

Network of general and specialty J protein chaperones of the yeast cytosol

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Contributed by Elizabeth Anne Craig, March 13, 2007 (sent for review January 15, 2007)

J proteins are obligate cochaperones of Hsp70s, stimulating their ATPase activity and thus allowing them to function in multiple cellular processes. In most cellular compartments, an Hsp70 works with multiple, structurally divergent J proteins. To better understand the functional specificity of J proteins and the complexity of the Hsp70:J protein network, we undertook a comprehensive analysis of 13 J proteins of the cytosol of the yeast *Saccharomyces cerevisiae*. Phenotypes caused by the absence of four proteins, Ss1, Jj1, Jj3, and Cwc23, could not be rescued by overexpression of any other cytosolic J protein, demonstrating the distinctive nature of J proteins. In one case, that of Zuo1, the phenotypic effects of the absence of a J protein could be rescued by overexpression of only one other J protein, Jj1, which, like Zuo1, is ribosome-associated. In contrast, the severe growth phenotype caused by the absence of the cytosol's most abundant J protein, Ydj1, was substantially rescued by expression of J domain-containing fragments of many cytosolic J proteins. We conclude that many functions of Hsp70 chaperone machineries only require stimulation of Hsp70's ATPase activity by J protein partners. However, a subset of Hsp70 functions requires specific J protein partners, likely demanding either sublocalization within the compartment or binding to specific client proteins.

Hsp40 | Hsp70 | molecular chaperone | multigene family

Through their action in protein folding, degradation, translocation across membranes, and disassembly of protein complexes, molecular chaperones are important participants in many crucial cellular processes (1, 2). Hsp70s and their J protein partners (at times referred to as Hsp40s) constitute an important component of the cellular “chaperone” in virtually all living systems (3). In most cellular compartments, an Hsp70 has multiple J protein partners. The cytosol, a hub of activity for a variety of crucial cellular processes, is no exception. With a goal of better understanding the degree of functional overlap among J proteins, as well as their individual specificities, we choose the cytosol of *Saccharomyces cerevisiae* as the focus of a comprehensive analysis of J protein function.

Although J proteins are obligate partners of Hsp70s, it is the Hsp70 that has long been considered the core of the Hsp70:J protein chaperone machine by virtue of their client protein interaction, which is modulated by nucleotide binding (4). ATP binding fosters rapid binding to client proteins, whereas nucleotide hydrolysis stabilizes the interaction. The cytosol of *S. cerevisiae* contains two predominant classes of Hsp70s, Ssa (SSA1–4) and Ssb (SSB1–2) (5). Yeast cells expressing only one representative of a class grow very similarly to wild-type cells under a variety of conditions. But the classes are functionally distinct, because a member of the Ssa class cannot substitute for an Ssb Hsp70 and vice versa (6).

A role of all J proteins is stimulation of the ATPase activity of their partner Hsp70s. Such stimulation fosters productive interaction of Hsp70s with their client proteins (4). The ≈65-aa J domain, the defining feature of all of the J proteins, is responsible for the stimulation. The J domain is characterized by a highly conserved histidine–proline–aspartic acid (HPD) tripeptide signature motif that is important for J domains' stimula-

tory activity. Despite the omnipresent J domain, J proteins, as a group, are strikingly dissimilar, varying significantly in their domain organization and localization within the cytosol (7). Other domains have been shown to associate directly with client proteins, thereby fostering client protein interaction with Hsp70 or promoting localization to a particular site within a cellular compartment.

Historically, J proteins have been divided into three classes (I, II, and III). The class I designation is based on the motifs/domains present in the founding member of this group, DnaJ of *Escherichia coli*: an N-terminal J domain, followed by a glycine/phenylalanine (G/F)-rich region, four repeats of the CxxCxGxG-type zinc finger, and a C-terminal extension known to bind client proteins (8). The cytosol of *S. cerevisiae* has three J proteins that have been placed in class I: Apj1, Xdj1, and Ydj1 (7). Class II J proteins, by definition, have a similar structure, in that they have a J domain and a G/F region. However, class II J proteins lack the CxxCxGxG-type zinc finger domain. Cytosolic class II J proteins include Caj1, Djp1, Hlj1, and Sis1. All of the other J proteins have been arbitrarily placed in class III. A few *S. cerevisiae* class III J proteins contain zinc-binding domains, but these are not of the type found in class I proteins. Jj1 contains C₂H₂-type zinc fingers, whereas Jj3 has a CSL-type zinc finger (Fig. 1). Apart from structural differences among J proteins, the relative abundance of cytosolic J proteins also varies dramatically (Fig. 1). Thus, it is possible that potential functional overlap between J proteins is masked by differences in their expression levels.

To better understand the functional diversity displayed by J proteins, we undertook an analysis of cytosolic J proteins of *S. cerevisiae*. Several J proteins of the cytosol appeared to be functionally unique, because overexpression of no other J protein was able to substitute for their functions. We also found that the function of the most abundant J protein of the cytosol, Ydj1, could be carried out by expression of the J domain from several diverse J proteins. Our analysis of a single cellular compartment likely provides a paradigm for understanding the general principles of diversity of function of J protein:Hsp70 machineries in other cellular compartments and in other organisms.

Results

Deletion Phenotypes of Genes Encoding Cytosolic/Nuclear J Proteins.

As a starting point for our analysis of cytosolic J protein function, we constructed or obtained strains having a deletion of each of the 10 genes that encode a predominantly cytosolic J protein, as defined in ref. 9: Apj1, Djp1, Jj1, Jj2, Jj3, Sis1, Swa2, Xdj1, Ydj1, and Zuo1. We also analyzed three other J proteins: Cwc23,

Author contributions: C.S. and E.A.C. designed research; C.S. performed research; C.S. and E.A.C. analyzed data; and C.S. and E.A.C. wrote the paper.

The authors declare no conflict of interest.

Abbreviations: CCV, clathrin-coated vesicle; DPH, diphthamide; DT, diphtheria toxin; G/F, glycine/phenylalanine; GPD, glyceraldehyde 3-phosphate dehydrogenase.

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This article contains supporting information online at www.pnas.org/cgi/content/full/0702357104/DC1.

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| Protein | Mol/Cell | Phenotype Class | Domain structure |
|---------|----------|--------------------------------------|----------------------------------|
| Apj1 | 125 | - | I J QS Zn 528 |
| Caj1 | - | - | II J G 391 |
| Cwc23 | - | Lethal | III J 283 |
| Djp1 | 8970 | - | II J G 432 |
| Hlj1 | 1840 | - | II J GF TM 224 |
| Jjj1 | 2310 | C ^S | III J Z Zn CH Zn 590 |
| Jjj2 | 184 | - | III J 482 |
| Jjj3 | 2010 | DTR | III J Zn 172 |
| Sis1 | 20300 | Lethal | II J GF/GM 352 |
| Swa2 | 768 | C ^S (CCV def.) | III CBD UBA TPR 668 |
| Xdj1 | 1210 | - | I J G Zn 459 |
| Ydj1 | 119000 | Very slow gr | I J GF Zn 409 |
| Zuo1 | 86400 | C ^S ; cation ^S | III J Z CH 433 |

Fig. 1. Cytosolic J proteins of *S. cerevisiae*. Molecules per cell are as reported by Ghaemmaghami *et al.* (11). No data available is indicated by -. Phenotypes tested are as follows: cold sensitivity (C^S), resistance to diphtheria toxin (DT^R), very compromised for growth (very slow gr.), sensitive to cations (cation^S), and defective in uncoating CCVs (CCV def.). Information was obtained from the *Saccharomyces* Genome database, www.yeastgenome.org. J protein classes were taken from Walsh *et al.* (7). In the domain structure, arrows indicate the J domain fragments used (Apj1₁₋₁₆₁, Caj1₁₋₁₃₉, Cwc23₁₋₁₃₉, Djp1₁₋₁₂₇, Hlj1₁₋₁₇₉, Jjj1₁₋₁₂₈, Jjj2₁₋₁₃₂, Jjj3₁₋₁₂₄, Sis1₁₋₁₆₇, Swa2₃₆₂₋₆₆₈, Xdj1₁₋₁₄₆, Ydj1₁₋₁₃₄, and Zuo1₁₋₂₃₄). Numbers indicate amino acids in each full-length J protein. Regions in which these amino acids are highly represented include QS, G, GF, and GF/GM (listed by their standard single-letter amino acid code). J, J domain; Zn, zinc finger; CBD, clathrin-binding domain; UBA, ubiquitin association; TPR, tetratricopeptide repeat; Z, zuotin-like; CH, charged; TM, transmembrane.

which, although predominantly nuclear, is in the cytosol as well; Hlj1, an endoplasmic reticulum (ER) membrane protein whose J domain faces the cytosol (10); and Caj1, whose localization has not been reported, but which lacks apparent sequences for targeting to any organelle. The phenotypes of the knockout strains, all in the W303 genetic background, were similar to those previously reported, with the exception of $\Delta swa2$, which grew slowly at low temperatures. Seven deletion strains, those lacking Cwc23, Jjj1, Jjj3, Sis1, Swa2, Ydj1, and Zuo1, had easily assayable phenotypes and became the focus of our studies. The reported abundance of J proteins within the cytosol ranges from >100,000 molecules per cell of Ydj1 to only 125 molecules per cell of Apj1 (11). Because of this large variation, we reasoned that functional overlap might exist among these proteins that is not evident from the phenotypes displayed by the single gene knockouts. Therefore, overexpression constructs were made for all 13 cytosolic J proteins by cloning the corresponding ORFs under the glyceraldehyde 3-phosphate dehydrogenase (GPD) promoter in a high-copy plasmid. Transformants harboring constructs containing Caj1, Djp1, Hlj1, and Xdj1 were either not obtained or grew very poorly, presumably because overexpression of these J proteins was deleterious. Therefore, these expression plasmids were not included in the experiments discussed below.

J Domain Fragments Are Sufficient for Robust Growth of $\Delta ydj1$. We began our analysis with Ydj1, because it is the most abundant cytosolic J protein. Ydj1 partners with Ssa Hsp70s in protein folding and translocation of proteins into endoplasmic reticulum (ER) and mitochondria (12–15). $\Delta ydj1$ cells are viable but have a severe growth defect at all temperatures. As expected from earlier reports (12), increased expression of Sis1 improved

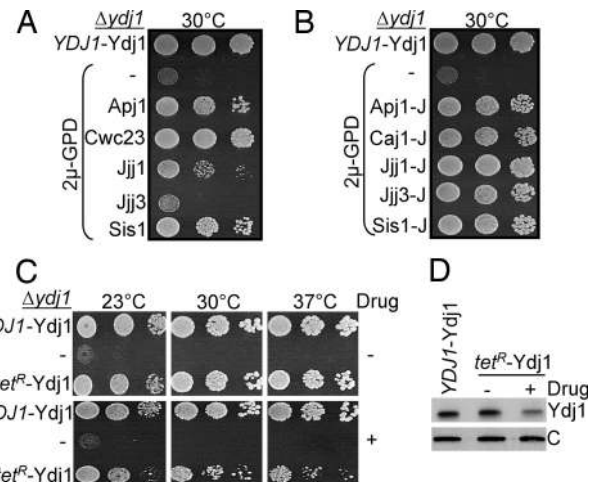


Fig. 2. Ability of other J proteins and lower levels of Ydj1 to rescue $\Delta ydj1$ growth phenotype. (A–C) Ten-fold serial dilution of $\Delta ydj1$ cells expressing the indicated proteins were plated onto minimal medium and incubated for 3 days at the indicated temperatures. The following were used as controls: Ydj1 under control of its own promoter in a centromeric vector, pRS314 (YDJ1-Ydj1); and empty vector (-). (A and B) Overexpressing full-length (A) or J domain fragments of cytosolic J proteins, as indicated in the legend to Fig. 1 (B), driven by the GPD promoter from a 2 μ plasmid. The complete set of plates is shown in SI Fig. 6A. (C) Ydj1 under control of the tetracycline-regulatable promoter (*tet*^R-Ydj1) or empty vector (-) in the absence (-) or presence (+) of 0.5 μ g/ml doxycycline (drug). (D) Total lysates prepared from cells grown in the absence (-) or presence (+) of 0.5 μ g/ml doxycycline (drug) were resolved by SDS/PAGE, electro-blotted, and probed with anti-Ydj1 antibodies and, as loading control, anti-Ssc1 antibodies (indicated by "C").

growth of $\Delta ydj1$ cells [Fig. 2A and supporting information (SI) Fig. 6A]. Overexpression of Apj1 also dramatically rescued the growth defect of $\Delta ydj1$ cells (Fig. 2A). Such rescue was not particularly surprising, because Apj1, like Ydj1, is a class I J protein. However, overexpression of the class III J protein, Cwc23, also rescued $\Delta ydj1$ cells significantly (Fig. 2A), even though it only has a J domain in common with Ydj1.

Because of rescue by divergent J proteins that had no obvious sequence similarity beyond their J domains, we reasoned that the J domain alone might be sufficient. Therefore, we designed constructs to express J domain-containing fragments at high levels. In contrast to the full-length proteins, J domain fragments of many cytosolic J proteins were able to substantially rescue $\Delta ydj1$ (Fig. 2B and SI Fig. 6B). At a variety of temperatures, the J domain fragment of Jjj1 permitted the most robust growth of any fragment tested. This rescue depended on J domain function, because Jjj1-J_{H32Q}, having an alteration in the histidine–proline–aspartic acid (HPD) motif, did not rescue (SI Fig. 6C). Even though expression of full-length Caj1 and Djp1 were deleterious (data not shown), their J domain fragments were able to rescue $\Delta ydj1$ (SI Fig. 6B). In addition, J domain fragments from a number of class III J proteins, such as Jjj3-J, rescued, although the full-length construct did not (Fig. 2A and B). We are unable to make a comprehensive summary regarding the ability of each full-length protein or J domain fragment to rescue because of the lack of tools to measure levels of expression of each polypeptide. However, we can draw the general conclusion that a variety of J domain-containing fragments are competent to carry out functions of Ydj1.

Normal Levels of a J Domain Fragment Are Sufficient for Rescue of $\Delta ydj1$. The rather efficient rescue of $\Delta ydj1$ by J domain fragments raised two questions: (i) What level of full-length Ydj1 is required for wild-type growth? (ii) How robust is the growth of

are tolerant to DT. For testing Jjj3 function *in vivo*, the overexpression constructs were transformed into a $\Delta j j j 3$ strain harboring a plasmid encoding the catalytic subunit of DT under the control of a galactose-inducible promoter. None of the J domain fragments, including that of Jjj3 itself, or full-length proteins could perform Jjj3's function *in vivo*, as evidenced by the ability of transformants to grow on galactose-based media (Fig. 4A and data not shown).

The inability of the J domain fragment of Jjj3 to substitute for the full-length protein (Fig. 4A) suggested to us that sequences in addition to the J domain may be critical for Jjj3's role in DPH biosynthesis. Jjj3 is a small protein, having only 94 residues, encompassing a CSL zinc finger in addition to the J domain. To determine whether this motif is important for Jjj3 function, we tested a mutant protein, Jjj3_{C156Y}, having a tyrosine substituted for the cysteine in the conserved CSL tripeptide. Jjj3_{C156Y} cells were tolerant to DT (Fig. 4A), indicating that the CSL domain is critical for Jjj3's function. In addition, we tested whether chimera Y^JJjj3, containing the J domain of Ydj1 (amino acids 1–63) and C-terminal region of Jjj3 (amino acids 70–172), could substitute for Jjj3. This construct was functional in DPH biosynthesis, because $\Delta j j j 3$ cells harboring this plasmid were sensitive to DT (Fig. 4C). Thus, although a J domain is required for Jjj3's function (SI Fig. 8), it is not a Jjj3 specificity determinant.

A Jjj3 J domain fragment, but not full-length protein, was competent to substitute for Ydj1 when overexpressed (Fig. 2A and B). We asked whether alteration of the zinc finger would allow full-length Jjj3 to rescue $\Delta y d j 1$ cells. Jjj3_{C156Y} rescued growth of $\Delta y d j 1$ cells as the Jjj3-J domain fragment (Fig. 4B).

Specialization of Essential J Proteins, Cwc23 and Sis1. We also tested two essential J proteins: Cwc23, which is implicated in RNA splicing (7), and Sis1, which is required for the maintenance of the [RNQ⁺] prion and is thought to play an important role in translation initiation (18, 19). $\Delta c w c 23$ and $\Delta s i s 1$ haploids carrying their respective wild-type genes on a *URA3*-based plasmid, as well as one of the expression plasmids, were plated on 5-fluoroorotic acid (5-FOA)-containing plates to select for cells having lost the wild-type gene. None of the heterologous full-length or J domain-containing fragments rescued either deletion strain (data not shown), suggesting that Sis1 and Cwc23 contain sequences specifically required for their essential functions.

Specialization of the Ribosome-Associated J Proteins, Zuo1 and Jjj1. Zuo1 and Jjj1 are both ribosome-associated J proteins. Zuo1 is the J protein partner of Ssb, the specialized ribosome-associated Hsp70 that binds nascent chains exiting the ribosome (20). Jjj1 functions in 60S subunit biogenesis with Ssa (21). All constructs were tested for rescue of $\Delta j j j 1$ and $\Delta z u o 1$. None of our expression plasmids rescued the cold sensitivity of cells lacking Jjj1, indicating a specialized function of this J protein (data not shown). As reported recently (21), overexpression of full-length Jjj1 partially rescued the cation and cold sensitivity of $\Delta z u o 1$ cells. However, no other construct had an effect on the growth of $\Delta z u o 1$ cells, indicating that Jjj1 is unique in its ability to partially substitute for Zuo1.

Swa2, the Auxilin Homolog. Swa2 is involved in the uncoating of clathrin-coated vesicles (CCVs) (22). Swa2 is a complex protein, having clathrin-binding domains (CBDs), a ubiquitin association domain (UBA), tetratricopeptide repeat (TPR) motifs, and a C-terminal J domain. All of the full-length and the J domain-containing constructs were transformed into $\Delta s w a 2$ cells. None of the full-length J protein constructs were able to rescue the cold sensitivity of $\Delta s w a 2$ (data not shown). However, several J domain-containing fragments, including at least one from each class, partially rescued the cold sensitivity of $\Delta s w a 2$ (Fig. 5A). As

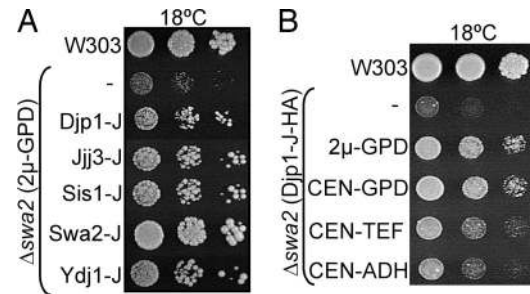


Fig. 5. J domain-containing fragments of some cytosolic J proteins partially rescued the cold sensitivity of $\Delta s w a 2$. Serial dilutions of $\Delta s w a 2$ cells containing the indicated plasmids were spotted on minimal media, and plates were incubated at 18°C for 10 days. Shown are wild-type control strain (W303) and $\Delta s w a 2$ harboring empty vector (–). (A) 2 μ -GPD plasmids expressing indicated J domain fragments. (B) Djp1-J HA fragment expressed from indicated promoters of variable strengths were transformed in $\Delta s w a 2$.

observed in the case of $\Delta y d j 1$, the level of rescue positively correlated with expression levels (Fig. 5B).

Discussion

We began a broad analysis of cytosolic J proteins to better understand the degree of functional diversity that exists within a single cellular compartment. As discussed below, a complex picture emerges, with the cytosolic compartment having a mixture of “general” and “specialty” J proteins.

The General J Protein, Ydj1. Surprisingly, the severe growth defects caused by the absence of the most abundant cytosolic J protein, Ydj1, could be substantially alleviated by the expression of J domains from a variety of J proteins. This sufficiency of the J domains to substitute for Ydj1 implies that the core ability of a J protein, the ability to stimulate the ATPase activity of its partner Hsp70, is sufficient for many cellular processes. Such rescue required neither a J domain from a particular class of J proteins nor expression at levels much higher than normal Ydj1 levels. Thus, even though full-length Ydj1 is competent to bind and deliver client proteins to Ssa Hsp70 and is a requirement for *in vitro* refolding of luciferase (8), we propose that this activity is not required for many of its *in vivo* functions.

However, we do not mean to imply that client protein binding is never critical for the function of “general” J proteins. First, a J domain, even when expressed at the level at which Ydj1 is normally expressed, does not fully rescue the growth defect caused by the absence of Ydj1. Second, our laboratory previously reported that the C-terminal substrate-binding domain of either Ydj1 or Sis1 is required for robust growth of yeast cells (23). It is likely that certain client proteins require interaction with a J protein for efficient presentation to Ssa but that binding to either Ydj1 or Sis1 often suffices in these cases. It is likely that other cytosolic J proteins functionally overlap with Ydj1 *in vivo*, as well. For example, both Ydj1 and class II Hlj1 participate in the degradation in the cytosol of proteins extracted from the endoplasmic reticulum (ER) (10).

The G/F Region of Class I and II J Proteins. The ability of the Ydj1 J domain, without any of the amino acids normally flanking it being present, to substantially rescue the growth defect of cells lacking Ydj1 established the sufficiency of the J domain itself. However, this rescue also raised a question concerning the importance of the G/F region, a defining feature of class I and II J proteins that lies adjacent to the J domain. Its functional significance has been an ongoing point of debate, in part due to the report that the growth defects of $\Delta y d j 1$ could be rescued by a fragment containing the J domain plus the G/F region, but not

by a construct encoding only the J domain of Ydj1 (24). However, the results reported here indicate that the need for such sequences is nonspecific, but rather may be needed for structural stability.

Nevertheless, the sufficiency of the J domain does not mean that, in the context of full-length proteins, the region rich in glycines never serves an important purpose. In analyses of both DnaJ of *E. coli* and Sis1 of *S. cerevisiae*, the G/F region has been shown to be functionally important. However, in these two cases, the specific sequences found to be critical were neither glycines nor phenylalanines. For example, the defects caused by the deletion of the G/F region of DnaJ of *E. coli* were mimicked by alterations of the D or I/V of DI/VF repeats found at the end of the G/F region (16). In the case of Sis1, alteration of a single D or N residue in a small, 12-aa “insertion segment” that distinguishes its G/F region from that of Ydj1 abolished the ability of a J domain plus G/F fragment, Sis1_{1–121}, to carry out Sis1’s essential functions (25).

Consistent with the idea that it is not the glycines and phenylalanines themselves in a G/F-rich region that are functionally important, a mutant *YDJ1* protein lacking the entire G/F region rescued a $\Delta ydj1$ strain and full-length Ydj1 (26). Furthermore, a closer look at the amino acid sequence of J proteins found in the yeast cytosol revealed that even the presence of a G/F-rich region as the distinguishing feature of class I and II J proteins is somewhat arbitrary. For example, in reviews (3, 7), Apj1, Caj1, Djp1, Hlj1, Sis1, Xdj1, and Ydj1 were classified as class I or II proteins, meaning that, by definition, a G/F region was present. The predominance of glycines and phenylalanines in the 33 aa C-terminal to the J domain of Ydj1 is obvious, with 15 glycines and seven phenylalanines. However, the presence of a G/F region adjacent to the J domain of Apj1 and Djp1 is not as clear. Apj1 has two glycines and three phenylalanines; Djp1 has six glycines and four phenylalanines.

Specialist J Proteins. As a counterpoint to the generality of Ydj1 function, the degree of specificity among J proteins is also illuminated by the results of our study. Seven of the 13 J protein deletion strains analyzed had assayable phenotypes. Of these, the phenotypes of four: $\Delta cwc23$, $\Delta sis1$, $\Delta jji1$, and $\Delta jji3$ could only be rescued by expression of the deleted genes, suggesting a high degree of specificity. What is the basis of such specificity? Although there is still much to be learned, the data presented here and elsewhere provide some clues. Tethering to a particular location within a particular cellular compartment may be important. For example, $\Delta zuo1$ is not listed above, because its phenotype could be partially rescued by one other J protein, Jjj1. Both Jjj1 and Zuo1 stably associate with ribosomes (21, 27). In cases in which tethering is important, an extremely high local concentration of the J protein may be required to recruit an Hsp70 partner to a particular site of action, a criteria, which in the case of Zuo1, would only be met by the other ribosome-associated J protein, Jjj1.

Interestingly, although both Jjj1 and Zuo1 are ribosome-associated, and Jjj1 can partially substitute for Zuo1, these two J proteins partner with different Hsp70s, Zuo1 with Ssb (27) and Jjj1 with Ssa (21). This ability of Ssa to function with an alternative J protein and substitute for Ssb is reminiscent of the ability of the human Zuo1 ortholog, Mpp11, to substitute for Zuo1 (28). In doing so, Mpp11 partners with Ssa, consistent with Ssb being present only in fungi and with Ssa orthologs being found in all eukaryotes. This ability also underscores the specificity that lies within the J protein group of proteins that, at least in many cases, eclipses the specificity of Hsp70s. Indeed, in the case of the yeast cytosol, both comparison of J domain sequences and functional information is consistent with Zuo1 being the only J protein partner of Ssb, with the other 12 working with Ssa.

One effect of the sequestering of a J protein, either by

localization to a particular site or by interaction with a particular client protein, may be its inability to substitute *in vivo* for another J protein, even if it is otherwise functionally competent. This idea is consistent with the observation that, unlike full-length Jjj3, a Jjj3_{C156Y} construct, which is unable to perform Jjj3’s specialized function in DPH biosynthesis, was able to rescue $\Delta ydj1$ as the Jjj3 J domain construct. Although, the role of Jjj3’s zinc finger in DPH synthesis is not known, one can speculate that it might either bind to a client protein or to a protein complex involved in DPH biosynthesis. Therefore, in cases such as Jjj3, the sequestration might prevent functional overlap.

However, increased expression of full-length J proteins such as Caj1, but not their J domains, is deleterious to cells. In fact, the J domain fragment of Caj1 rescued the growth defects caused by the absence of Ydj1 quite effectively. At this point, we can only speculate about the cause of the toxicity. Perhaps these J proteins are deleterious because they bind client proteins “inappropriately” or because they recruit Hsp70 to specific sites, depleting the pool available to function with specialized J proteins.

Swa2 presents an unusual case and, on the basis of our data, cannot be easily classified as a general or specialty J protein. It is structurally complex and highly specialized for uncoating of CCVs (29). However, the cold sensitivity of $\Delta swa2$ could be rescued by a number of J domain-containing constructs. Our result is consistent with the ability of a fragment containing the J domain and the adjacent tetratricopeptide repeat (TPR) motifs of Swa2 itself to partially complement the defect of $\Delta swa2$ in α -factor processing (30), even though it cannot bind clathrin. The simplest explanation is that such specialization, that is direct binding to clathrin, is not absolutely critical under the conditions tested. It should also be noted that Swa2 is present in <1,000 molecules per cell, whereas even $\approx 100,000$ molecules per cell of a J domain does not allow wild-type growth of $\Delta swa2$ cells. Similarly, the apparent importance of the substrate-binding domain of Ydj1 at higher temperatures may be due to the higher demand for refolding of partially denatured proteins.

Conclusions

In summary, our results suggest that important general functions of J proteins can be carried out by J domains, indicating only a requirement for stimulation of Hsp70s’ ATPase activity. The specificity of J proteins is largely governed by regions outside the J domain. Mechanistically understanding the basis of specificity of individual J protein in the yeast cytosol demands further study, but substrate specificity and sequestration to particular sites within a cellular compartment likely play significant roles. However, it is also possible that subtle but important alterations in the cycle of binding and release of Hsp70s from particular client proteins may be important as well.

Materials and Methods

Genetic Methods. Knockout strains were constructed in the W303 genetic background by first swapping the *KanMX* cassette in the respective deletion strains from the knockout library collection (Open Biosystems, Huntsville, AL) (31) with *LEU2* by using a linearized marker-swap plasmid (32). The disrupted gene:marker cassette was then PCR amplified by using specific flanking primers and then used for one-step disruption in W303. In the case of the *DJPI* deletion, a disruption cassette plasmid was constructed that included the 5’ UTR-*LEU2*-3’ UTR in pBluescript (Stratagene, La Jolla, CA) by using standard protocols (33) and transformed directly into W303. $\Delta jji1$ (21), $\Delta sis1$ (34), and $\Delta zuo1$ (20) strains were previously described. $\Delta cwc23$ and $\Delta hlj1$ strains [from P. Ahlquist (University of Wisconsin, Madison) and J. Brodsky (University of Pittsburgh, Pittsburgh, PA), respectively] were back-crossed six times with W303. *In vivo* diphthamide biosynthesis was monitored by scoring viability of

yeast cells upon conditional expression of a galactose-inducible diphtheria toxin (*GAL-DT*) plasmid pLMY101 (35).

Construction of J Protein Overexpression Plasmids. ORFs corresponding to the full-length and J domain-containing fragments (see legend to Fig. 1) of all of the cytosolic J proteins were PCR amplified by using appropriate gene-specific primers and were cloned into 2 μ - or centromere (CEN)-based plasmids under different promoters (36). For regulated expression of Ydj1, the complete ORF was cloned in a tetracycline-repressible vector pCM184 (37). Selected J proteins were HA-tagged by in-frame cloning of DNA encoding a 3xHA tag after the insertion of a NotI site by QuikChange PCR (Stratagene) before the stop codon in each coding sequence. The 2 μ GPD-Y^J Jjj1 chimera was constructed by in-frame fusion of the DNA encoding amino acids 1–60 of Ydj1 with amino acids 61–128 of Jjj1 protein. For making 2 μ GPD-Y^J R-HA, a 32-aa sequence unrelated to the J proteins (LPPWWQQLALAASLVPLAWLSHQKHCPGLNLS), followed by a 3xHA tag, was inserted after codon 63 of Ydj1 by PCR sewing

using overlapping primers. This resulted into a 128-aa protein fragment containing the J domain of Ydj1, with the other segment having no relationship to a J protein. The 2 μ GPD-Y^J Jjj3 chimera was constructed by in-frame fusion of amino acids 1–63 of Ydj1 to amino acids 70–172 of Jjj3 by PCR sewing.

Other Methods. Total proteins were isolated by treating cells with 0.1 N NaOH and resuspended in SDS sample buffer (62.5 mM Tris·HCl, pH 6.8, 5% glycerol, 2% SDS, 2% β -mercaptoethanol, and 0.01% bromophenol blue). Protein was detected as previously described by using anti-HA mouse 12CA5 (Roche Biochemicals, Indianapolis, IN) and anti-Ydj1 rabbit antibodies (34). Quantification was done with ImageQuant software (Molecular Dynamics, Sunnyvale, CA).

We thank J. Brodsky and P. Ahlquist for yeast strains; R. J. Collier (Harvard Medical School, Boston, MA) for plasmid pLMY101; and A. Meyer, R. Aron, P. Yang, A. Andrew, and J. Marszalek for helpful discussions and comments on the manuscript. This work was supported by National Institutes of Health Grant GM31107 (to E.A.C.).

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