

Analysis of Reconstituted Tripartite Complex Supports Avidity-based Recruitment of Hsp70 by Substrate Bound J-domain Protein

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Abstract

Hsp70 are ubiquitous, versatile molecular chaperones that cyclically interact with substrate protein(s). The initial step requires synergistic interaction of a substrate and a J-domain protein (JDP) cochaperone, via its J-domain, with Hsp70 to stimulate hydrolysis of its bound ATP. This hydrolysis drives conformational changes in Hsp70 that stabilize substrate binding. However, because of the transient nature of substrate and JDP interactions, this key step is not well understood. Here we leverage a well characterized Hsp70 system specialized for iron-sulfur cluster biogenesis, which like many systems, has a JDP that binds substrate on its own. Utilizing an ATPase-deficient Hsp70 variant, we isolated a Hsp70-JDP-substrate tripartite complex. Complex formation and stability depended on residues previously identified as essential for bipartite interactions: JDP-substrate, Hsp70-substrate and J-domain-Hsp70. Computational docking based on the established J-domain-Hsp70(ATP) interaction placed the substrate close to its predicted position in the peptide-binding cleft, with the JDP having the same architecture as when in a bipartite complex with substrate. Together, our results indicate that the structurally rigid JDP-substrate complex recruits Hsp70(ATP) via precise positioning of J-domain and substrate at their respective interaction sites - resulting in functionally high affinity (i.e., avidity). The exceptionally high avidity observed for this specialized system may be unusual because of the rigid architecture of its JDP and the additional JDP-Hsp70 interaction site uncovered in this study. However, functionally important avidity driven by JDP-substrate interactions is likely sufficient to explain synergistic ATPase stimulation and efficient substrate trapping in many Hsp70 systems.

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Introduction

Via cyclic interaction with substrate proteins, Hsp70 molecular chaperones play key roles in general protein homeostasis, as well as many critical cellular pathways. Regardless of the particular substrate or cellular pathway, the substrate-Hsp70 interaction is driven by the same conformational changes of Hsp70 - induced by hydrolysis, reversed by ATP nucleotide exchange.^{1–4} In the ATP bound conformation Hsp70's peptide binding cleft, located in the β subdomain of the substrate binding domain (SBD β), is open - thus available for rapid binding of substrate, but also allowing its rapid release. ATP hydrolysis leads to trapping of the substrate - covering of the cleft by the "lid" (i.e., the SBD α subdomain), minimizing escape. In this ADP-conformation only the linker connects the nucleotide binding domain (NBD) and SBD, while in the ATP conformation, the linker and SBD β dock onto the NBD, as can SBD α . as well.

A key to this efficient nonequilibrium binding of substrates by Hsp70 is the timing of ATP hydrolysis such that the transiently interacting substrate is trapped in the cleft.⁵ However, though the presence of substrate in the cleft does result in an increase in ATPase activity, it is very modest. Efficient substrate trapping requires a J-domain protein (JDP) cochaperone.^{6,7} Together J-domains and substrates, synergistically stimulate Hsp70's ATPase activity.⁸⁻¹¹ J-domains interact at the NBD-linker-SBD β interface of Hsp70(ATP).¹² The invariant, His, Pro, Asp (HPD) motif critical for ATPase stimulation is in a loop following helix II, whose residues are also critical for Hsp70 binding. As many JDPs bind proteins that are also Hsp70 substrates, this step of the Hsp70 substrate interaction cycle is often referred to as substrate delivery.³ However, experimental information about this step is scant.^{11,13-17} primarily because of the transient nature of the interactions - both between the Jdomain and Hsp70(ATP), and between substrate and Hsp70(ATP). To investigate this step in the cycle we have taken advantage of a specialized JDP/Hsp70 system in which both the JDP and the Hsp70 bind a single substrate ^{18,19} We reasoned that if delivery by JDPs to Hsp70 is a mechanistically important feature of JDP/Hsp70 systems a specialized one would be most likely to have evolved such characteristics, as it lacks constraints conferred by having multiple substrates.

The JDP/Hsp70 system we chose for analysis is in mitochondria of *Saccharomyces cerevisiae*, functioning in the biogenesis of iron-sulfur clusters (Fe-S), prosthetic groups critical for activity of many proteins.²⁰ The substrate of the JDP Hsc20 and the specialized Hsp70 Ssq1 is Isu1, a small (15 kDa), single domain scaffold protein on which clusters are synthesized, and then - upon interaction with Hsc20/Ssq1 - transferred onto recipient

proteins.¹⁸ Hsc20 is a simple, 21 kDa JDP. The 11 kDa C-terminal Isu1 binding domain (CTD) is juxtaposed to the J-domain forming a rigid Lshaped structure.²¹⁻²³ The Hsc20-Isu1 binding interface involves evolutionary conserved surfaceexposed hydrophobic and charged residues.²³ While Hsc20 functions as a specialized JDP in Fe-S biogenesis in bacteria and mitochondria, Ssq1 emerged as a Fe-S specialized Hsp70 in the S. *cerevisiae* lineage.²⁴ It is the result of a duplication of the multifunctional mitochondrial (mt)Hsp70 present in all mitochondria.²⁵ Ssq1 maintains structural and functional features of its ancestor, but evolved unusually high specificity toward its substrate Isu1 and JDP partner Hsc20.24,26,27 Ssg1 recognizes a single site on Isu1 that is distinct from the site of Hsc20 binding – LPPVK pentapeptide on a surface exposed loop.2

Here, building on our understanding of the bipartite interactions amongst JDP Hsc20, Hsp70 Ssq1 and substrate Isu1,^{23,29-32} we report the isolation and analysis of a tripartite Ssq1-Hsc20-Isu1 complex. Our results are consistent with precise positioning of the native substrate Isu1 via the rigid JDP – driving avidity based simultaneous J-domain and substrate interaction with Hsp70 for efficient substrate trapping.

Results

Tripartite Ssq1-Hsc20-Isu1 complex is dependent on established bipartite interaction sites

To test whether a tripartite Ssq1-Hsc20-Isu1 complex could be isolated we took advantage of well chracterized Hsp70 mutational variants, in which an evolutionary conserved Thr residue involved in ATP hydrolysis is replaced by Ala such variants bind ATP but their ATPase is strongly reduced.³³ We carried out pulldown assays using Ssq1^{T239A} (henceforth called Ssq1*), which, as expected, is defective in ATP hydrolysis (Figure S1). Isu1 tagged at the C-terminus with glutathione-S-transferase (Isu1^{GST}), allowing it to be pulled down by glutathione resin, was used (Figure 1(A)).³⁴ As expected, little Ssq1* was pulled down when added to Isu1^{GST} alone. However, when Hsc20 was included, Ssq1*, as well as Hsc20, were pulled down (Figures 1(B) and S2 (A)). When wild-type Ssq1 rather than Ssq1* was used, it was pulled down, but Hsc20 was not - consistent with the ability of catalytic amounts of Hsc20 to drive stable binding of Isu1 by Ssg1 after ATP hydrolysis (Figure S2(C)).²⁵

To preliminarily assess the dependence of Ssq1^{*} association with Isu1^{GST} on Hsc20 concentration, the amount of Hsc20 was titrated, while holding Ssq1^{*} and Isu1^{GST} concentrations fixed. Proportionally increasing amounts of Ssq1^{*} (as well as Hsc20) were pulled down, until saturation was reached when Hsc20 reached the



Figure 1. Reconstitution of tripartite Ssq1*-Hsc20-Isu1 complex. (A-C) To assess tripartite complex formation pulldown assays were performed. Isu1^{GST} (2.5 μ M) was incubated in the presence of Hsc20 and/or Ssq1* (2.5 μ M and 5 µM, respectively unless otherwise indicated). Glutathione resin was added to pull-down GST and associated proteins, which were then separated by SDS-PAGE and stained with Coomassie blue (entire gels and controls in indicated Supplemental Figures). Molecular weight markers in kDa on left. *****; Isu1^{GST} degradation product. Amounts of pulled down Ssg1 were quantified by densitometry and corrected for background binding to GST alone for three independent experiments, Error bars represent SD, Maximum binding set at 100%, (A) Schematic of basic pulldown assay. (B) Dependence of Ssg1* pulldown on presence of Hsc20. Representative gel at top, quantitation at bottom (for details see Supplmental Materials and Methods). (left) in presence and absence of Hsc20. (right) with increasing concentrations of Hsc20 (0.0, 0.25, 0.5, 1.5, 2.5, 5.0 µM). Entire gels and controls, Figure S2. (C) Disruption of individual protein-protein interactions affects formation of tripartite complex. +, WT protein; see Table 1 for key to residues of indicated substitutions and Figure S3 and S4 for gels. (D) Computational model of the tripartite Ssg1 (ATP)-Hsc20-Isu1 complex, generated by docking model of Isu1 to the previously published³² model of Hsc20-Ssq1 complex and refining the obtained dominant structural state by all-atom MD simulations (8.6 µs). (left) Surface representation shows overall architecture of the complex: Ssq1 in the ATP bound conformation with SBD β and SBD α (dark green) docked to NBD (light green) and interdomain linker (yellow) placed inside the NBD. Hsc20 (cyan) interacts with Ssq1 via J-domain (JD) at the NBD/SBD β , linker interface and with Isu1 via C-terminal domain (CTD). (right) cartoon representation of Hsc20 and Isu1 in the tripartite complex show Hsc20-Isu1 and Ssq1-Isu1 interfaces; L63, V72 and F94 of Isu1 in contact with L105, L109 and Y163 of Hsc20 and LPPVK (PVK) of Isu1 in proximity to the substrate binding cleft of SBD β . (Hsc20-Ssq1 interface, Figure S5(A).

concentration of Ssq1 in the assay (Figures 1(B) and S2(B)), consistent with a stable tripartite Ssq1*-Hsc20-Isu1^{GST} complex. To ensure that complex formation does not depend of the presence of the GST tag on Isu1 we carried out pull-down experiments using Hsc20 or Ssq1* tagged by GST at the C- and N-termini, respectively. In both cases the untagged components were pulled down as efficiently as when Isu1 was tagged (Figure S3).

We then asked if amino acid substitutions known to disrupt individual protein-protein interactions (Table 1) affect the efficiency of Ssq1* pulldown by Isu1^{GST}. When Hsc20 was replaced by a variant having substitutions of residues at the Hsc20-Isu1 binding site (Hsc20^{LLY}),²³ the amount of Ssq1*pulled down was reduced by ~70% (Figures 1(C), and S4(A)). To test the effect of disruption of the interaction of Isu1 in the peptide binding cleft of Ssq1*, two variants were used – Isu1^{GST} having a triple alanine substitution in the LPPVK Ssq1* binding site (Isu1^{GST-PVK}) and Ssq1* having the key phenylalanine in the substrate binding cleft changed to serine (Ssq1*^{-F462S}) (Figures 1(C) and S4(B) and (C)).^{29,35} In both cases Hsc20 was pulled down efficiently, but less than 10% of Ssq1* was

Protein	Variant	Location	Affected interaction		
Ssq1	F462S	substrate binding cleft	Isu1 (Ref. ³⁴)		
Hsc20	R37A	J-domain helix II	Ssq1 (Ref. ³²)		
	H48A	J-domain HPD	Ssq1 (Ref. ³¹)		
	D50A	J-domain HPD	Ssq1 (Ref. ³¹)		
	LLY (L105A, L109A, Y163A)	CTD	Isu1 (Ref. ²³)		
	K132A	CTD	Ssq1 (this manuscript)		
	K172A	CTD	Ssq1 (this manuscript)		
	KK (K132A, K172A)	CTD	Ssq1 (this manuscript)		
lsu1	PVK (P135A, V136A, K137A)	LPPVK (Ssq1 substrate)	Ssq1 (Ref. ²⁹)		

	Table 1	Protein	variants	used that	have	disturbed	interaction	between	tripartite	e compl	ex subi	units.
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recovered with Isu1^{GST}. Importance of the Jdomain-Ssq1 interaction was assessed using alanine-substitutions in Hsc20's J-domain, two in the HPD (H48A and D50A) and one in Helix II (R37A) (Figures 1(C) and S5).^{31,32} As expected, Hsc20 was pulled down by Isu1^{GST}, but Ssq1* was reduced by ~70% in each case, indicating the importance of the J-domain-Ssq1 interaction. Taken together, the results support the idea that formation of a tripartite Ssq1*-Hsc20-Isu1 complex requires multivalent interactions between its constituents, as disruption of any of the established interactions interfere with its formation.

The dependence on Hsc20's known bivalent interaction sites for pulldown of Ssq1* by Isu1^{GST} motivated us to generate a structural model of the tripartite complex - using molecular docking and all-atom molecular dynamics (MD) simulations. Isu1 was docked to the previously characterized Hsc20-Ssq1(ATP) structural model in which the Jdomain positions at the conserved NBD-linker-SBD β interface of Ssq1.³² In the obtained dominant structural state - accounting for 54% of the structures, with no other states accounted for more than 8% - the J-domain binding interface was the same as in the initial Hsc20-Ssq1(ATP) complex (Figures 1(D) and S6(A)).³² Hsc20 also maintained its original, rigid structure.²³ Isu1 positioned having two contact points: (1) the known Hsc20-Isu1 binding interface; (2) the LPPVK of Isu1 located in proximity to the SBDβ substrate binding cleft of Ssq1 Figures 1(D) and S6(B).^{23,29,30} Furthermore, the spatial arrangement of Hsc20 and Isu1 in the tripartite complex closely resembled that in the modeled bipartite complex (Figure S6(C)). Overall, the modeling suggests that upon formation of the tripartite complex none of the components undergo major conformational changes.

Hydrogen-deuterium exchange analysis of tripartite complex

To obtain the tripartite complex in quantities suitable for biophysical analysis we co-expressed the three components in *Escherichia coli*. The major peaks obtained from size exclusion chromatography contained Ssq1, Hsc20, Isu1 (peak A) and Hsc20, Isu1 (peak B) (Figures 2 and S7). Native mass spectrometry (MS) revealed a molecular weights of 107.6 kDa for the tripartite complex and 35.8 kDa for the Hsc20-Isu1 complex consistent with a 1:1:1 and 1:1 stoichiometries.

To experimentally assess dynamic features of Ssq1*, alone and in the tripartite complex, we used hydrogen-deuterium exchange (HDX) in conjunction with MS,^{36,37} monitoring the exchange of main chain amide protons over time upon addition of deuterium (Figure 3(A)). Expected differences emerged from a global comparison of the 30 s exchange time point (Figures 3(B), (C) and S8(A)); for other time points see Figures S8(B) and S9(A) and (B). Compared to Ssg1* alone, deuterium uptake for most of the NBD, SBD β and interdomain linker of Ssg1 in the tripartite complex was slower - indicating conformational stabilization, as would be expected with binding of Hsc20's Jdomain at the NBD-linker-SBDB interface and LPPVK of Isu1 at the binding cleft of SBDB (Figures 3(B-D) and S8-S9). A different picture emerged from analysis of SBDa. First, the rates of exchange for its N- and C- terminal α-helices were the same for the tripartite complex and Ssg1* alone. Second, the exchange rate for the N-terminal helix was very rapid, consistent with this portion of the lid being highly dynamic, regardless of whether in the complex or not.

We also carried out HDX on Hsc20 and Isu1 alone, in complex with each other and in the tripartite complex. The exchange rates for Hsc20 alone (Figures S10 and S11) were consistent with its rigid structure. The junction region between the two domains displayed the slowest rates, which were retarded even further upon interaction with Isu1 and Ssq1*. As expected, rates of exchange for CTD and J-domain segments involved in interaction with Isu1 and Ssq1*, respectively, were also slower in the tripartite complex (Figure S11). In striking contrast to Ssq1* and Hsc20, very fast exchange rates were observed for lsu1 (Figure S12), consistent with its conformational previously flexibility reported in NMR experiments.^{38,39} Although this feature curtailed detailed analysis of HDX patterns, slower exchange



Figure 2. Isolation of the tripartite Ssq1*-Hsc20-Isu1 and bipartite Hsc20-Isu1 complexes from *E. coli* coexpressing their constituents. (left) Size exclusion chromatography of Ni-NTA purified complexes; peak A, tripartite and peak B, bipartite. For identification of proteins in peak A and B see Figure S7. (middle, right) Deconvoluted native mass spectrometry (MS) spectra of the tripartite (middle) and bipartite (right) complexes- predicted molecular weights are indicated. For the original native MS spectra see Figure S7.

rates were observed for Isu1 when in complex with Hsc20 and in the tripartite complex, consistent with structural stabilization.

Kinetics of tripartite complex formation

Equilibrium pull-down and HDX-MS experiments allow monitoring interaction of individual proteins and their conformational dynamics, but do not provide information about the kinetics of complex formation. Because a biologically meaningful tripartite complex must form rapidly, we analyzed the complex formation usina biolaver interferometry (BLI), which allows time resolved measurements. Isu1^{GST} was immobilized on anti-GST antibody-coated BLI sensors, which were then introduced into solutions containing ATP and one or more test proteins, that is the association phase (Figure 4(A)). When Hsc20 or Ssq1* was added individually the results were consistent with the pulldown experiments: Hsc20 was rapidly recruited to the sensor; no binding signal was observed for Ssg1* alone. In addition, consistent with efficient formation of a tripartite complex, the

binding signal of Ssq1* and Hsc20 together was higher than Hsc20 alone, with the magnitude of the difference being proportional to the amount of Hsc20 added (Figure S13). This signal increased rapidly, reaching 50% of maximum in less than 9 seconds. When the sensor was moved to a solution containing no protein (i.e., the dissociation phase), the signal decreased.

On closer inspection, two phases of binding were apparent- a rapid initial phase followed by slower binding. We therefore divided association into two steps, first one component, then both together to binding equilibrium of the first is ensure maintained (Figure 4(B)). When Isu1^{GST} loaded sensor was placed in Hsc20 solution (first step) rapid binding was observed, followed by slower binding when the sensor was moved to a mixture that contained Ssq1* as well (second step). We interpret this result as indicating rapid formation of the Hsc20-Isu1 complex (first step), followed by slower binding of Ssq1* to the Hsc20-Isu1 complex (second step). When, the order of addition was reversed, no binding of Ssq1* was observed in the first step, as expected, while

Figure 3. HDX-MS analysis of Ssq1 alone and in the tripartite Ssq1*-Hsc20-Isu1 complex. (A) Scheme of HDX-MS experiment. Proteins, alone or in complexes, were incubated in D₂O buffer allowing deuterium incorporation into the polypeptide backbone. After quenching exchange at a number of points in time and subsequent proteolysis, peptides were subjected to MS analysis to determine the increase in mass resulting from deuterium uptake. (B-C) Deuterium uptake of Ssq1* alone and in the tripartite Ssq1*-Hsc20-Isu1 complex. Data shown are for 30 s exchange time point. (B) Relative fractional deuterium uptake mapped on the structural model of Ssq1*(ATP). Nucleotide binding domain (NBD), substrate binding subdomains (SBD β , having substrate binding cleft; SBD α "lid"), interdomain linker (linker); N-terminal (α^{N}) and C-terminal (α^{C}) lid helices. D246, E253 – residues of the NBD interacting with Hsc20; F462 residue of the substrate binding cleft of SBD β interacting with LPPVK of Isu1. (C) Difference in deuterium uptake between Ssq1* alone and in the tripartite complex. Horizontal lines indicate peptides observed; differences in deuterium uptake for a given peptide. (D) Kinetics of relative deuterium uptake into selected peptides of Ssq1* alone (green), tripartite complex. (black).





Figure 4. Kinetics of the tripartite complex formation. (A-D) BLI analysis of Isu1^{GST} interaction with Ssq1* and Hsc20, and their variants. BLI sensors were loaded with Isu1^{GST} or GST (background control). Association: loaded sensors were inserted into solutions containing Ssq1* and/or Hsc20. If two step was performed (B-D), 10 min later sensors were moved to a solution containing indicated combination of Ssq1* and/or Hsc20 for an additional 10 min. Dissociation phase was initiated by placing sensors into solutions without proteins. (A) One step association analysis of Ssq1* and Ssq1*/Hsc20 using Isu1^{GST} and, as a control, GST. (B) Two step analysis of Isu1^{GST} interaction with Ssq1* and Hsc20, assessing effect of order of addition of Hsc20 and Ssq1*. (C) Two step analysis of interaction of Isu1^{GST} with Ssq1* and indicated variants defective in Ssq1* interaction: Hsc20 (R37A or D50A); Isu1^{LLY} (LLY). (D) Two step analysis of Isu1^{GST} interaction with Hsc20 and Ssq1* variant (F426S) defective in interaction with Isu1.

efficient formation of the tripartite complex was observed in the second step when Hsc20 was also present.

Next, we tested how disruption of bipartite protein-protein interactions affect the kinetics of tripartite complex formation (Figure 4(C) and (D)). Consistent with our pulldown results: (1) no signal above background was observed for an Hsc20 variant defective in Isu1 interaction; (2) no signal above that observed for Hsc20 alone for variants defective in J-domain-Ssq1 or Ssq1-Isu1 interaction. We conclude that the tripartite

complex forms rapidly, but disruption of any of the bipartite interactions between its constituents severely impedes its formation.

An additional interaction site between Hsc20 and Ssq1 stabilizes the tripartite complex

The surprising stability of the tripartite complex led us to reinspect the structural model. Positively charged K132 and K172 of Hsc20's CTD and negatively charged D364 of Ssq1's NBD (Figures 5(A) and S14) stood out as potentially forming an



additional interaction site. A peptide encompassing D364 had retarded deuterium uptake in the tripartite complex compared to Ssg1* alone (Figure 5(B)) supporting this idea. Therefore, we carried out pulldown and BLI experiments. When Hsc20 K132A and K172A variants were tested for tripartite complex formation the amount of Ssq1* pulled down with Isu1^{GST} was reduced by $\sim 40\%$ and \sim 60%, respectively (Figures 5(C) and S15). When the double variant was used, the amount of pulled down Ssq1* was reduced to the background level. In the BLI assay the signal for each single variant was substantially less than for wild-type (WT) Hsc20. When the double Hsc20 variant was used, no signal was detected above that observed when no Hsc20 was present (Figure 5(D)). We conclude that the Hsc20 CTD/ Ssg1 NBD interaction plays a role in formation of the Ssg1*-Hsc20-lsu1 tripartite complex and may in part explain its unusual stability.

To gain insight into the origin and evolution of the additional interaction site we first analyzed Ssq1, mtHsp70, and Hsc20 sequences from a wide variety of eukaryotes. Finding that positions homologous to D364 of Ssg1 and K132/K172 of Hsc20 are not universally conserved (Figure. S16 (A)), we focused on fungi closely related to S. cerevisiae, including both species in which Hsc20 functions with Ssq1 (i.e., post-duplication) and those in which Hsc20 functions with mtHsp70 (i.e., non-duplication) (Figure 5(E)). In non-duplication species positions 364 of Ssg1 and 132/172 of Hsc20 are not conserved. However, in postduplication species Ssg1s have mostly acidic amino acids (D, E) at position 364 and Hsc20s have mostly basic amino acids (K, R) at position 172. This pattern of conservation suggests that a new electrostatic interaction between an acidic residue at position 364 of Ssq1 and a basic residue at Hsc20 position 172 evolved in the last common ancestor of post-duplication species. Indeed, maximum likelihood phylogeny predicted, with 90% probability, that the common ancestor of Ssq1 had D364 while its Hsc20 partner had K172 (Figure S16(B) and (C)). The conservation pattern of Hsc20 position 132 in post-duplication species is more complex. It is occupied by basic amino acids in most species belonging to the Saccharomyces clade (Figure 5(E)), but is variable in the sister Candida clade. This pattern suggests that the strength of this new electrostatic interaction varies among Hsc20/Ssq1 systems – strong in *S. cerevisiae* and other species in which positions 172 and 132 are occupied by basic amino acids, but weaker in others.

Discussion

Here we report the biochemical isolation of a tripartite complex consisting of Hsp70 Ssg1, JDP Hsc20 and their substrate protein Isu1. Formation of this 1:1:1 complex requires not only the sites necessary for the subsequent hydrolysis of bound ATP that leads to substrate "trapping" (J-domain-Hsp70 and substrate-Hsp70), but also JDPsubstrate interaction sites. This dependence underscores the importance of avidity in driving Hsp70-substrate interaction cvcle the bv promoting simultaneous interaction of J-domain and substrate. Though the specialized system analyzed here evolved to be highly specialized, allowing isolation of the complex, it likely represents an initial step in the substrate binding cycle key to JDP/Hsp70 systems more generally.

While the dramatic effects on complex formation/ stability of substitutions at the binding sites make the importance of these interactions evident, the conformation of Ssq1* in the complex is less straightforward. As expected, the results of HDX point to the docking of the NBD and SBD β , a

Figure 5. Novel interaction between CTD of Hsc20 and NBD of Ssq1 is required for the tripartite complex formation. (A) Fragment of the structural model of the tripartite complex showing interaction between K132 and K172 of Hsc20's CTD and D364 of Ssg1's NBD- the novel interaction site. (B) Kinetics of relative deuterium uptake into the D364containing Ssq1* peptide (354–369). Ssq1* alone (green); tripartite complex (black). (C) (top) Isu1GST (2.5 µM) was incubated with Ssq1* (5 μM) and Hsc20 WT or K132A, K172A or double K132A/K172A (KK) variants (2.5 μM). Glutathione resin was added to pull-down GST and associated proteins, which were then separated by SDS-PAGE and stained with Coomassie blue; entire gel, loading controls and quantification in Figure S12. Molecular weight markers in kDa on left \star – marks the Isu1^{GST} degradation product. (bottom) Amounts of Ssq1* pulled down were guantified by densitometry and corrected for background binding to GST alone; Ssg1* levels for Isu1GST interacting with Hsc20 and Ssq1* were set at 100%. Error bars represent SD. (D) BLI analysis of Isu1^{GST} interaction with Ssq1* and Hsc20 variants defective in the novel Hsc20-Ssq1 interaction. Association-1, BLI sensors loaded with Isu1^{GST} were placed in solution containing Hsc20 WT or variants K132A, K172A or double K132A/K172A (KK) (1 μM); association-2, the sensors were placed in solution containing Ssq1* (1 μ M) and Hsc20 WT or variant (1 μ M). Dissociation, the sensors were placed in solution without proteins. (E) Sequence conservation of Hsc20 and Hsp70 positions involved in the novel Hsc20-Ssa1 interaction in S. cerevisiae across fungi post-duplication species (green branches) in which Hsc20 functions with Ssg1 and non-duplication species (black branches) in which Hsc20 functions with mtHsp70. Star indicates emergence of Ssq1 as result of mtHsp70 gene duplication.

requirement for J-domain interaction.^{32,40} The positioning of the SBD α lid is less clear, not only for the Ssq1*(ATP)-Hsc20-Isu1 complex discussed here, but for ATP-bound Hsp70s more generally. The relatively high rate of exchange revealed in the HDX experiments and the similarity of HDX data for Ssq1* in isolation and in the complex, suggests that the SBD α lid is undocked and dynamic in both. Thus, we conclude that the lid does not play a critical role in stabilization of the tripartite complex via interaction with Isu1. Although lid docking has been well established for some ATP-bound Hsp70s,⁴ undocked lids have been reported for others.^{41–43}

Our results are informative as to how synergy of ATPase stimulation by J-domain and substrate is accomplished in this specialized system, 25,44 as well as the role of avidity in forming this remarkably stable complex. The precision of the architecture positioning the J-domain and the substrate's Hsp70 binding site is striking. Hsc20 is a rigid molecule forming a structurally rigid complex with Isu1.23 When the J-domain of Hsc20 interacts with Ssg1* (ATP) via long distance electrostatic interactions, the bound Isu1 is positioned such that the LPPVK is in very close proximity to the substrate binding cleft of Ssq1*(ATP). This positioning results in extremely high local concentration, enabling avidity, that is functionally high affinity. The "fourth interaction", that between the CTD of Hsc20 and the NBD of Ssg1 illustrates the evolvability of JDP/Hsp70 systems. We think that this additional interaction. though weak, increases avidity and allows Hsc20/ Ssq1 system to function at the very low concentrations present in mitochondria. While mtHsp70 is one of the most abundant mitochondrial protein, Ssg1 functions at an approximately 1000-fold lower concentration.45 On the other hand, our evolutionary analysis shows that this interaction is restricted to S. cerevisiae and closely related species, suggesting that other Hsp20/Ssq1 and Hsc20/mtHsp70 systems function without this additional interaction. It should also be kept in mind that additional interaction(s) between JDP and its Hsp70 partner may not only increase avidity but further regulate the timing of hydrolysis.46-

It is easy to envision how precise positioning of the Hsc20-Isu1-Ssq1 system promotes avidity driven substrate trapping, via syneraistic stimulation of ATP hydrolysis. Other systems likely achieve sufficient avidity by increasing local concentration of substrate and J-domains using less precise positioning.^{48–52} JDP rigidity, though advantageous in the case of a single substrate, would likely constrain adaptability to function with many substrates. Having a more flexible, yet restricted distance between J-domain and JDP bound substrate sufficiently increases local concentrations. In the case of JDPs that do not bind substrate on their own, avidity could be achieved with help of additional "adapter" protein(s) or additional domains of the JDP itself that position substrate or

JDP (or both) in close proximity to the Hsp70 partner. Such mechanisms likely operate for JDP/ Hsp70 systems involved in protein import into the mitochondrial matrix and at the tunnel exit of the ribosome.⁵³

Materials and Methods

Protein purification

Mutants of *SSQ1* were constructed by sitedirected PCR mutagenesis. Recombinant mature Ssq1 (residues 19-657) WT, Ssq1^{T239A} (Ssq1*) and Ssq1*^{-F462S} with a polyhistidine (His) tag at the C-terminus were purified as described in.⁵⁴ Recombinant mature Hsc20, also termed Jac1 (residues 10-184) WT and alanine substitution variants (Table 1) with the C-terminal His tag were purified as described in Supplementary Methods. Recombinant mature Isu1 (residues 36-165) with the C-terminal His tag and Isu1^{GST} with C-terminal GST tag were purified as described in²⁶ and,²³ respectively.

GST pulldown assay

Pull-down experiments were performed as described in.³⁴ In short, in a 150 μ I reaction, 2.5 μ M Isu1^{GST} or purified GST (control for background binding) was incubated with Ssq1* and/or Hsc20 (typically 5 µM and 2.5 µM, respectively) in buffer A (40 mM HEPES-KOH, pH 7.5, 5% (v/v) glycerol, 100 mM KCl, 1 mM dithiothreitol, and 10 mM MgCl₂) for 15 min at 25 °C to allow complex formation. An aliquot of the reaction (7.5 µL) was taken as a loading control "input 5%". 40 µl of glutathione-immobilized agarose beads (preequilibrated with 0.1% (w/v) bovine serum albumin, 0.1% (v/v) Triton X-100 and 10% (v/v) glycerol in buffer A) were added to each reaction and incubated at 4 °C for 1 h with rotation. The beads were washed one time with 500 μ l of buffer A and then three times with 200 µl of buffer A. Proteins bound to the beads were incubated with 20 ul of two-fold concentrated Laemmli sample buffer (125 mM Tris-HCl, pH 6.8, 5% (w/v) sodium lauryl sulfate (SDS), 10% (v/v) 2-mercaptoethanol, 20% (v/v) glycerol) for 10 min at 100 °C and aliquots were loaded on SDS-PAGE (Bolt™ 12%, Bis-Tris, 1.0 mm, Mini Protein Gels, Invitrogen) and visualized by Coomassie blue staining.

Structural model of the tripartite Ssq1(ATP)-Hsc20-lsu1 complex

All simulations were performed using GROMACS 2021.3,⁵⁵ if not stated otherwise. The CHARMM36 force field⁵⁶ was used for proteins, ions and ATP-Mg, and the TIP3P model was used for water. In each of the simulation boxes, the numbers of Na⁺ and Cl⁻ ions were adjusted to 0.15 M. If not stated otherwise, temperature was kept at 298.15 K with

the v-rescale algorithm⁵⁷ using a coupling constant of 0.1 ps. Pressure was kept at 1 bar using the Parrinello-Rahman algorithm⁵⁸ with a coupling time of 5 ps. Periodic boundary conditions were applied and the Particle Mesh Ewald summation⁵⁹ was used to calculate long-range electrostatic interactions, with a cut-off radius of 1 nm and a Fourier grid spacing of 0.12 nm. Van der Waals interactions were calculated with Lennard-Jones potential with a cut-off radius of 1 nm. All bonds involving hydrogen were constrained using the LINCS algorithm. Leap-frog Verlet algorithm was used to integrate equations of motion with a time step of 2 fs.

A structural model of the tripartite complex was obtained by docking the homology model of Isu1 to the structural model of the Ssg1(ATP)-Hsc20 complex using ClusPro⁶⁰: a) without restraints on protein-protein interactions; b) with attraction restraint on Isu1 LPPVK - Ssq1 F462 interaction; c) with attraction restraint on Isu1 L63,V72, F94 -Hsc20 L105, L109, Y163 interaction; d) with both restraints simultaneously. From the 382 structures obtained from docking we selected 72 that had Isu1 located near Hsc20 CTD. Redundancy among these 72 models was removed using gmx cluster tool from GROMACS package, with single-linkage clustering method and 0.5 nm root mean square deviation (RMSD) cutoff on Isu1 position, resulting in 11 distinct models of the Hsc20-Ssq1(ATP)-Isu1 complex. Next, each of these models was placed in a $16 \times 16 \times 16$ nm dodecahedron box. solvated with \sim 90.000 water molecules and energy-minimized. The systems were then simulated in equilibrium MD for at least 500 ns - total simulation time was 8.6 µs. Trajectories were superimposed on Ca atoms of Ssq1 NBD domain, and clustered based on backbone of Hsc20 and Isu1 using gmx cluster with Jarvis-Patrick method and 0.3 nm RMSD cutoff.

Purification of tripartite Ssq1*-Hsc20-Isu1 complex from *Escherichia coli* co-expressing its components

Rosetta 2 strain with plasmid E. coli pLysSRARE2 (chloramphenicol resistance) was transformed with plasmid pRSF HSC20-ISU1 resistance) (kanamycin harboring Hsc20 sequence without tag and Isu1 sequence tagged with six histidine codons at C-terminus and SSQ1*-HEP1 plasmid pETDuet1 (ampicillin resistance) harboring Ssq1 sequence tagged at Cterminus with six histidine codons and no tagged Hep1 sequence. Hep1 stabilizes Ssq1 when expressed in bacterial cells,^{54,61} but does not form stable complex with Ssq1 and is not present in the tripartite complex preparation. The strain was grown in LB medium (with 50 µg/ml ampicillin, 25 μg/ml kanamycin and 10 μg/ml chloramphenicol) at 37 °C. Expression was induced by addition of 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at $A_{600} = 0.45$. After 3 hours at 30 °C, cells

were harvested and lysed in a French Press in buffer B (25 mM HEPES-KOH pH 7.5, 150 mM KCl. 1 mM PMSF, 10% (v/v) glycerol 2 mM magnesium acetate and 30 mM imidazole pH 7.5). After clarifying spin, the supernatant was loaded on a Novagen His-Bind Resin column (2.5 ml) equilibrated in buffer B. The column was washed with buffer B (50 ml) and subsequently with buffer B with the addition of 10 mM MgCl₂ and 1 mM ATP (25 ml). Proteins were eluted with 30 mM-500 mM imidazole gradient in buffer B (20 column volumes). Fractions containing Ssq1*, Hsc20 and Isu1 and fractions containing Hsc20 and Isu1 were concentrated using Amicon[®] Ultra – centrifugal filters with 100 kDa and 30 MWCO, respectively. Concentrated tripartite complex was incubated at 4 °C with 1 mM dithiothreitol for 10 min before addition of 2 mM ATP and 10 mM MgCl₂ and further 20 min incubation. Then the complex was subjected to size exclusion chromatography in buffer C (25 mM HEPES-KOH pH 8.0, 100 mM KCl, 5% (v/v) glycerol, 1 mM DTT and 10 mM MgCl₂) using HiLoad 16/600 Superdex 200 prep grade column. Fractions containing all three proteins were concentrated using ${\sf Amicon}^{\circledast}$ Ultra – 4 centrifugal filter with 100 kDa MWCO, and stored at -70 °C.

Hydrogen/deuterium exchange mass spectrometry (HDX-MS)

Protein samples (5 μ l) were diluted 10-fold in D₂O buffer D (25 mM HEPES-KOD, 100 mM KCl, 5% (v/ v) glycerol, 10 mM MgCl₂, 1 mM DTT, pD = 8.0) and incubated for 10 s, 1 min, 5 min, 25 min, 120 min, and additionally 30 s for samples containing Ssq1*, at room temperature. The exchange was quenched by transferring 50 µl of reaction mixture to ice-cold eppendorf tube containing 10 ul of quench buffer (2 M glycine, 4 M guanidium hydrochloride. 1 M Tris(2-carboxyethyl)phosphine (TCEP), pD = 2.4). Samples were then vortexed and flash frozen in liquid nitrogen and kept frozen at -80 °C until MS data acquisition. Reactions were performed in quadruplicates (in triplicates for reactions containing Ssq1*). Control samples with maximal deuteration level were incubated in D₂O buffer for 24 h and guenched as described, while minimal deuteration controls were prepared by adding the protein to the quenched reaction conditions. Deuteration levels were not corrected for back exchange.

Samples were injected into a nanoACQUITY UPLC system (Waters) and digested online using immobilized pepsin resin column (Poroszyme) with 0.07% (v/v) formic acid in water as mobile phase (200 μ l/min flow rate). Peptides were loaded onto the 2.1 mm \times 5 mm C18 trapping column (ACQUITY BEH C18 VanGuard precolumn, 1.7 μ m resin, Waters) and eluted onto a reversed phase C18 column (Acquity UPLC BEH, 2.1 \times 50 m m, 1.7 μ m resin, Waters) using a 7–35% gradient of acetonitrile in 0.1% (v/v) formic acid at 90 μ l/min flow

rate. Temperature of all fluidics, columns, and valves, except the pepsin digestion column kept at 13 °C, was maintained at 0.5 °C using HDX Manager (Waters). Outlet of the C18 column was directly coupled to the ion source of SYNAPT G2-HDMS mass spectrometer (Waters) operating in IMS mode. Leucine-enkephalin (Sigma) was used for carrying out lock mass and activation. Mass spectra were acquired in IMS mode over the m/z range of 50–2000. The spectrometer parameters were set as described in.⁶²

Peptides were identified using ProteinLynx Global Server software (Waters) using a randomized database, based on a peptide list obtained using non-deuterated proteins, processed as described above and measured in MS^E mode. The peptide list was filtered by minimum intensity (3000) and minimum product per amino acid (0.3) and analyzed using DynamX 3.0 (Waters). All MS spectra were manually inspected. Percentage of peptide deuteration was calculated using HaDeX tool,⁶³ using minimal and maximal deuteration controls described above as 0% and 100%, respectively. Difference in number of exchanging protons was calculated using HaDeX. Exchange levels and differential deuterium exchange of peptides exhibiting exchange kinetics representative for the protein region (Supplementary Excel files) were then mapped on Ssq1, Hsc20 and Isu1 structural models using VMD.⁶

Bio layer interferometry (BLI)

BLI measurements were performed in buffer E (40 mM HEPES-KOH, pH 7.5, 5% (v/v) glycerol, 100 mM KCl, 1 mM dithiothreitol, 10 mM MgCl₂, 0.05% (v/v) Triton X-100) using a single channel BLItz instrument (Pall ForteBio) operating at room temperature. Sensograms were recorded as a function of time. After the initial equilibration step (30 s) Isu1^{GST} or GST (10 μ M) was immobilized on anti-GST biosensors in the presence of bovine serum albumin at 0.5 mg/ml (10 min). The sensors were then washed with buffer C (10 min). Following a baseline step (30 s), the sensors were immersed into solutions containing analytes to measure association, followed by immersion into protein-free solutions to record dissociation (10 min). To assess the background binding, sensors with immobilized GST were used.

CRediT authorship contribution statement

Marcin Jelen: Investigation. Igor Grochowina: Investigation, Software, Formal analysis, Data curation, Visualization. Aneta Grabinska-Rogala: Investigation. Szymon J. Ciesielski: Investigation, Methodology. Katarzyna Dabrowska: analysis. Investigation, Formal Bartlomiej Tomiczek: Formal analysis, Software. Lukasz Nierzwicki: Formal analysis, Software. Wojciech Delewski: Investigation. Brenda

Schilke: Investigation. Methodology. Jacek Czub: Supervision. Formal analysis. Michal Dadlez: Supervision, Formal analysis. Rafal Supervision, Dutkiewicz: Investigation, Methodology, Writing - original draft. Elizabeth A. Craig: Conceptualization, Supervision, Writing review & editing, Funding acquisition. Jaroslaw Marszalek: Conceptualization, Supervision, Writing - original draft, Writing - review & editing, Funding acquisition.

DECLARATION OF COMPETING INTEREST

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jmb.2023. 168283.

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† Equal contribution.

Abbreviations:

 JDP, J-domain protein; SBD, substrate binding domain of Hsp70; NBD, nucleotide binding domain of Hsp70; CTD, C-terminal domain of Hsc20; Ssq1*, Ssq1(T239A) variant defective in ATPase activity; GST, glutathione-Stransferase; MD, molecular dynamics simulations; HDX, hydrogen-deuterium exchange; MS, mass spectrometry; BLI, bio-layer interferometry; RMSD, root-mean-square deviation; LB, Luria Broth media

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