Chaperones Skp and SurA dynamically expand unfolded OmpX and synergistically disassemble oligomeric aggregates

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Edited by Susan Marqusee, University of California, Berkeley, CA; received October 25, 2021; accepted January 19, 2022

Periplasmic chaperones 17-kilodalton protein (Skp) and survival factor A (SurA) are essential players in outer membrane protein (OMP) biogenesis. They prevent unfolded OMPS from misfolding during their passage through the periplasmic space and aid in the disassembly of OMP aggregates under cellular stress conditions. However, functionally important links between interaction mechanisms, structural dynamics, and energetics that underpin both Skp and SurA associations with OMPS have remained largely unresolved. Here, using single-molecule fluorescence spectroscopy, we dissect the conformational dynamics and thermodynamics of Skp and SurA binding to unfolded OmpX and explore their disaggregation activities. We show that both chaperones expand unfolded OmpX distinctly and induce microsecond chain reconfigurations in the client OMP structure. We further reveal that Skp and SurA bind their substrate in a fine-tuned thermodynamic process via enthalpy–entropy compensation. Finally, we observed synergistic activity of both chaperones in the disaggregation of oligomeric OmpX aggregates. Our findings provide an intimate view into the multifaceted functionalities of Skp and SurA and the fine-tuned balance between conformational flexibility and underlying energetics in aiding chaperone action during OMP biogenesis.

Molecular chaperones are key cellular components that play fundamental roles in maintaining cellular proteostasis (1, 2). Essential activities of chaperones include the assistance of de novo protein folding, the stabilization of nonnative proteins in folding competent or unfolded states, and the rescue of misfolded and aggregated proteins (3–5). Chaperones are an integral part of a wide range of protein quality-control systems, and their activities are intimately coupled to the biogenesis networks that aid the structural and functional maturation of proteins from their site of cellular synthesis to their target cellular compartments.

One network where chaperone activity is of particular relevance is the biogenesis of outer membrane proteins (OMPs) (6–7). OMPs are a diverse group of β-barrel membrane proteins found in the outer membrane of Gram-negative bacteria, mitochondria, and chloroplasts. They fulfill a plethora of functions in cell signaling, metabolism, and transport (8–10); are indispensable to the survival of bacteria (10, 11); and constitute important virulence factors and drug targets (12–14). The OMP biosynthesis pathway is highly complex and conserved across all kingdoms of life (15) and involves the coordinated action of a multicomponent protein machinery that aids in overcoming the many hurdles that these proteins have to surmount on their way to their target outer membrane (4, 7, 16).

In Gram-negative bacteria, OMPs are translated in the cytoplasm, from where they are translocated across the inner bacterial membrane via the Sec machinery to the periplasmic space (17, 5).

Within this aqueous compartment, OMPs are escorted to the outer membrane in an unfolded state (denoted as the uOMP state) with the aid of various chaperones that maintain the largely insoluble and aggregation-prone uOMP polypeptide chains in a protected, partially unfolded state (18, 19). At the outer membrane, uOMPs are transferred to the β-barrel assembly machinery (BAM), which facilitates their native folding and insertion into the membrane (20). Noteworthy, the periplasm is devoid of any known source of energy-providing molecules, such as adenosine triphosphate (ATP); hence, all chaperones as well as the entire folding machinery likely operate without the aid of external energy, following thermodynamic principles (21).

Two chaperones which have been shown to be indispensable for the biogenesis of bacterial OMPS are the 17-kilodalton protein (Skp) (22) and survival factor A (SurA) (23). Skp and SurA, both located in the periplasm, exhibit antifolding activity (also known as holdase activity), whereby they sequester uOMP substrates to prevent aggregation until they reach the bacterial outer membrane.

Significance

Outer membrane proteins (OMPs) are crucial for the survival of bacteria. The two chaperones 17-kilodalton protein (Skp) and survival factor A (SurA) play key roles in OMP maturation by keeping unfolded OMP proteins soluble in the periplasm. However, their functionalities are incompletely understood. Here, we establish connections between structural and energetic features employed by the two chaperones when interacting with unfolded OmpX. We find that expansion, accompanied with fast polypeptide chain reconfiguration, prevents unfolded OmpX from misfolding and aggregating. Moreover, chaperone interaction with unfolded OmpX is thermodynamically calibrated, allowing for a fine-tuned association of chaperones with OMPs in the adenosine triphosphate-depleted periplasm. We further discovered that Skp and SurA act together as disaggregases and are able to disassemble oligomeric OMP aggregates, revealing remarkable functionalities of this periplasmic chaperone system.

Author contributions: N.C., A.H., G.K., and M.S. designed research; N.C. and A.S. performed research; N.C., A.H., M.Q.M., and G.K. contributed new reagents/analytic tools; N.C., A.H., and M.S. analyzed data; and N.C., A.H., G.K., and M.S. wrote the paper.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at http://www.pnas.org/lookup/suppl/doi:10.1073/pnas.2118919119/-/DCSupplemental.

Published February 25, 2022.

PNAS 2022 vol. 119 no. 9 e2118919119 https://doi.org/10.1073/pnas.2118919119 | 1 of 11
Energetic mechanisms underlying Skp and SurA chaperone–OMP activities of Skp and SurA and find synergistic activity of both chaperones differ strongly and are dictated entropic contributions of complex formation and find that the interaction modes of both chaperones expand the unfolded polypeptide chain upon binding.

Studying Chaperone Effects on uOmpX Conformational Dynamics

In addition to the well-described holdase activities of Skp and SurA that protect OMPs from misfolding or aggregating, a recent study has suggested that Skp disaggregates oligomeric uOMP structures (38). While SurA has not been directly implicated as a disaggregator, modeling studies propose a synergistic interaction among these chaperones especially under conditions of stress (18, 39), thus raising the question of the role that both chaperones played in disassembling OMP aggregates.

To gain insights into the multifaceted functionalities of Skp and SurA and their action mechanisms, we study here the conformational dynamics and thermodynamics of the eight β-stranded protein OmpX in the presence of the chaperones and explore their disaggregation activities. Using single-molecule Förster resonance energy transfer (smFRET), we resolve the heterogeneities, structural dynamics, and thermodynamics underlying the different states of uOmpX at near-native conditions. Strikingly, we find that both chaperones expand the unfolded polypeptide chain upon binding.

The degree of expansion is concentration dependent for SurA, but not for Skp. Probing structural changes and chaperone interaction at different temperatures, we gain insights into the enthalpic and entropic contributions of complex formation and find that the interaction modes of both chaperones differ strongly and are dictated by entropy−enthalpy compensation. Finally, we use fluorescence correlation spectroscopy (FCS) to probe the disaggregation capabilities of Skp and SurA and find synergistic activity of both chaperones in the disassembly reaction of oligomeric OmpX aggregates.

Our findings provide fundamental insights into the structural and energetic mechanisms underlying Skp and SurA chaperone–OMP interactions and their role in OMP biogenesis.

Results

Studying Chaperone Effects on uOmpX Conformational Dynamics under Native-Like Conditions by smFRET. Probing chaperone effects and conformational dynamics of uOMPs under native-like conditions (i.e., in the absence of or at very low denaturant concentrations) has been difficult to achieve due to the aggregation-prone nature of OMPs in aqueous environments (4, 7, 40). Single-molecule methods, in particular single-molecule fluorescence techniques such as smFRET, operate at very low concentrations (i.e., in the pM range) and thus alleviate protein aggregation and precipitation challenges (41–43). smFRET measurements further provide access to subpopulation-specific conformational heterogeneity, even at very low denaturant concentrations or the absence thereof (44) and allow probing of long-range polypeptide chain dynamics across a wide spectrum of timescales in the presence of binding partners and without the need of synchronizing the ensemble with perturbation/relaxation techniques (45–47).

We therefore set out to develop an smFRET assay that allowed us to study the conformational dynamics of single uOmpX polypeptide chains in native-like aqueous environments and the effect that the chaperones Skp and SurA have on the structural and dynamic properties of uOmpX.

To begin, we prepared an N- and C-terminal fluorescently labeled double-cysteine OmpX variant (OmpX1,149) furnished with donor and acceptor dyes (Fig. 1A and Methods). The placement of the FRET-dye pair at the terminal ends of the protein allowed us to monitor the conformational properties of the uOmpX polypeptide chain. After preparation, labeling, and refolding of the protein in the presence of the surfactant lauryldimethylamine-N-oxide (LDAO) (SI Appendix, Fig. S1 A and C and Methods), we subjected OmpX1,149 to a multistep denaturation and dilution protocol to transfer it to a micelle-free aqueous buffer environment at picomolar concentrations (Fig. 1A). Briefly, we first denatured refolded OmpX1,149 with 6 M guanidinium chloride (GdmCl) at micromolar concentrations and subsequently diluted it in the presence of 6 M GdmCl to a concentration of 20 nM. This was followed by a 1,000-fold dilution step into GdmCl-free buffer or into GdmCl-free buffer supplemented with the chaperones Skp or SurA. The resultant [GdmCl] after the final dilution step was 6 mM and the remaining [LDAO] was <100 nM, thus yielding uOmpX at 20 pM in a micelle-free native-like aqueous buffer environment at picomolar concentrations (Fig. 1A).

The resultant [GdmCl] after the final dilution step was 6 mM and the remaining [LDAO] was <100 nM, thus yielding uOmpX at 20 pM in a micelle-free native-like aqueous buffer that is largely devoid of denaturant (denoted as uOmpXaq), smFRET measurements were performed directly after the dilution protocol, thus mimicking the scenario when newly secreted OMPs enter the periplasm and encounter their interacting chaperones. Subsequently, fluorescence bursts were recorded from a large number of individual protein molecules diffusing through the confocal detection volume to generate FRET efficiency (E) histograms (Methods). Because smFRET provides intramolecular distance information in the nanometer range by measuring the energy transfer between fluorescent donor and acceptor dyes attached to the polypeptide chain, E histograms report on the diversity of conformations with different levels of compactness (i.e., distance between the two dyes) and provide information on intrachain dynamics (43, 48).

uOmpX in the Absence of Chaperones Is Structurally Heterogeneous and Exhibits Multitier Dynamics. Under native-like buffer conditions and in the absence of chaperones, the E histogram of uOmpXaq at 25 °C in the first 2 h of measurement exhibited a broad distribution with a FRET efficiency peak of the aqueous uOmpX state (Eaq) located at Eaq ∼ 0.77 and a tail spreading toward low transfer efficiencies (Fig. 1B, Left). Broadened E distributions are generally ascribed to conformational heterogeneity with interconversion dynamics on timescales comparable to or longer than the observation time (~1 to 2 ms). Indeed, the width of the unfolded state population is in excess of that expected for a distribution that is limited only by detection shot noise (red cityscape in Fig. 1B), indicating the existence of a heterogeneous ensemble of conformations. Even at elevated temperatures of 31°C and 37°C (Fig. 1B, Center and Right, respectively), we observed a broadened E distribution further
indicative of a rough energy landscape of the uOmpX aq state (34). Additionally, recurrence analysis (49) revealed slow interconversions among the various substates on timescales longer than the burst duration ($\geq 100$ ms; Fig. 1C). This suggests collective and transient conformational changes of remaining secondary motifs or tertiary long-range interactions, as have been described before for other OMPs under denaturing conditions (34, 50). Fluorescence lifetime and sufficient rotational averaging allowed us to exclude the possibility that the broadening originates from changes in dye quantum yields or a restricted rotational freedom of the dyes (SI Appendix, Table S1). We also observed a modest expansion of the uOmpX aq chain on the timescale of 4 to 6 h (SI Appendix, Results, Fig. S1 D–H, Table S2), in line with reports which suggest that this mechanism...
prevents aggregation in the absence of both a membrane mimetic environment and chaperones (40). Thus, our results indicate that uOmpXaq might also possess an intrinsic nonaggregating ability.

In addition to the slow interconversion dynamics that give rise to broadened uOmpXaq distributions, the unfolded polypeptide chain exhibited fast submillisecond reconfiguration dynamics at early time points (0 to 2 h), as indicated by a displacement of the uOmpXaq population from the static FRET line in the relative donor lifetime ($\tau_{(D/A)/D(0)}$, i.e., the ratio of the fluorescence lifetime of the donor in the presence and absence of acceptor, respectively) versus $E$ plot (Fig. 1D). The uOmpXaq population, however, did not fall onto a line that describes the intrachain distance distribution of a Gaussian chain, used to model completely unfolded polypeptides (51) (Fig. 1D, dashed line). This substantiates our recurrence analysis (49), in that, uOmpXaq does not behave like a fully unfolded, random coil polypeptide. uOmpXaq thus exhibits a multitracer dynamical character with fast and slow intrachain motions, as has been previously observed for other proteins and membrane proteins (52–54).

Interestingly, for the measurement conducted at 25 °C, at late time points (4 to 6 h), the $E$ population of uOmpXaq coincided with the static FRET line (SI Appendix, Fig. S1I, Left), indicating that uOmpXaq’s fast reconfiguration dynamics vanish over time. This points toward the formation of long-range interactions in the unfolded-state and a dedynamization of the uOmpXaq polypeptide chain while preserving the broad conformational heterogeneity. Such a behavior was absent in the case of the measurement performed at 37 °C (SI Appendix, Fig. S1I, Right) and was reversible upon temperature increase from 25 °C to 37 °C (SI Appendix, Fig. S1J), likely due to temperature-induced chain dynamics. Of note, in all experiments, an additional minor population (~10%) at a high FRET efficiency ($E \sim 0.92$) was observed. This very compact state is particularly observed after long measurement times of 4 to 6 h (SI Appendix, Fig. S1I) and likely constitutes a misfolded state, denoted as uOmpXcompact throughout the article (SI Appendix, Fig. S2A).

In conclusion, we find that uOmpXaq adopts a broad, heterogeneous ensemble of structures and populates a rugged energy landscape leading to a multitracer dynamical behavior with both fast and submillisecond chain reconfiguration dynamics and slow conformational interconversion. An overall expansion of the uOmpXaq polypeptide chain is observed, in line with a conformational loosening mechanism that likely serves to alleviate uOMP aggregation (40).

**Skp- and SurA-Bound uOmpX Is Dynamically Expanded.** Next, we probed the effects that the chaperones Skp and SurA have on the structural and dynamic properties of the uOmpX polypeptide chain. To this end, we subjected denatured OmpX to a multistep dilution and denaturation protocol to transfer the protein into native-like buffer in the presence of the chaperones Skp or SurA and subsequently recorded protein into native-like buffer in the presence of the chaperone and appeared structurally distinct and more expanded than the uOmpXaq and uOmpXcompact states.

To extract populations from the FRET efficiency histograms, we found that three Gaussian distributions were sufficient to describe the overall shape and change of shape upon chaperone titration (see SI Appendix, Fig. S3 and Methods). Gaussian fitting showed that the peak positions of the two populations in the high FRET regime exhibited $E$ values of $E \sim 0.75$ and $E \sim 0.92$, which are in very good agreement with the $E$ values of the uOmpXaq ($E \sim 0.77$) and uOmpXcompact ($E \sim 0.92$) states. This suggests that the presence of Skp does not affect the conformational properties of these states and implies that these do not directly interact with Skp. We therefore identify these states, even in presence of chaperones, as uOmpXaq and uOmpXcompact (SI Appendix, Fig. S2B).

The FRET efficiency peak of Skp-bound uOmpX was centered around $E_{\text{Skp3-uOmpX}} \sim 0.45$, a considerably larger interferophore distance than the unbound uOmpXX (i.e., uOmpXaq) state. A careful analysis of the donor lifetime, the fluorophore anisotropies, and the rotational correlation times of the fluorophores ensured that the lower $E$ values reflect a global conformational change arising from the increased end-to-end distance of the uOmpX polypeptide chain upon binding to Skp (SI Appendix, Table S1). Moreover, Gaussian fitting revealed that, at [Skp] = 2.5 μM, ~80% of uOmpX molecules were in a Skp3–uOmpX complexed state, suggesting that the presence of Skp shifts the equilibrium from the unbound to the bound form.

In a second set of experiments, we diluted uOmpX in the presence of the chaperone SurA at 37 °C. Upon addition of SurA, we observed, similarly to Skp3, an additional FRET efficiency peak at lower $E$ values (Fig. 2C). Increasing [SurA] from 46 nM to 25 μM lead to an increase of the low $E$ state population (i.e., uSurA–uOmpX). Gaussian fitting allowed us to identify the peak position and fraction of SurA-bound uOmpX with $F_{\text{uSurA-uOmpX}} \sim 0.6$ at a concentration of 25 μM, which is close to the intracellular concentration of SurA (18, 58). At this concentration, ~87% of uOmpX molecules are bound to the chaperone. The lower $E$ value of uSurA–uOmpX also indicated an expanded conformation as compared to uOmpXaq and yet a less expanded conformation as compared to Skp3–SurA. The distinct peak positions of the two chaperone-bound states ($F_{\text{Skp3-uOmpX}} \sim 0.45$ and $F_{\text{SurA-uOmpX}} \sim 0.6$) thus suggest that they themselves occupy a distinguishable configurational space likely due to the difference in the interaction mechanisms between the chaperone and its substrate (7, 16).

Since temperature affected the conformation and reconfiguration dynamics of uOmpXaq, we asked how temperature influences the structural ensemble and dynamics of chaperone-complexed uOmpX. To this end, we performed titration experiments at 31 °C and 25 °C and evaluated $F_{\text{Skp3-uOmpX}}$ and $F_{\text{SurA-uOmpX}}$ at different [Skp3] and [SurA] (SI Appendix, Fig. S4 and Table S3). The FRET efficiencies of the unbound (uOmpXaq) and compact (uOmpXcompact) states did not vary markedly across the chaperone concentration series performed at different temperatures (SI Appendix, Figs. S5 and S2 A–C). For the chaperone-bound fraction of uOmpX, $F_{\text{Skp3-uOmpX}}$ remained constant, within error, across nearly three orders of magnitude of [Skp3] at 25 °C (Fig. 2D, Left). This suggests a concentration-independent, possibly 1:1 stoichiometry, of Skp3 and uOmpX, in accordance with previous studies (32, 59). Increasing the temperature of the complex, however, shifted $F_{\text{Skp3-uOmpX}}$ to higher $E$ values (Fig. 2D), while at each temperature, $F_{\text{Skp3-uOmpX}}$ itself remained
constant again across all concentrations. We infer that, at elevated temperatures, uOmpX in complex with the chaperones appears to form a more compact structural ensemble, possibly due to the increased dynamics of uOmpX in the Skp3 cavity at 37 °C as compared to 25 °C, similar to recent observations made by NMR spectroscopy (32). By contrast, for SurA–uOmpX complexes, $E_{\text{SurA–uOmpX}}$ decreased with increasing chaperone concentrations at 25 °C (Fig. 2E, SI Appendix, Table S3). Interestingly, at elevated temperatures, the dependency on [SurA] was enhanced and we observed a strong decrease of $E_{\text{SurA–uOmpX}}$ with increasing [SurA] at 37 °C from $E_{\text{SurA–uOmpX}}$ (1.2 μM) ∼ 0.66 to $E_{\text{SurA–uOmpX}}$ (25 μM) ∼ 0.55. This suggests that uOmpX is more expanded at elevated [SurA], which can be explained by either a different interaction mode or that uOmpX is sequestered by multiple SurAs as also suggested by previous studies (31, 33).

Next, we modeled the structural ensemble of the states globally using a log-normal distribution and obtained a standard deviation (SD) of the donor–acceptor distance ($\sigma_R$) and the expected interdye distance ($R_{\text{interdy}}$) (Fig. 2F, inset; Methods). A measure of heterogeneity among a population of varying end-to-end distance is the coefficient of variance, $\text{CV} = \sqrt{\langle \sigma^2 \rangle}$, where a larger CV indicates an increased heterogeneity. We determined CVs of 0.204 ± 0.036 and 0.304 ± 0.022 for Skp3-bound and SurA-bound uOmpX, respectively. Compared to SurA–uOmpX, the CV of the Skp3–uOmpX complex is reduced and indicates less conformational heterogeneity for the bound substrate, likely due to the binding of uOmpX in the cavity of Skp3 (55, 56). By contrast, the higher CV for the SurA–uOmpX complex marks an increased heterogeneity. This increase in heterogeneity could originate from the transient multisite binding of SurA resulting in diverse unstructured and expanded uOmpX conformations with none, one, or even multiple SurAs bound. We further used the log-normal distribution to extract interdye distances in order to quantify the end-to-end distances of the chaperone-bound uOMP complexes. Compared to $R_{\text{interdy}}$, the interdye distance grows from $R_{\text{interdy}}$ at 1.2 μM ∼ 4.6 nm to $R_{\text{interdy}}$ at 25 μM ∼ 5.8 nm at 25 °C and to $R_{\text{interdy}}$ at 12 μM ∼ 5.4 nm at 25 °C.

In a last step, to gain further insights into the structural and dynamical properties of chaperone-bound uOmpX, we analyzed the underlying dynamics of the chaperone-complexed uOmpX polypeptide chain. To this end, we extracted single-molecule events of the chaperone-bound uOMP complex using a combined FRET efficiency and relative donor fluorescence lifetime filter (SI Appendix, Fig. S3). We determined the peak positions for $E$ and $\tau_{\text{D(A)}}$ for each condition and plotted the values in a two-dimensional (2D) diagram (Fig. 2F and SI Appendix, Tables S3 and S4). The bound FRET efficiencies ($E_{\text{Skp3–uOmpX}}$ and $E_{\text{SurA–uOmpX}}$) across different chaperone concentrations were offset in distinct clusters from the static FRET line. The offset indicates that the polypeptide chain undergoes fast configurational changes on the submillisecond timescale while being bound to chaperones.

To explore the timescale of dynamics further, we implemented species-filtered two-dimensional fluorescence lifetime correlation spectroscopy (2D FLCS) (60, 61) and extracted characteristic chain reconfiguration times. Briefly, in a first step, photon pairs of selected FRET efficiency-filtered fluorescence bursts with a time gap matching the time interval ($\Delta T$) and a window size of $2 \times \Delta \Delta T$ (Fig. 3A) were sorted according to the microtime of the initial
Fig. 3. Conformational dynamics of the Skp3- and SurA-bound uOmpX. (A) Representation of species-filtered 2D FLCS using SurA–uOmpX complexes as an example. Here, \( t \) is the microtime of the emitted photons, \( \Delta T \) is the time interval, and \( \Delta \Delta T \) is the window size. Emission-delay histograms corresponding to uOmpXaq (B), Skp3–uOmpX (C), and SurA–uOmpX (D). The legend indicates the short and long initial microtimes.

### PNAS Chamachi et al. 2022
Chaperones Skp and SurA dynamically expand unfolded OmpX and synergistically disassemble oligomeric aggregates

and final photon \((t_1 \text{ and } t_2, \text{ respectively})\) in the 2D emission-delay histogram (SI Appendix, Fig. S6). The timescale of interconversion dynamics between different molecular states were then extracted by comparing slices of the 2D histogram with either short or long initial microtimes (blue and red, respectively, in the center panel of Fig. 3A and in the histograms of Fig. 3 B–D). If the chosen \( \Delta T \) matches the time of a conformational change, the emission-delay histograms for two initial microtime windows will start to separate. For uOmpXaq, all delay time windows \( \Delta T \pm \Delta \Delta T \) produced separated emission-delay histograms (Fig. 3B). The amplitude of separation was greater at a lower time point \( \Delta T = 5 \mu s \) as compared to that at higher time points of 5 \( \mu s \) or 240 \( \mu s \). This reinforced our finding that uOmpXaq exhibits both fast and slow conformational dynamics by virtue of its rough energy landscape. For both Skp3–uOmpX (Fig. 3C) and SurA–uOmpX (Fig. 3D) complexes, by contrast, the emission-delay histograms started separating at lower time points \( \Delta T = 2.5 \mu s \) and \( \Delta T = 6 \mu s \), respectively. Skp- or SurA-bound uOmpX thus shows a dominating microsecond conformational change, suggesting that both chaperones modulate the energy landscape of uOmpX such that the energy barriers between the subpopulation of their bound substrate are reduced, leading to a dynamization of the polypeptide chain in the chaperone-bound state.

To conclude, we found that binding of the chaperones Skp3 and SurA leads to an increased end-to-end distance of the uOmpX polypeptide chain. Skp3-complexed uOmpX showed a decreased conformational heterogeneity, while binding of SurA increased the conformational heterogeneity of uOmpX, indicating very distinct interaction mechanisms of the chaperones on their substrates. Complexed uOmpX exhibited fast chain reconfiguration dynamics on timescales <10 \( \mu s \), unlike uOmpXaq alone, which showed multitier dynamics on a large range of timescales (submillisecond to ≥100 ms) originating from its rugged energy landscape. The fast dynamics and the possible expansion of chaperone-bound uOmpX may aid in preventing uOmpX from getting trapped in a fortuitous conformation which might be folding incompetent or inaccessible to the BAM complex. A wealth of different conformations also indicates a small entropic penalty for complexing uOmpX to Skp3 or SurA.

### Entropic and Enthalpic Contributions of Chaperone–uOmpX Interactions

We next set out to obtain insights into the energetic contributions of Skp and SurA chaperone–uOmpX interactions. Interaction isotherms at different temperatures allow for the extraction of enthalpic, \( \Delta H \), and entropic, \( \Delta S \), contributions to the free energy change of interaction, \( \Delta G \), which are connected to the association constant, \( K_a \), for a bimolecular association reaction via:

\[
K_a(T) = \exp\left(\frac{-\Delta H}{R \cdot T} + \frac{\Delta S}{R}\right) \tag{1}
\]

In order to extract entropic and enthalpic contributions, we developed a global analysis approach that allows the reconstruction of FRET efficiency distributions at different temperatures and chaperone concentrations (Fig. 4 A and Methods).
Briefly, by varying the optimization parameters $\Delta H$ and $\Delta S$ and considering the fraction of the compact state ($f_{c}(T)$), which is likely unable to be chaperone complexed, as well as the maximum fraction of the bound state ($f_{b,\text{max}}(T)$) and minimizing the reduced $\chi^{2}$, the best estimators for $\Delta H$ and $\Delta S$ were obtained. Here, we chose a Hill coefficient of 1 for Skp3 and 1.5 for SurA, based on earlier reports (32, 33, 59, 62). We further assumed that Skp3 remained trimeric under our experimental conditions (SI Appendix, Methods). As $c_{1/2}$ (inverse of $K_{D}$) is dependent on temperature, at 37°C, we obtained a $c_{1/2}$-value of 359 ± 0.1 nM and 407 ± 0.1 nM for Skp3–uOmpX and SurA–uOmpX, respectively. This would convert to dissociation constants, $K_{D} \sim 359$ nM and $K_{D} \sim 82$ μM for Skp3–uOmpX and SurA–uOmpX, respectively. At 25°C, we obtained a $c_{1/2}$-value of 3.554 ± 0.001 nM and 102.89 ± 0.02 nM for Skp3–uOmpX and SurA–uOmpX, respectively. This would convert to $K_{D} \sim 3.6$ nM and $K_{D} \sim 1.0$ μM for Skp3–uOmpX and SurA–uOmpX, respectively, which is in very good agreement with previously reported values for the chaperone–OMP interactions (33, 36, 62). More interestingly, for Skp3–uOmpX complex formation, the binding enthalpy and entropy changes of the interaction were $\Delta H(Skp3–uOmpX) = -298 \pm 19$ kJ mol$^{-1}$ and $\Delta S(Skp3–uOmpX) = -0.84 \pm 0.06$ kJ mol$^{-1}$ K$^{-1}$ (Fig. 4B), respectively, yielding a $\Delta G(Skp3–uOmpX) = (-38.3 \pm 0.2)$ kJ mol$^{-1}$ at 37°C. The change of enthalpy upon binding is large, which is in agreement with recent findings demonstrating a rich interaction of the uOmpX polypeptide chain with the cavity formed by Skp3 (32). At the same time, the change in interaction entropy is strongly negative, suggesting a stark reduction of the overall configurational space for uOmpX upon its encapsulation by Skp3. Hence, Skp3 binding to uOmpX is entropically unfavorable and yet this entropic penalty is counterweighted by a large enthalpic contribution, enabling the binding of Skp3 to the client OMP.

For SurA–uOmpX complex formation, the binding enthalpy and entropy changes of interaction were $\Delta H(SurA–uOmpX) = -132 \pm 24$ kJ mol$^{-1}$ and $\Delta S(SurA–uOmpX) = -0.24 \pm 0.08$ kJ mol$^{-1}$ K$^{-1}$ (Fig. 4B), respectively, yielding an overall stabilization of the complex of $\Delta G(SurA–uOmpX) = (-57 \pm 1)$ kJ mol$^{-1}$ at 37°C, which is slightly more favorable than that for Skp3. The change of enthalpy was twofold lower than that for Skp3, likely due to more transient binding as suggested in recent studies (31, 33). At the same time $\Delta S$ upon SurA binding was threefold lower than with Skp3 and close to zero, indicating only a minimal change of configurational space for the binding of SurA to uOmpX. Although our observed interaction enthalpy and entropy are composed of multiple components (SI Appendix, Methods), they agree well with our finding of an increased coefficient of variation (i.e., structural heterogeneity) for SurA–uOmpX as compared to Skp3–uOmpX. Thus, binding of SurA to its client uOmpX is markedly different in nature than Skp3 binding to uOmpX. The reduction of conformational freedom of uOmpX in the cavity of Skp3 needs to be compensated by an increased interaction enthalpy (Fig. 4C). By contrast, uOmpX binding to the surface of SurA reduces the conformational freedom to a lesser extent and thus a lower enthalpic contribution is sufficient to enable binding (Fig. 4C).

**SurA and Skp3 Act Synergistically as Disaggregases on Oligomeric OmpX Structures.** Aside from the known uOMP holdase activities of Skp and SurA, recent work has suggested that Skp also has the ability to act as a disaggregase to disassemble oligomeric OMP structures (38). By contrast, uOmpX binding to the surface of SurA reduces the conformational freedom to a lesser extent and thus a lower enthalpic contribution is sufficient to enable binding (Fig. 4C).

To address this question, we investigated the action of Skp3 and SurA toward aggregated OmpX ($\text{OmpX}_{\text{Agg}}$) and devised...
an FCS-based assay that allows us to probe the disaggregase efficacy of the two chaperones. To this extent, we diluted labeled and denatured uOmpX at pM concentrations with unlabeled and denatured uOmpX in buffer to yield a final OmpX concentration of 1 μM (Fig. 5A). The residual GdmCl and LDAO concentrations were 12 mM and >200 nM, respectively. It is important to note, that due to the 100,000-fold concentration difference of labeled and unlabeled uOmpX, only a small fraction of potential aggregates will be fluorescently marked and thus visible to FCS measurements. We incubated the mixture for 10 min, subsequently performed FCS measurements at 37 °C, and obtained a multimodal autocorrelation function with two major components suggesting the presence of a fast and a slowly diffusing species (Fig. 5B, green). A comparison with an FCS curve recorded from a highly diluted sample of only labeled uOmpXaq (~10 pM) indicates that the fast-diffusing component in the correlation curve of the aggregation sample stems from monomeric uOmpXaq, with a characteristic diffusion time of $\tau_{\text{Diff}} = (0.27 \pm 0.01)$ ms (Fig. 5B, gray; SI Appendix, Table S6 and Fig. S7A). The second, slowly diffusing species in the OmpXAgg measurements showed a $\tau_{\text{Diff}} = (31.6 \pm 1.3)$ ms (SI Appendix, Fig. S7B), which is about two orders of magnitude slower than that of single uOmpXaq molecules. Notably, OmpXAgg is possibly an ensemble of differently sized aggregates; hence, values reported here reflect average values. We determined that in our aggregation mixture, about 24 ± 2% of the labeled uOmpX molecules are part of aggregates (termed apparent fraction; Fig. 4D). Interestingly, at lower temperatures, the propensity of OmpX aggregation was reduced (Fig. 5B).

Next, we studied the effect that the chaperones Skp and SurA have on OmpX aggregation. To this end, we first incubated the mixture containing 1 μM unlabeled uOmpX and 10 pM labeled uOmpX for 10 min and then added either 2.5 μM Skp3 or 5.8 μM SurA to the mixture. We recorded FCS curves at 37 °C directly after the dilution and obtained multimodal curves, indicating the presence of more than two components in solution. In addition to the fast-diffusing species with $\tau_{\text{Diff}} = (0.27 \pm 0.01)$ ms reflecting free uOmpXaq and the aggregated OmpXAgg species with very high diffusion times ($\tau_{\text{Diff}} \sim (32 \pm 1)$ ms), we expected to also observe chaperone-complexed uOmpX species with intermediate diffusion times. By comparison with an FCS curve recorded from a highly dilute sample of labeled uOmpXaq (~10 pM), we assigned diffusion times of $\tau_{\text{Diff}} = (1.79 \pm 0.05)$ ms and $\tau_{\text{Diff}} = (1.38 \pm 0.03)$ ms to Skp3- and SurA-bound monomeric uOmpX, respectively (SI Appendix, Fig. S7B and C). To quantify the fractional amounts of the three components (i.e., uOmpXaq, chaperone-bound monomeric uOmpX, and OmpXAgg), we fitted the autocorrelation data of our mixture experiment with three components. Here, the respective diffusion times were kept constant to extract the amplitudes of each component as they are inversely proportional to the concentrations and fractional amounts of the species. The addition of Skp3 to the aggregation mixture reduced the apparent fraction of OmpXAgg to 20 ± 2% and the addition of SurA even to 13 ± 1%. A reduction of apparent OmpXAgg was only observed at chaperone concentrations above 1 μM (SI Appendix, Fig. S7F).

Interestingly, in both cases, the disaggregation of OmpX was found to be accompanied by an increase in the free uOmpXaq population (SI Appendix, Fig. S7G) and only negligibly in the chaperone-uOmpX complex. Assuming uniform mixing of the labeled and unlabeled uOmpX sample, the increase of the free uOmpX population suggests that the chaperones show a higher affinity toward OmpXAgg as compared to uOmpX. This finding is significant, as it suggests...
that the careful adaptation of affinities is crucial during physiological stress conditions. In such a scenario, disaggregation activity might be more important than holdase activity to prevent the formation of large toxic OMP aggregates in the periplasm.

Finally, we asked the question whether there are synergistic effects on the disaggregation reaction, given that both chaperones are present in the periplasm. We performed our disaggregation assay in which both chaperones were added simultaneously to the mixture containing OmpXAgg and labeled uOmpX at pH concentrations after a 10-min incubation at 37 °C. Strikingly, the fraction of OmpXAgg was reduced to ∼5% (Fig. 5D). Taking into account our single-chaperone measurements, for an independent, additive reduction of the apparent OmpXAgg, we would have expected a ∼9%. The increased disaggregation by a factor of two for both chaperones together suggests therefore a cooperative action of Skp3 and SurA. Furthermore, the additional presence of the chaperone-bound uOmpX fraction (∼8%) in the presence of both Skp and SurA could originate from handing over of uOmpX between the chaperones or from a dynamic adaptation of the chaperone functionalities between disaggregation and chaperoning activities.

Conclusions

Periplasmic chaperones Skp and SurA are essential players in OMP biogenesis. They act as holdases, preventing uOMP substrates from misfolding and exhibit disaggregase functionalities, aiding in the clearance of OMP aggregates (32, 38). Understanding the molecular action mechanisms of these chaperones is fundamental to unraveling the role that these chaperones play during OMP biogenesis. Here, we have provided an intimate view on the structural, dynamic, and thermodynamic aspects of uOmpX in complex with Skp and SurA and explored their disaggregase activities.

Using smFRET, we have characterized the structural dynamics of chaperone-free and chaperone-bound uOmpX under near-native conditions and show that both chaperones Skp and SurA expand uOmpX upon binding. While previous reports have observed an expansion for SurA-bound OMPs (31, 38), an expansion upon Skp binding of an OMP in its cavity has not been observed. More surprisingly, the Skp3-bound uOmpX state exhibits a greater end-to-end distance as compared to the SurA-uOmpX state over the concentration range tested, suggesting that uOmpX interacts with Skp3 at multiple sites by spreading across the inner surface of the chaperone cavity and appears conformationally expanded within the chaperone cavity (Fig. 6A). The expansion of uOmpX upon Skp3 binding was independent of the [Skp3], indicating that the valency of Skp3 likely does not change as expected for the small eight β-stranded OmpX (59). The SurA–uOmpX complex (Fig. 6B), by contrast, showed an increased expansion with increasing [SurA], which is in line with multiple SurAs binding to one uOmpX. The higher valency was previously already suggested with a Hill-coefficient of n = 1.5 for the SurA–uOmpX association (33). Interestingly, chaperone-induced expansion was also observed for the ATP-dependent DnaJ–DnaK system for cytosolic proteins as clients (63), by forming a chain of DnaK molecules on the denatured substrate. However, unlike for Skp3 and SurA, for DnaK, the dynamics of interaction were largely dominated by ATP hydrolysis. Mechanistically, an expansion of OMPs by either Skp or SurA likely prevents intrachain contacts and, by extension, misfolding of uOmpX in the aqueous periplasmic space. Additionally, chain expansion can aid in exposing the β-signal peptide sequence, which facilitates recognition by BAM (64, 65). Hence, our results point to a general mechanism of chain expansion, which in the case of Skp and SurA, enables OMP chains to defer misfolding even in the absence of a high-energy substrate.

We further quantified the dynamics of both Skp3- and SurA-bound uOmpX using 2D-FLCS. We demonstrate that the intrachain dynamics of the chaperone-complexed uOmpX chain are on a timescale of 5 to 10 μs (Fig. 6 A and B), whereby the fast chain reconfiguration dynamics are possibly facilitated through local transient interactions with the chaperones. This is in contrast to the multier (i.e., both slow conformational and fast reconfiguration chain) dynamics observed for the aqueous uOmpX substrate (Fig. 6C). Periplasmic chaperones presumably help to lower the energy barrier and, by extension, misfolding of uOmpX into the unfolded state. We hypothesize that such a manipulation and dynamization of the energy landscape of the OMP substrate might be beneficial for BAM interactions (64). Notably, transient binding interactions and multivalency interactions were also observed for the cytoplasmic chaperone Trigger Factor (TF), which is structurally homologous to SurA, and like the latter is also involved in ATP-independent transfer of clients to the downstream chaperone machinery for folding (66).

Given that the periplasm is deficient of ATP as an energy source, protein folding and chaperoning, involving binding to the client uOMPs, have to be driven entirely by a change in the free energy of the system. An analysis of binding isotherms at different temperatures revealed that the change of entropy upon binding to the client uOmpX substrate was low for Skp3 and close to zero for SurA, suggesting that the conformational freedom is affected only minimally. A low entropic change upon binding has been hypothesized to be favorable for locally transient chaperone–OMP interactions, due to a reduced enthalpy–entropy compensation (29). Strikingly, comparing Skp3 and SurA, we find that the chaperone–uOMP interactions are fine-tuned exquisitely, meaning a higher entropic interaction change with Skp3 is compensated by a higher interaction enthalpic term, or vice versa in the case for SurA, to ensure a favorable, yet low change of free energy upon binding (Fig. 6). Hence, the different contributions of enthalpy and entropy to both complexes suggest that the thermodynamics of

Fig. 6. Model of Skp3 and SurA chaperone action on uOmpX and OmpXAgg. (A) Skp3–uOmpX is expanded due to numerous intermolecular interactions between the chaperone and its substrate, while the substrate itself undergoes fast chain reconfiguration on timescales <5 μs. (B) The increased expansion of uOmpX upon incrementing [SurA] indicates that more than one molecule of SurA binds to uOmpX. Similar to Skp3, SurA expands its substrate and induces uOmpX chain reconfiguration on a timescale of <10 μs. Interactions of both chaperones with uOmpX are energetically calibrated through an exquisite entropy–enthalpy compensation, as indicated by the values of entropy and enthalpy change (∆H and ∆S, respectively), along with ∆G. (C) uOmpX appears both submillisecond chain reconfiguration dynamics and >100 ms timescale conformational changes in aqueous buffer. (D) Both chaperones can disassemble aggregated OmpX, which can emerge under stress conditions.

Chamachi et al.
Chaperones Skp and SurA dynamically expand unfolded OmpX and synergistically disassemble oligomeric aggregates
the Skp–uOmpX and SurA–uOmpX interaction are differently calibrated in accordance with their chaperoning mechanisms.

Lastly, we found that both chaperones have the ability to disassemble aggregated OmpX (Fig. 6D). Interestingly, both chaperones show a higher affinity toward aggregated OmpX than uOmpX. A higher affinity toward the aggregate might be particularly useful when aggregated OmpX emerges, for example, under stress conditions to avoid the accumulation of OMP aggregates. Surprisingly, we found that Skp3 and SurA act synergistically on uOmpX aggregates, suggesting additional thermodynamic aspects of the Skp–uOmpX and SurA–uOmpX interaction differ and SurA binding results in a reduced loss of entropy compared to Skp binding. In a biological context, expansion and fast chain reconfiguration of the bound polypeptide may serve to ensure that the uOMP remains accessible to the BAM complex. The enthalpy–entropy compensations consent with the binding modes of the two chaperones to maintain an energetically favorable interaction with uOmpX. Lastly, both chaperones synergistically disaggregate OmpX aggregates, suggesting additional interactions between both chaperones that enhance the disaggregation. We hypothesize that the tightly regulated multifaceted functionalities of both chaperones enable cellular vitality to be maintained and regulated under normal and stress conditions.

Methods

Protein Production. A tag-free double-cysteine variant of OmpX (OmpX1-14a) without a signal sequence was expressed as inclusion bodies in BL21 (DE3) cells and purified following standard procedures including anion exchange chromatography, as previously reported (69, 70). After refolding, the protein was labeled with FRET donor (ATT0532, Atto-Tec) and acceptor (Abberior STAR 635P, Abberior) dyes. We further produced two other OmpX variants; however, they were not used for the main study due to their misbehavior possibly caused by higher propensity toward aggregation (SI Appendix, Fig. S8). Details are given in SI Appendix.

Skp and SurA were produced as N-terminal hexahistidine (His6) fusion proteins in E. coli BL21 (DE3) cells and purified using immobilized metal affinity chromatography under denaturing conditions, as previously reported (32, 59). For FCS experiments involving OmpX aggregates, as shown in Fig. 5A, the concentration of GdmCl and LDAO was 12 mM and 175 mM, respectively, after addition of 1 μM uOmpX in the same buffer as that used for smFRET experiments. Details are given in SI Appendix.

Sample Preparation for Single-Molecule Measurements. A multistep dilution protocol, as shown in Fig. 1A, was followed to prepare samples for smFRET and FCS measurements. In both FCS and smFRET experiments, the labeled protein was present at a concentration of 20 μM in buffer containing 20 mM Tris HCl (pH 8.0) and 150 mM NaCl. The remaining concentration of GdmCl and LDAO was 6 mM and 87.5 mM, respectively. For FCS experiments involving OmpX aggregates, as shown in Fig. 5A, the concentration of GdmCl and LDAO was 12 mM and 175 mM, respectively, after addition of 1 μM uOmpX in the same buffer as that used for smFRET experiments. Details are given in SI Appendix.

Melting curves of labeled and unlabeled OmpX obtained through circular dichroism spectroscopy measurements overlapped within error indicating that labeling did not affect the protein stability markedly (SI Appendix, Fig. S9A).

smFRET and FCS Measurements. Experiments were carried out using a single-molecule confocal fluorescence microscope as previously described (71) and detailed in SI Appendix. Data analysis was performed with custom-written Matlab scripts (Mathworks). Single-molecule events were identified from the acquired photon stream by a burst search algorithm. Details about smFRET and FCS analysis procedures, including burst selection, data reduction, calculation of FRET efficiencies, and fluorescence, are given in SI Appendix.

Data Availability. Raw h3 data (time-correlated single-photon counting data) and preanalyzed data from our single-molecule FRET and FCS experiments data have been deposited in a publicly accessible database, OpARA (DOI: http://dx.doi.org/10.25532/OPARA-162).

Acknowledgments. We thank Sandro Keller (University of Graz) and all members of the Schlierf group for fruitful discussions. This work was supported by the Technische Universität Dresden institutional funds (M.S.), German Federal Ministry of Education and Research (Bundesministerium für Bildung und Forschung) BMBF with Grant 03Z2E11 and 03Z2E51 (M.S.), the Deutsche Forschungsgemeinschaft with SCHL1896/3-1 and SCHL1896/6-1 (M.S.), and Dresden International Graduate School for Biomedicine and Bioengineering (DIGS-BB) granted by the Deutsche Forschungsgemeinschaft (DFG) in the context of the Excellence Initiative to N.C. A.S. was supported by Joachim Herz Stiftung. G.K. acknowledges support by the European Research Council (ERC) under the European Union’s Horizon 2020 Framework Programme through the Marie Skłodowska-Curie Grant MicroSPARK (agreement no. 841460 (G.K.), the Herchel Smith Funds of the University of Cambridge, and the Wolfson College Junior Research Fellowship.

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