Real-Time Monitoring of Protein Complexes Reveals their Quaternary Organization and Dynamics

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SUMMARY

The dynamics of protein complexes are crucial for their function yet are challenging to study. Here, we present a nanoelectrospray (nESI) mass spectrometry (MS) approach capable of simultaneously providing structural and dynamical information for protein complexes. We investigate the properties of two small heat shock proteins (sHSPs) and find that these proteins exist as dodecamers composed of dimeric building blocks. Moreover, we show that these proteins exchange dimers on the timescale of minutes, with the rate of exchange being strongly temperature dependent. Because these proteins are expressed in the same cellular compartment, we anticipate that this dynamical behavior is crucial to their function in vivo. Furthermore, we propose that the approach used here is applicable to a range of nonequilibrium systems and is capable of providing both structural and dynamical information necessary for functional genomics.

INTRODUCTION

Understanding the function of proteins requires information about three of their basic properties: structure, interactions, and dynamics. MS presents an established approach toward the study of the first two facets. Primary structural details, regarding sequence and posttranslational modifications, are the focus of numerous MS-based proteomics studies (Aebersold and Mann, 2003). Furthermore, given that the vast majority of proteins perform their cellular roles in the form of higher-order assemblies (Sali et al., 2003) and that aspects of their quaternary structure can be maintained in the gas phase (Ruotolo and Robinson, 2006), the application of MS to the direct study of protein complexes is an extremely exciting avenue of research (Benesch et al., 2003). The application of such an online MS monitoring approach, adapted by using nESI (Fligge et al., 1999), has in the last few years been extended to studying the dynamics of multimeric proteins (Aquílna et al., 2005; Sobott et al., 2002). An alternative strategy is to perform such time-resolved experiments in an offline mode, whereby a reaction mixture is sampled repeatedly, rather than continuously. Such an approach has previously been used in several studies, including monitoring the assembly (Fändrich et al., 2000; Stockley et al., 2007) and dynamics (Keetch et al., 2005) of protein complexes. In these cases, the dead time (i.e., the time between mixing of components and detection) is ~1 min. Early time points have become accessible using an ESI capillary mixer (Wilson and Konermann, 2003), but until this technology can be transferred to an nESI platform, the applicability to the study of macromolecular assemblies is limited (Benesch et al., 2007). Continuous monitoring of a reaction mixture by means of nESI suffers from the difficulty of maintaining a stable flow over the duration of the experiment; in addition, prolonged electrospraying can induce electrochemical changes in the solution within the capillary, which may interfere with the reaction kinetics (van Berkel et al., 1997). Repeated manual sampling of a reaction can also prove problematic because there is a limit to the number of time points obtainable, as dictated by the experimentalist’s capacity, and the capillary-to-capillary irreproducibility can lead to impairment in data quality (Keetch et al., 2003).

Here, we present an automated nESI method for rapid repeated monitoring of the dynamics of protein complexes in real time. This approach obviates many of the difficulties associated with the real-time monitoring of reactions by MS, as described above, and is capable of providing reproducible data for a wide range of dynamic species. We describe in detail the development of the method and demonstrate the advantages of the technology by application to the quantitative monitoring of the kinetics of an enzymatic digestion reaction. We subsequently examine the dynamics of two previously uncharacterized sHSPs—HSP18.1 and HSP17.6 from Arabidopsis thaliana. The sHSPs are a widely diversified family of molecular chaperones that are thought to prevent irreversible protein aggregation by holding destabilized substrates soluble for subsequent refolding by...
ATP-dependent chaperones (Haslbeck et al., 2005; Narberhaus, 2002; van Montfort et al., 2002). The importance of these proteins is evidenced by their being found in almost all organisms, and they have been implicated in a range of disease states, including cataract, cancer, myopathies, motor neuropathies, and neurodegeneration (Horwitz, 2003; Sun and MacRae, 2005; Welsh and Gaestel, 1998). We find HSP18.1 and HSP17.6 to be involved in several different well, so as to generate improved statistics, or different reactions altogether, to improve overall duty cycle.

There are numerous reactions that proteins undergo in the cell. Among the most important of these involve enzymes and their substrates, and we therefore chose such a system to characterize our method. The enzyme, Trypsin, and substrate, Cytochrome C (CytC), were incubated together, and three aliquots were deposited onto the sample plate of the robot, which was programmed to sample each well repeatedly over a 5 hr period (Figure 1A). The total-ion chromatogram displays “blocks” of signal, each corresponding to a separate infusion. These blocks are grouped in threes, as a result of the sequential sampling of the wells. A high degree of consistency was found in the intensity of signal within each group, evidencing the high level of reproducibility between emitters on the nESI chip. The total ion current, however, increases over the course of the reaction, suggesting that more species are observed as the reaction progresses. Mass spectra obtained at the beginning, middle, and end of the time course are shown in the inset of Figure 1A. At the first time point, only three major peaks, corresponding to the five, six, and seven charge states of intact CytC, are observed. After 300 min, these peaks are no longer observed: instead, the spectrum is dominated by peaks at lower m/z corresponding to tryptic peptides, the most intense of which were identified as CytC119-123, CytC28-38, and CytC14-22 (Table S1). At intermediate times, however, both these reactant and product species can be observed simultaneously. Moreover, a species corresponding to a 9.5 kDa fragment (CytC71-79) that is not observed in either the initial or final spectrum can be identified, demonstrating how this real-time approach can be used to detect transient intermediates that would not be identified in studies performed at equilibrium.

To get a quantitative description of the reaction, we plotted the relative intensities of the most intense peaks corresponding to reactants, intermediates, and products in single MS scans as a function of time (Figure 1B). The very small deviation between measurements demonstrates the excellent reproducibility afforded by this approach. Exponential decay of the intact CytC is observed, with a concomitant sigmoidal increase of the tryptic peptide products. The intermediate ions show an initial increase, followed by gradual decay as they are themselves digested. From these data, a rate constant of 7.6 ± 0.6 × 10⁻³ min⁻¹ was extracted, which is consistent with previously reported time-scales (Stone et al., 2001) and current models of enzymatic digestion (Srividhya and Schnell, 2006). Because the rate of many biological reactions is temperature dependent, we tested the thermo-control capabilities of the platform by performing the same experiment at a range of temperatures up to 50°C (Figure 1C). At elevated temperatures, the reaction proceeded much more rapidly: at 37°C, 45°C, and 50°C, the reaction was complete in ~84, ~48, and ~23 min, with rate constants of 2.5 × 10⁻² min⁻¹, 5.7 × 10⁻² min⁻¹, and 1.6 × 10⁻¹ min⁻¹, respectively. Even at the highest temperature, monitoring transient intermediates was still possible (Figure S3). This increase in rate of digestion with temperature is a result of more molecules overcoming the activation energy barrier. Furthermore, by assessing the temperature dependence of the rate constants, we extracted an activation energy of 88.8 kJmol⁻¹ (Figure 1C, inset).

To investigate the difference between the repeated sampling used above and a continuous sampling approach, we set the robot to perform both these monitoring approaches on a CytC digest reaction in 33% methanol (Figure 1D). This solution condition induces a molten-globule form of CytC (Bychkova et al., 1996), which is particularly susceptible to acid-induced
denaturation (Konermann and Douglas, 1997; Konermann et al., 2001). When this digestion reaction was monitored continuously (i.e., as an uninterrupted infusion through a single nanospray emitter), it reached completion within 30 min. However, with repeated sampling (i.e., multiple short infusions through a new nozzle every time), the rate of reaction was much slower, such that ~85% of the protein remained undigested after the 30 min, and the reaction took an order of magnitude longer to reach completion (Figure 1D). In the case of continuous sampling, we attribute these differences to prolonged electrospraying causing a decrease in pH in the aqueous solution within the emitter (van Berkel et al., 1997), which results in a further destabilization of the CytC (Konermann and Douglas, 1997; Konermann et al., 2001). The resulting increase in frequency and magnitude of local protein unfolding events consequently leads to an increased susceptibility to digestion. This result therefore shows that, depending on solution conditions, continuous sampling may be unreliable. In contrast, repeated sampling is not affected by electrochemical changes that might arise from prolonged electrospraying. As such, the automated nESI platform we have described and characterized here provides a means for producing reliable and reproducible kinetic data. Furthermore, because using nESI is essentially a prerequisite for the routine analysis of macromolecular assemblies by means of MS (Benesch et al., 2007), this system has the considerable advantage of being applicable to the study of such species.

Dynamic Subunit Exchange of the sHSPs

Several members of the sHSPs family have been shown to be very dynamic, in that oligomers are capable of freely exchanging subunits (Haslbeck et al., 2005; Narberhaus, 2002; van Montfort et al., 2002). Moreover, as is the case for the mammalian...
α-crystallins, this exchange can also occur between closely related protein oligomers, and it is the resulting hetero-complexes that are the functional entities in vivo (Horwitz, 2003). In Arabidopsis thaliana, 19 genes encode sHSPs, with several members being localized in the same cellular compartments (Siddique et al., 2008). Here, we apply our automated nESI MS approach to two cytosolic class I sHSPs from Arabidopsis thaliana—HSP17.6 and HSP18.1. nESI mass spectra of these proteins display principal charge state series centered around 6900 m/z, corresponding to masses of 210,258 Da and 216,301 Da for HSP17.6 and HSP18.1, respectively (Figure 2A). Additional peaks around 2200 m/z are also observed in both cases. These peaks correspond to charge states of monomeric species, and their low charge state suggests that they arise from a solution-phase equilibrium with the oligomers (Benesch et al., 2003). Comparison with the spectra of the proteins under denaturing conditions reveals that both are composed of 12 noncovalently bound subunits.

Incubating the two proteins at an equimolar ratio at room temperature for 2 hr before analysis resulted in the spectrum shown in the upper panel of Figure 2B. The lower panel shows the overlaid spectra of the individual proteins. The equilibrated mixture, however, gave a very different spectrum, suggesting the occurrence of a subunit exchange reaction. To assist with the assignment of the peaks arising from this heterogenous ensemble, we simulated spectra for the 31+ charge state of the different candidate heterododecamers. The composite peaks that would be expected for unrestricted exchange of monomeric subunits to an equilibrium position given by a binomial distribution of equimolar components (top panel), and with the added constraint that the exchange units were dimers (lower panel), are shown in Figure 2C (see Experimental Procedures). These simulations are compared with an expansion of the 31+ region of the experimental data (middle panel). Clearly, the profile of the peaks observed is not fitted by monomeric exchange but is very well represented by dimeric exchange. Differences between the relative intensities of the different peaks in the modeled and experimental data likely arise from a fractionally greater proportion of HSP18.1 than HSP17.6 in the reacting solution. Overall, however, the persistence of dimeric exchange shows that the dimer interfaces are not compromised on the timescale of the experiment.

To monitor the kinetics of this subunit exchange reaction, we used our automated nESI approach. Figure 3A shows the evolution of the spectra obtained over a 60 min time course at 24°C, for the 31+ charge state (Figure 2C). At the first time point (2 min), the dominant peaks correspond to the homododecamers 12:0 and 0:12 (expressed as the number of subunits of HSP17.6:HSP18.1). Some signal arising from the 10:2 and 2:10 heterododecamers is also observed. After 6 min, the signal corresponding to these.
heterododecamers has increased, and 8:4 and 4:8 are also clearly observed. After 20 min, almost no homododecamers are observed, and a significant population of 6:6 is observed. At the end of the time course, the signal has converged so that the most prevalent species is 6:6, reflecting the equimolar mixture of reactant sHSPs.

The intensity of the peaks corresponding to 12:0 and 0:12 are plotted as a function of time in the upper panel of Figure 3B. The abundance of the homododecamers follows an exponential decay function for approximately the first 15 min of the reaction, well represented by first-order kinetics. By plotting the natural logarithm of this decay, we get a first-order rate constant of 0.16 min$^{-1}$ (inset). When we repeated the experiment at 30°C, the reaction reached completion after ~20 min, with a rate constant of 0.40 min$^{-1}$. The lower panel of Figure 3B shows the relative abundance of the different heterododecamers formed. The plots corresponding to 10:2 and 8:4 both go through maxima, at approximately 8 min and 22 min, respectively, whereas that for 6:6 rises steadily to a maximum. At 30°C, the reaction profile is unchanged except in its time frame. Overall, these observations are consistent with a reaction wherein individual dimeric units are incorporated in a sequential fashion and dissociation of the oligomers is rate determining (Sobott et al., 2002).

As such, this rapid experiment has provided details as to oligomer stoichiometry (dodecamers), structural composition (robust dimeric building blocks), and dynamics (rapid sequential dimeric exchange).

**DISCUSSION**

Here, we have described a robust and versatile automated nESI approach for studying the dynamics of protein complexes, allowing the quantitative monitoring of their reactions, which occur on the timescale of minutes. The reproducibility due to the chip-based nESI, as well as the ability to perform measurements in multiplicate, results in a precision of measurement not readily attainable by conventional nESI approaches. In the first instance, we demonstrated the applicability of this platform for monitoring enzyme kinetics, by monitoring digestion of CytC by Trypsin. Rate constants were determined, and, moreover, a proteolysis intermediate, CytC$_{1-79}$, was observed. This result raises the intriguing possibility of determining the relative susceptibility of different cleavage sites to digestion, thereby gaining information as to the accessibility and flexibility of certain polypeptide regions. Such a real-time approach amounts to an extension of the limited proteolysis approach (Hubbard, 1998) and holds exciting potential for probing both structure and conformational dynamics of proteins and their complexes.

Subsequently, we applied this reaction-monitoring strategy to the subunit exchange of two previously uncharacterized and closely related sHSPs from *Arabidopsis thaliana*. nESI-MS analysis revealed these proteins to be dodecameric, and their subunit exchange reaction was successfully monitored at both 24°C and 30°C, even in the latter case when the reaction was complete within a few minutes. The apparent adherence to first-order kinetics suggests that dissociation of the oligomers is the rate-limiting step in the exchange reaction, a feature also observed in other members of this protein family (Bova et al., 1997). Furthermore, a rate constant at 24°C of 0.16 min$^{-1}$ is very similar to that observed...
for a different pair of dodecameric plant sHSPs (Sobott et al., 2002) and highlights the remarkably dynamic nature of members of this protein family. Moreover, the extremely high resolution of separation afforded by MS allowed the monitoring of the relative populations of the heterododecamers formed. We therefore determined that subunit exchange is achieved by the sequential incorporation of dimeric units of one sHSP into a dodecamer of the other. This dynamical observation allows us to draw the structural conclusion that the protein dodecamers must be composed of dimeric units. Such dimeric “building blocks” have been suggested as being a common feature of sHSPs (Haslbeck et al., 2005; van Montfort et al., 2002), but, interestingly, there is a marked contrast with previous subunit exchange data for two other dodecameric plant sHSPs, Pisum sativum HSP18.1 and Triticum aestivum HSP16.9, where heterododecamers composed of an odd number of each of the components (e.g., [HSP18.1]_o[HSP16.9]_o) were also observed (Sobott et al., 2002). This difference implies variability in the interfaces between subunits across even these evolutionarily closely related sHSPs. Because the subunit interfaces are thought to be at least partly responsible for the substrate binding function of these proteins (Haslbeck et al., 2005; van Montfort et al., 2002), this suggests that adaptation has occurred in these interfacial areas of the sequence to regulate substrate specificity.

This ability to elucidate information as to oligomeric organization via real-time nESI subunit monitoring is very attractive. Other strategies have been used in combination with nESI to achieve this goal—notably, gas phase dissociation (Benesch et al., 2006; Benesch and Robinson, 2006) and destabilization in solution through either temperature regulation (Benesch et al., 2003) or the addition of perturbants (Hernández et al., 2006). A major advantage of our approach is that it involves the monitoring of the protein complexes in their native state, with the proviso of an MS-compatible buffer. As such, potentially nonspecific interactions are avoided. Furthermore, the subunit exchange is nondissociative, in the sense that intact oligomers, rather than subolgimeric species in solution, are being monitored. Suboligomeric units may be elucidated, therefore, independent of their stability outside the intact oligomer. This advantage is exemplified in the results shown in Figure 2A, which reveal only dodecameric and monomeric forms of the sHSPs. The presence of dimeric substructure is revealed only upon performing the subunit exchange experiment.

Other methods have been used for the study of subunit exchange reactions, including fluorescence resonance energy transfer (Bova et al., 1997), native gel electrophoresis (van den Oetelaar et al., 1990), and affinity chromatography (Schneider et al., 2001). These methods have the disadvantages of requiring the use of a potentially invasive tag, providing an “average” of the whole reaction mixture, or resulting in a poor resolution of separation, or a combination thereof. In contrast, the high-resolution separation afforded by an MS approach allows the relative quantitation of the different species within heterogenous ensembles (Aquilina et al., 2003). We have demonstrated here that it is the real-time monitoring of these populations that allows the elucidation of dynamical and structural details. Furthermore, monitoring the exchange of a protein complex with its isotopically labeled but isostructural equivalent obviates the need for a tag (Keetch et al., 2005) and represents a completely generalized strategy for future investigations. As such, the approach described here provides an exciting, robust, and universal method for the study of the subunit exchange reactions and nonequilibrium states of protein complexes in general.

**SIGNIFICANCE**

The sHSPs are a family of molecular chaperones found in almost all organisms studied to date. Detailed structural studies on these proteins are comparatively scarce because of the frequently polydisperse and dynamic character of these proteins (Haslbeck et al., 2005; Horwitz, 2003; Narberhaus, 2002; van Montfort et al., 2002). In fact, it appears that this dynamic nature, particularly their ability to exchange subunits and form hetero-oligomeric species, may itself be crucial to their function in vivo (Haslbeck et al., 2005; Horwitz, 2003; Narberhaus, 2002; van Montfort et al., 2002). Here, we have developed a robust and universal nESI-MS approach for monitoring the reactions of protein complexes in real time and have applied it to the subunit exchange of two sHSPs from Arabidopsis thaliana. By quantifying the relative populations of the different homo- and hetero-oligomeric species as the reaction proceeds, this report represents the first detailed study, to our knowledge, of a subunit exchange reaction between two species that are found in the same cellular compartment in vivo.

Through the simultaneous determination of both structural and dynamical properties of these proteins, showing them to be highly dynamic dodecamers with dimeric substructure, this study also demonstrates the versatility of nESI-MS. Functional genomics is concerned with combining structural and dynamical information so as to understand the roles and mechanisms of action of proteins and their noncovalently bound complexes in the cell. Most established structural genomics approaches, however, are not well suited to studying the dynamics of protein complexes, because they are performed at equilibrium or on quenched states. In contrast, nESI-MS has not only become an established structural genomics approach (Robinson et al., 2007) but also, as we have shown here, is capable of providing complementary dynamical insights crucial to understanding protein function by generating a large amount of information in real time.

**EXPERIMENTAL PROCEDURES**

**Automated nESI- MS**

All mass spectral measurements were performed using a Nanomate HD nESI system (Advinion BioSciences Ltd., Ithaca, NY) (van Peit et al., 2002). Mass spectra were recorded on either an LCT or Q-ToF II mass spectrometer (both from Waters, UK). Spectra were calibrated externally using 33 mg/ml cesium iodide. Data were acquired and processed with MassLynx software (Waters, UK) and are shown with no background subtraction.

**Enzyme-Substrate Reaction**

Fifty microliters of 100 μM horse-heart CytC (Sigma C-2506), in water, was combined with 50 μl of 200 mM ammonium bicarbonate and 6.25 μl of 0.5 mg/ml sequence-grade modified Trypsin (Promega V5113), giving a final enzyme:substrate ratio of 1:38.

Mass spectra were obtained on the Q-ToF II in positive ion mode, with sample cone, 180 V; extractor cone, 10 V; ion transfer stage pressure,
6.5 × 10⁻³ mbar; quadrupole analyzer pressure, 5.5 × 10⁻³ mbar, and ToF analyzer pressure, 4.7 × 10⁻⁷ mbar. nESI was achieved with a spray voltage of 1.65 kV and a head pressure of 0.4 psi (28 mbar) set on the Nanomate.

To extract kinetic information, each single 5 s acquisition time point was analyzed without smoothing. Relative abundances were assessed from the peak heights of the species being monitored (Table S1). Line-fitting was achieved using SigmaPlot 2001 (SPSS Science, Chicago, IL), and error bars represent three standard deviations from the mean.

Subunit Exchange Reaction

HSP18.1 and HSP17.6 were expressed in Escherichia coli and were purified as described elsewhere (Basha et al., 2006). Samples were buffer exchanged into 200 mM ammonium acetate by using a Superdex 200HR10/30 column (GE Healthcare). To monitor the subunit exchange kinetics in real time, the proteins were combined at a molar ratio of 1:1, to give a final protein concentration of 10 μM (monomer) for each component, according to UV absorbance at 280 nm.

Mass spectra were obtained on the LCT in positive ion mode, with sample cone, 150 V (individual components) or 200 V (subunit exchange experiment); extractor cone, 5 V; ion transfer stage pressure, 7.33 mbar; and ToF analyzer pressure, 1.1 × 10⁻⁸ mbar. nESI was achieved with a spray voltage of 1.8 kV and a head pressure of 0.55 psi (38 mbar) set on the Nanomate. The reaction mixture was sampled 34 times over the course of the 1 hr experiment, with each 1 μl aliquot being electrospayed for 1 min.

Simulated spectra were constructed using SigmaPlot 2001, as described elsewhere (Sobott et al., 2002). Theoretical intensities of the different dodecamers were calculated on the basis of a binomial distribution of the components HSP18.1 and HSP17.6 at a ratio of 1:1, assuming an equal preference for each composition (Figure 2, top panel). A second simulation was performed with the restriction that the exchanging units were homodimers (Figure 2, lower panel). Kinetics were monitored by signal-averaging all scans in each separate infusion, and the peak heights of the 31+ charge states of the various dodecamers were monitored. Data were normalized to the final relative distributions dictated by the binomial distribution, and kinetic parameters were extracted using SigmaPlot 2001.

SUPPLEMENTAL DATA

Supplemental Data include 3 figures and 1 table and can be found with this article online at http://www.chembiol.com/cgi/content/full/15/3/246/DC1/.

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