40mer and 42mer, could be clearly identified, although other minor peaks remained unassigned. In the unheated protein, peaks were distributed between 9000 and 14000 m/z, and were considerably more numerous than for the heat-treated sample. While some species could be clearly identified, notably a 24mer and 40mer, the considerable peak overlap rendered further assignment problematic.

To facilitate the identification of all constituent species we employed a collision-induced dissociation approach which we have developed in our laboratory for the analysis of polydisperse assemblies [22]. In such an experiment the protein complexes are submitted to numerous high-energy collisions with an inert gas, which results in their dissociation into monomers and stripped oligomers (that is, oligomers having lost one or more monomers). The charge removed by the monomers is disproportionate to their mass, and thereby leads to a dramatic increase in the m/z ratio of the residual stripped oligomers. This dissociation process can occur repeatedly, in a sequential manner, leading to an effective charge
reduction of the oligomers with a consequent increase in separation between peaks [41, 42]. As such, the oligomeric distributions of the heterogeneous ensembles can be readily deconvoluted.

The results of this approach applied to the heat-treated ScHSP26 are illustrated in figure 4B. Activation of the oligomers gave rise to the appearance of four distributions of peaks: one centred on 2000 m/z corresponding to monomers, and others around 17000 m/z, 24000 m/z, and 35000 m/z, corresponding to singly, doubly and triply stripped oligomers respectively. Figure 4C shows a portion of the spectrum from the doubly stripped oligomer region of the spectra for heat treated (lower) and non-heat treated (upper) ScHSP26. The peak separation is such that the different oligomers comprising the heterogeneous ensemble can be identified. Nine distinct peaks were observed in the case of the non-heated protein, corresponding to oligomers containing even numbers of subunits in the range 24 to 40. For the heat-treated protein, the range was narrowed to 32 to 42.

Summing the intensities of the signal corresponding to the different species across the entire doubly stripped oligomer region allowed us to quantify the relative populations of the species which comprise the polydisperse assemblies (Fig. 4D). The oligomeric distribution of unheated ScHSP26 was found to have a range of 24 to 42 subunits, with no species containing an odd number of subunits. Notably, the distribution was non-Gaussian, with the 24mer species being considerably more abundant than the ‘neighbouring’ 26mer. After heat treatment, however, the distribution looked considerably different (Fig. 4D, lower). No species smaller than 32mers were observed, and the distribution was dominated by an oligomer comprised of 40 subunits.

*ScHSP26 exchanges subunits by different pathways*
In attempt to gain insight into the subunit dynamics of ScHSP26 we used $^{15}$N-labeled protein which is 288 Da heavier per subunit than the wild-type ($^{14}$N) protein, yet of identical sequence and structure. Previously, we developed a technique of measuring the rate of subunit exchange for a mixture of two polydisperse proteins, in which the major overlapping peak of the doubly-stripped oligomer region for each protein is monitored [31]. Because the major overlapping peak for a homo-oligomer in this region occurs at an $m/z$ value equal to the mass of the monomer, i.e., where all oligomers carry one charge per subunit [22], it is possible to follow gross changes in the subunit composition of two heterogeneous ensembles over time.

An equimolar mixture of $^{14}$N and $^{15}$N ScHSP26 was prepared, and infused into a thermo controlled nanospray probe set at 34°C (Fig. 5A, upper panel). At the earliest time-point two distinct peaks were observed, one at $\approx$23770 m/z and the other at $\approx$24090 m/z which correspond to the doubly stripped oligomers of all of the polydisperse assemblies of $^{14}$N and $^{15}$N ScHSP26 respectively. Over a period of 95 mins, these peaks were observed to broaden and converge towards a midpoint at $\approx$23940 m/z, consistent with a fully exchanged mixture of ScHSP26. This indicates exchange occurring in a stepwise manner, presumably via the incorporation of dimers. When this experiment was repeated at 39°C (Fig. 5 middle panel), this convergence proceeded considerably more rapidly, and was complete within 30 minutes. A third experiment performed at 45°C (Fig. 5, bottom panel) showed a further increase in the exchange rate, such that the reaction mixture was equilibrated after approximately 12 minutes. In this case the subunit exchange reaction characteristics were quite distinct from those observed at the lower temperature. Notably during the reaction (e.g. after five minutes) the presence of both fully exchanged species as well as the original homogeneous isotope populations were observed.

To establish a more quantitative measure of the relative rates of exchange we developed a model for the subunit exchange of ScHSP26 based on dimers dissociating from the oligomers with a rate constant
of $k_1^-$, and associating with the next oligomer it encounters, such that the equilibrium distribution is maintained at all times. By simulating this subunit exchange process for a range of $k_1^-$, and comparing the outputs we can determine the $k_1^-$ value which best fits the data. Remarkably, but consistent with the qualitative change in subunit exchange profile we observe experimentally, we find it necessary to invoke a second rate constant, $k_2^-$, acting on a subpopulation of oligomers, to adequately fit the data at all temperatures (Fig. 5B&C). This second rate constant is particularly large at 45 °C, in accord with the notable ‘W-shaped’ distribution of the heterooligomers. From the temperature dependence of subunit exchange, we can extract activation energies of $97 \pm 18$ kJmol$^{-1}$ and $483 \pm 120$ kJmol$^{-1}$ for $k_1^-$ and $k_2^-$ respectively (Fig. 5D). This analysis shows that there is an increase in the rate of the overall reaction with temperature, however with the temperature dependence of $k_2^-$ being considerably more acute than for $k_1^-$.

While it is necessary to invoke two rate constant to explain the form of the data, it is not possible to unambiguously elucidate the nature of the second process. For example, two independent populations of oligomer, or different oligomers having different dissociation rates could reasonably explain the data. More complicated models involving additional free parameters could be constructed and will fit the data equally well, but cannot be statistically justified with the current data over this two-parameter model. The need to invoke a minimum of two rate constants in fitting the data clearly demonstrates that either at least two different modes of subunit exchange, or two types of oligomer with disparate exchange characteristics, exist.
DISCUSSION

Here we have reported on investigations into the oligomeric organization and dynamics of ScHSP26. Using both SEC-MALS and MS approaches we have shown that at ambient temperature this protein exists as a polydisperse ensemble comprising numerous oligomeric states. Previous reports have suggested the presence of multiple oligomeric states for ScHSP26 [43, 44], but here we have quantified their relative populations. Using a tandem MS technique we have demonstrated the 24mer to be ‘anomalously’ abundant at 25 °C, consistent with previous reports which have culminated in a cryo-electron microscopy-derived 3D structure of this oligomeric state [17]. However we show here that ScHSP26 also forms larger oligomers, up to 42 subunits in size at room temperature. The oligomers observed were composed exclusively of an even number of subunits. This strongly suggests there to be a basic dimeric ‘building block’, an observation consistent with the cryo-electron microscopy reconstruction of the ScHSP26 24mer [17], as well as structural studies of other sHSPs [2, 4, 5].

The activity of ScHSP26 is known to be temperature dependent, such that it only exhibits chaperone function at heat shock temperatures [30]. Here we investigated the effect of temperature on the oligomeric structure of ScHSP26 by using a combined approach of tryptophan fluorescence, NMR, and thermo-controlled MS. We found that an increase in temperature leads to the equilibrium shifting to monomers and dimers with partially flexible C-terminal extensions, such that at the highest temperature measured these sub-oligomeric species dominated the spectrum. Although these data show that the individual subunits undergo partial unfolding at higher temperature they also indicate that, unlike in mammalian sHSPs [37, 45], this region is not unstructured and highly mobile at 25°C. This agrees with the fact that a hydrophobic tryptophan is the pen-penultimate residue in ScHSP26 which, at ambient temperature, would be energetically costly to be located in a flexible and fully solvent-exposed region of the protein. Indeed, our previous studies showed that inserting a tryptophan residue into the C-
terminal extension of the sHSP αA-crystallin led to a reduction in flexibility of the extension [46]. As such it appears that a long and flexible C-terminal extension is a property of the polydisperse mammalian sHSPs.

Redistribution of HSP26 monomers into monomers and dimers at elevated temperature and in dilute solution has previously been observed for ScHSP26 [30] and other sHSPs [39, 47, 48]. Examination of the oligomeric region of the mass spectrum at elevated temperature revealed the remaining ScHSP26 to exist in forms of 32 subunits or larger, with the majority existing as a 40mer. These large oligomers therefore are thermodynamically favoured at higher temperatures, and represent the oligomeric form which are present under conditions during which ScHSP26 is active as a chaperone. Such an increase in oligomeric size at elevated temperatures has also been observed for the dodecameric sHSPs HSP16.9 from wheat [39], HSP18.1 from pea [28], and the polydisperse αB-crystallin [49]. This increase in size may therefore represent a mechanism of activation for the sHSPs.

To probe the inherent dynamics of this previously unreported range of oligomeric states which form at higher temperatures we monitored their subunit exchange in real time by means of MS. We found that ScHSP26 exchanged subunits in a stepwise manner such that at 34 °C complete equilibration occurred after approximately 95 minutes. At 39 °C this reaction was complete in about 30 minutes, which is very similar to the timescale for the exchange of the A and B subunits of α-crystallin at this temperature [31]. However, increasing the temperature to 45 °C resulted in the exchange reaction apparently occurring via a different regime such that at intermediate reaction times populations of both un-exchanged and fully exchanged protein were present. Detailed comparison with the data and in silico modelling of subunit exchange revealed that two parameters are required to adequately describe subunit exchange of ScHSP26 at all temperatures investigated here. The need for a two-parameter fit
may reflect the existence of multiple conformations of the same oligomers, potentially with differing activity or chaperoning roles.

An elegant fluorescence study has shown that the conformation of the middle domain of ScHSP26 undergoes rearrangement as the temperature is increased, resulting in chaperone-active oligomers [12]. Furthermore, this was shown to be a bi-phasic process. Taking our results in the context of these observations provides a possibility that the different middle-domain-conformers of ScHSP26 exchange subunits at different rates, with the active form being more dynamic. Two distinct conformers of the ScHSP26 24mer have been revealed by cryo-electron microscopy [17], and also for the related Archaeoglobus fulgidus HSP20.2 and Methanocaldococcus jannaschii HSP16.5 [50]. A temperature-dependent shift in the relative populations of these conformations was observed in the archaeal proteins [50], suggesting they may reflect forms of differing activity. Considering all these observations together, a complex picture of the mechanism of thermal activation of ScHSP26 emerges. At low temperatures the heterogeneous ensemble of ScHSP26 is dynamic, with subunits exchanging freely between oligomers. The rate of this exchange becomes more rapid as the temperature is increased, with the oligomers which comprise the ensemble being on average larger than those at low temperatures. Concomitant to this the middle domain undergoes a conformational change which leads to a highly dynamic oligomeric form with high chaperone activity.

The role of subunit exchange, via the dissociation and re-association of subunits, in sHSP chaperone action in vivo is not well understood. It appears, for ScHSP26 at least, that complete dissociation of dimers from the oligomers is not a pre-requisite for chaperone activity [32]. However dissociation is effectively a continuous process, and it may be that surfaces required for binding become exposed as a result of structural relaxation during the dissociation process. While only two sHSPs are present in Saccharomyces cerevisiae [11], it is clear that the genome of many species encode numerous sHSPs
[2], and that those members of the same evolutionary class found in the same cellular compartment will form a range of heterocomplexes \textit{in vivo} [25]. If the oligomers themselves are the active chaperones, then this would provide a large array of species, potentially all with different target protein affinities and specificities [28]. As such, the primary function of subunit exchange of the sHSPs could be to establish and maintain this chaperone ensemble, thereby providing broad stress and target protein specificity.
SIGNIFICANCE

Through the detailed interrogation of ScHSP26, this study highlights how fluctuations in both the quaternary structure and dynamics of proteins govern their function [51, 52]. Elucidating the details of these properties however is hampered by the heterogeneous nature of many protein assemblies. As we have demonstrated here, this can however be overcome by exploiting the high resolution of separation afforded by MS. Integrating MS measurements, which report on the properties of individual oligomers, with those derived from biophysical methods which report on the level of the constituent monomers (such as fluorescence and NMR spectroscopies here) therefore provides an attractive and general ‘hybrid’ approach for the study of proteins which populate heterogeneous oligomeric ensembles.
EXPERIMENTAL PROCEDURES

Protein Preparation

ScHSP26 was expressed in *E. coli* BL21 strain using standard methods. For uniformly $^{15}$N-labelled protein, cells were grown in minimal medium containing $^{15}$N-labelled ammonium chloride with 200 µg/ml carbenicillin at 32 °C. ScHSP26 purification from the soluble cell fraction was performed as described previously [53], with the following modifications. Protein was enriched in the 60–90% (w/v) ammonium sulfate fraction, and elution of the protein from the DEAE column (diethylaminoethyl-Sepharose Fast Flow resin; Sigma) was performed using a 0-400 mM sodium chloride gradient.

Size-Exclusion Chromatography and Light Scattering

ScHSP26 was analyzed by using size-exclusion chromatography with on-line light scattering, absorbance, and refractive index detectors [54]. A Bio-Sil 400 column (Bio-Rad) was connected in-line to a UV detector (Amersham Biosciences UV-900), a DAWN-EOS (Wyatt Technology) laser light scattering detector, and an Optilab-DSP (Wyatt Technology) refractive index detector. Samples were loaded onto the column at a concentration of 1.5 mg/ml and eluted with 50 mM phosphate buffer (pH 7.2) containing 100 mM sodium chloride (PBS).

Chaperone assay

Bovine pancreatic insulin (0.25 mg/ml) was incubated at 25 °C for 100 min in PBS (pH 7.2). Aggregation and precipitation were initiated by addition of DTT to a final concentration of 10 mM. ScHSP26 was added from a 5 mg/ml stock to give the final w/w ratios indicated.
**Fluorescence Spectroscopy**

ScHSP26 was diluted to 0.025 mg/ml in 100 mM ammonium acetate at pH 7.3 and heated from 20-90 °C at 1 °C/min on a Carey Eclipse fluorescence spectrophotometer. The sample was excited at 280 nm and emission was recorded at 330 nm. Fluorescence spectra were also recorded at three temperatures to provide more information on the structural transitions.

**NMR Spectroscopy**

$^1$H and $^{15}$N NMR spectra were acquired at magnetic field strength corresponding to a $^1$H resonance frequency of 500 MHz on a Varian Inova-500 spectrometer equipped with a triple-resonance pulsed-field gradient probe. 1.66, 0.33 or 0.17 mM $^{15}$N-labelled ScHSP26 was dissolved in 20 mM sodium phosphate buffer pH 6.5, containing 10% D$_2$O, 0.02% NaN$_3$. $^1$H-$^{15}$N 2D HSQC spectra were acquired at 25 °C, and 45 °C with 16 transients per increment in the direct dimension ($^1$H) and 256 increments in the $^{15}$N dimension. Spectral widths were 6000 Hz in the $^1$H dimension and 1000 Hz in the $^{15}$N dimension. $^1$H-$^1$H Watergate TOCSY spectra with $^{15}$N-decoupling were acquired at various temperatures between 20 and 60 °C over a spectral width of 6000 Hz in both dimensions, with 256 $t_1$ increments, 80 transients per increment using spin-lock periods of 30 and 50 ms. All NMR spectra were processed using Varian VNMR software (version 6.1c).

**Nanoelectrospray Mass Spectrometry**

Samples of $^{14}$N and $^{15}$N-labelled ScHSP26 were prepared for MS analysis by loading them onto a Superdex 200HR10/30 gel filtration column (GE Healthcare) and eluting at 0.3 ml/min with 200 mM
ammonium acetate at 6 °C. The resulting fractions corresponding to the protein oligomers were combined and concentrated with a centrifugal filtration device (Millipore Biomax), to give final protein concentrations of 1.3 mg/ml.

MS experiments were conducted using a previously described protocol [57] on a Q-ToF 2 mass spectrometer (Waters) which had been modified for high-mass operation [58]. Conditions were carefully chosen to allow the ionization and detection of the proteins without disrupting the noncovalent interactions that maintain the quaternary structure. The following instrument voltages and pressures were used: capillary 1.7 kV, sample cone 100 V, collision cell voltage 10 V, ion transfer stage pressure 9.0×10^-3 mbar, quadrupole analyzer pressure 9.5×10^-4 mbar, ToF analyzer pressure 1.7×10^-6 mbar and 10 μbar of argon in the collision cell. Sample heating was performed using an online thermo-controlled nanoESI device designed in-house [39]. To effect dissociation, gas-phase oligomers were collided with argon atoms in the collision cell of the mass spectrometer. This was achieved by increasing the accelerating voltage into the collision cell, and raising the gas pressure within to 35 μbar. Values in the range of 20-200 V were used for this parameter, depending on the extent of dissociation desired. The lower values were used to remove adducted solvent and buffer, thereby reducing peak width, and the higher values were used to effect actual dissociation of the oligomers [40]. Relative protein abundances were extracted by comparing peak intensities across the doubly stripped oligomer region [22].

**Subunit exchange**

Subunit exchange analyses were performed at 34 °C, 39 °C and 45 °C using the thermo-controlled nanoelectrospray probe. Equal volumes of the $^{14}$N and $^{15}$N-labelled $\text{S.}$chHSP26 were mixed immediately
prior to analysis, and acquisition of spectra was initiated within a 60 s equilibration period. To quantify our data we fitted the experimental data to a model for subunit exchange, described as follows.

The oligomerisation of sHSPs can be modelled as a series of consecutive equilibria of the form \( P_{i-1} + P_1 \rightleftharpoons P_i \), where \( P_i \) denotes an oligomer of size \( i \), as described previously [28]. For each step, a stepwise association constant can be defined as 
\[
K_i = \frac{[P_i]}{[P_{i-1}][P_1]} = \frac{k_i^+}{k_i^-},
\]
with a corresponding free energy 
\[
\Delta G_i = -RT \ln \frac{[P_i]}{[P_{i-1}][P_1]}. \]
The rate of change of each oligomer will then be given by:
\[
\frac{d[P_i]}{dt} = -ik_i^- [P_i] + k_i^+ [P_{i-1}][P_1] + (i + 1)k_{i+1}^- [P_{i+1}] - k_{i-1}^+ [P_i][P_1]
\]
where the dissociation of each individual subunit within an oligomer is assumed to be equally probable. From a series of experimentally determined oligomer populations together with the concentration of free subunits at a given condition it is possible, therefore, to determine the ratio \( \frac{k_i^+}{k_i^-} \). In the subunit exchange experiments here, \(^{14}\)N and \(^{15}\)N-labelled ScHSP26 is mixed. In such a situation, the rate of change of a given oligomer comprising \( i \) ‘light’ and \( j \) ‘heavy’ subunits will be given by:
\[
\frac{d[P_{i,j}]}{dt} = -(i + j)k_{i+j}^- [P_{i,j}] + k_{i+j}^+ ([P_{i,0}][P_{i-j,0} + [P_{0,i}][P_{j-1}]] + (i + j + 1)k_{i+j,1}^- ([P_{i,j+1}] + [P_{i-1,j+1}] - k_{i+j,0}^+ ([P_{i,0}] + [P_{0,i}] [P_{i,j}])
\]
Starting from the initial condition that oligomers of the form \([P_{i,0}]\) and \([P_{0,j}]\) have their equilibrium values, we can simulate the subunit exchange process that ultimately leads to heterogeneous oligomers with \( i=j \) being the most populated for any set of oligomers of size \( i+j \). The simplest method to fit the model to the kinetic data is to fit the data to one rate constant. We can accomplish this by assuming the dissociation rate for a monomer to leave an oligomer is independent of oligomer size (\( k_i^- = k_j^- \)).
following the variation of this constant as a function of temperature, we can estimate the activation parameters of the subunit exchange process.

While the 30 °C and 39 °C data fit well to this model, the 45 °C data does not, suggesting that the subunit exchange process is more complex. There are several ways in which this extra complexity can be introduced. The 45 °C data can be explained by specifying that a minor proportion of the oligomers have a significantly faster dissociation rate than the bulk. Thus the kinetic data has to be fitted globally to three constants: a ‘fast’ dissociation constant, a ‘slow’ dissociation constant and the proportion of fast/slow oligomers. The relative proportion of the two oligomer types is mainly specified by the 45 °C, data equating to ca. 25% of the total oligomers. Using the value obtained from fitting these data when fitting the 30 °C and 39 °C data gives a significantly reduced $\chi^2$ relative to the one parameter fits.
ACKNOWLEDGMENTS

The authors thank Alan Sandercock (now MedImmune) and Nelson Barrera (now Pontificia Universidad Católica de Chile) for helpful discussions on subunit exchange. JLPB is a Royal Society University Research Fellow; JAA an R. D. Wright National Health and Medical Research Council Fellow; AJB is a Canadian Institutes of Health Research Fellow; GD a C. J. Martin National Health and Medical Research Council Fellow; and CVR is a Royal Society Professor. JAC’s research is supported by the National Health and Medical Research Council and the Australian Research Council.
REFERENCES


FIGURE LEGENDS

Figure 1: Oligomeric distribution and chaperone activity of ScHSP26

(A) ScHSP26 examined by SEC-MALS illustrates the heterogeneity of the oligomeric ensemble. The molecular mass decreased across the peak from 950 kDa to 580 kDa and was centered around a maximum population species of 810 kDa, corresponding to an approximately 34 subunit oligomer. (B) Insulin reduction assay of the chaperone action of ScHSP26. At 25 °C, in the absence of ScHSP26, insulin B chain is reduced and exhibits an apparent increase in absorbance at 360 nm due to light scattering over a period of 100 min as a result of aggregation. At a subunit molar ratio of 0.2:1.0 ScHSP26:insulin, a 50% reduction in aggregation was observed due to the chaperone action of ScHSP26. At the higher ratios of 0.5:1.0 and 1.0:1.0, almost complete suppression of aggregation is achieved.

Figure 2: C-terminal flexibility of ScHSP26

(A) The dependence of tertiary/quaternary structure of the ScHSP26 oligomers on temperature was examined by observing the changes in intrinsic tryptophan fluorescence. ScHSP26 was heated from 20 - 90 °C at 1 °C/min (right). The sample was excited at 280 nm and emission was recorded at 330 nm. The resultant melt demonstrated that the tryptophan residues became significantly more solvent exposed between 20 °C and 45 °C whereas a much smaller, linear decrease in fluorescence was observed up to 90 °C. Fluorescence spectra were also recorded at three temperatures to provide information on the structural transitions (left). At 20 °C, the emission spectrum has a λ_max at ≈325 nm. At 45 °C, there was a decrease in the emission intensity concomitant with a dramatic shift in the λ_max to 352 nm. At 90 °C, a further decrease in fluorescence intensity but no λ_max shift occurred. These data
indicate that the majority of the tertiary structure in the vicinity of the tryptophan residues is lost during the initial unfolding transition. (B) The primary sequence of sHSPs in general is composed of an N-terminal region, the α-crystallin domain, and a C-terminal region. The latter is subdivided into a C-terminal tail, and a C-terminal extension, separated by the conserved I/VX/I/V motif. The sequence of ScHSP26 is aligned to those of animal sHSPs which have been demonstrated to contain flexible C-terminal extensions. Residues which are clearly identified by NMR spectroscopy measurements are highlighted in bold. (C) $^1$H-$^{15}$N 2D HSQC NMR spectra of ScHSP26 at I: 25°C and 1.66 mM; II: 45 °C and 1.66 mM; III: 45 °C and 0.33 mM; IV: 45 °C and 0.17 mM. At the same concentration (1.66 mM), solutions of HSP26 exhibit a minor enhancement of C-terminal flexibility with increased temperature, as evidenced by the greater number of cross-peaks resolved at random coil chemical shift values and an additional cross-peak from G212 (I and II). At 45 °C, 5- and 10-fold dilution an increased number of cross-peaks are observed (III and IV), suggesting greater C-terminal flexibility. This is consistent with an overall loosening of the structure and smaller, less-structured species of ScHSP26 being present under more dilute conditions.

Figure 3: MS demonstrates a temperature dependent dissociation of ScHSP26

(A) At 25 °C the mass spectrum of ScHSP26 was dominated by signal arising from a broad range of oligomers with a minor percentage of monomer and dimer. (B) At higher temperatures (43 °C and 60 °C) the relative amount of signal arising from the ScHSP26 oligomers was greatly diminished with a concomitant increase in the percentage of monomer and dimer.

Figure 4: Defining the oligomeric distributions of ScHSP26

(A) NanoESI-MS demonstrated that ScHSP26 exists as a broad range of oligomers at 25 °C (upper),
but forms mainly 40mers (open circles) and 42mers (shaded circles) at 43 °C (lower). Note that a 24mer is notably abundant at room temperature (black circles). (B) Collision-induced dissociation of monomers from the native oligomers gives rise to a series of stripped oligomers of reduced charge with sufficient peak resolution to accurately define their size distribution. Overlaid are schematic representations of the different species. (C) Portion of the doubly-stripped region highlighted in B for ScHSP26 at 25 °C (upper) and 43 °C (lower). From the peaks in this region of the spectra it was possible to assign charge states to individual oligomers, as well as calculate their relative abundance. (D) Histograms showing the most abundant oligomers present at 25 °C and 43 °C after analysis of the stripped oligomer region.

**Figure 5: Subunit exchange**

(A) The overlapping peaks of the doubly stripped oligomers were monitored at 34 °C (top), 39 °C (middle) and 45 °C (bottom) until exchange was complete. $^{14}$N and $^{15}$N-labelled ScHSP26 homo-oligomer peaks diminished with time, and in their place a peak corresponding to hetero-oligomers of the two subunits arose. This peak was broad, due to the variety of isotopic stoichiometries in the polydisperse assembly. As expected, the rate of exchange was found to increase with temperature, notably however, there appeared to be a change in the mechanism of exchange at 45 °C. This was evidenced by the fact that in the period between 2 and 4 min a population of fully exchanged ScHSP26 co-existed with the original homo-oligomers. (B) Alternative ‘top down’ contour plot representation of the data in A, left, compared with the simulated subunit exchange time-course which best fits each temperature, right. (C) Reduced $\chi^2$ surface for the fitting of subunit exchange data to two rate constants, $k_1^-$ and $k_2^-$. (D) Arrhenius plots for the temperature dependence of $k_1^-$ and $k_2^-$ allows the extraction of the activation energies of the two processes.
FIGURE 1
FIGURE 2
FIGURE 3

A

monomers & dimers

oligomers

B

43°C

60°C
FIGURE 4
FIGURE 5