Hsp40/70/110 chaperones adapt nuclear protein quality control to serve cytosolic clients

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Misfolded cytosolic proteins are degraded by the ubiquitin proteasome system through quality control (QC) pathways defined by E3 ubiquitin ligases and associated chaperones. Although they work together as a comprehensive system to monitor cytosolic protein folding, their respective contributions remain unclear. To bridge existing gaps, the pathways mediated by the San1 and Ubr1 E3 ligases were studied coordinately. We show that pathways share the same complement of chaperones needed for substrate trafficking, ubiquitination, and degradation. The significance became clear when Ubr1, like San1, was localized primarily to the nucleus. Appending nuclear localization signals to cytosolic substrates revealed that Ydj1 and Sse1 are needed for substrate nuclear import, whereas Ssa1/Ssa2 is needed both outside and inside the nucleus. Sis1 is required to process all substrates inside the nucleus, but its role in trafficking is substrate specific. Together, these data show that using chaperones to traffic misfolded cytosolic proteins into the nucleus extends the nuclear protein QC pathway to include cytosolic clients.

Introduction

Proteins are involved in all cellular functions. Accordingly, intricate intracellular pathways have evolved that govern the synthesis, quantity, delivery, activity, and lifetime of their protein constituents. As they work in concert, these mechanisms are collectively termed the protein homeostasis (or proteostasis) network (Balchin et al., 2016). Their importance is underscored by the prevalence and diversity of human disease when they become compromised (Labbadia and Morimoto, 2015). Although specialized mechanisms heighten the fidelity of information flow from genes to proteins, rare errors emanating from DNA replication, transcription, and translation can result in aberrant protein products. In addition, stochastic deviations from protein folding and maturation processes contribute to the total load. Because misfolded proteins can be cytotoxic through interference with cellular functions, quality control (QC) pathways are positioned throughout to monitor the folding state of nascent and preexisting polypeptides. Molecules beyond repair are targeted for degradation via the ubiquitin–proteasome system (UPS) or in lysosomes (Amm et al., 2014; Brandman and Hegde, 2016; Casson et al., 2016; Voos et al., 2016; Preston and Brodsky, 2017).

Aberrant proteins exist in two basic forms: aggregate and soluble. Protein aggregates, both ordered and amorphous, are associated with various human diseases including Alzheimer’s, Huntington’s, and Parkinson’s (Currais et al., 2017; Dubnikov et al., 2017; Pearce and Kopito, 2018). Although it remains controversial whether large aggregates are harmful or protective, cellular mechanisms exist to counter them. They can undergo targeted degradation through autophagy or the UPS (Gamerdingner et al., 2009; Scior et al., 2016). Alternatively, they can be found sequestered as ubiquitin-modified and/or chaperone-associated macromolecular structures (Kopito and Sitia, 2000; Kaganovich et al., 2008). Intracellular segregation presumably reduces their toxicity by limiting inappropriate interactions. Although many misfolded proteins have the tendency to aggregate on their own, they can be maintained in a soluble state through direct chaperone interactions (Balchin et al., 2016). Substrate solubility is particularly important for protein QC pathways that rely on the UPS (Amm et al., 2016; Comyn et al., 2016). In the better-understood ER-associated degradation (ERAD) pathways, luminal substrates must translocate across the ER membrane, and integral membrane proteins must be extracted for ubiquitination and degradation by the 26S proteasome (McCaffrey and Braakman, 2016; Preston and Brodsky, 2017). ER proteins forming large aggregates cannot use ERAD and are instead degraded in lysosomes through an autophagic mechanism (Kruse et al., 2006; Jackson and Hewitt, 2016).

Currently, the best understood protein QC mechanism is glycans-directed ERAD (Caramelo and Parodi, 2015; Cherepanova © 2018 Prasad et al. This article is distributed under the terms of an Attribution–Noncommercial–Share Alike–No Mirror Sites license for the first six months after the publication date (see http://www.rupress.org/terms/). After six months it is available under a Creative Commons License (Attribution–Noncommercial–Share Alike 4.0 International license, as described at https://creativecommons.org/licenses/by-nc-sa/4.0/).
Chaperones manage misfolded proteins in the nucleus

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Results

Ydj1 and Sse1 chaperones mediate trafficking of misfolded cytosolic proteins to the nucleus

The cytosolic/ER membrane J class chaperone Ydj1 is required for CytoQC, but its role in the pathway is unclear (Park et al., 2007; Fang et al., 2014; Maurer et al., 2016). To understand its contribution, we analyzed three model substrates in conditional and null mutants of YDJ1 (Caplan et al., 1992). Δ2GFP and ΔssPrA are San1-dependent substrates, and Ste6*C is a Ubr1-dependent substrate (Prasad et al., 2010, 2012). The conditional ydj1-151 mutant strongly stabilized each substrate at the restrictive temperature as expected (Fig. 1). At the permissive temperature, substrate stabilization remained strong in mutant cells. This suggests that the 37°C growth restriction reflects a loss of function other than San1/Ubr1-dependent CytoQC, which is nonessential (Park et al., 2007; Eisele and Wolf, 2008; Heck et al., 2010; Prasad et al., 2010; Maurer et al., 2016). We also observed that the higher incubation temperature also reduced substrate degradation rates in the WT cells (Fig. 1, A, C, and E). This likely reflects competition from endogenous substrates that increase under stress conditions or the formation of aggregates that are more difficult to degrade (Fang et al., 2014). In all cases, stabilization was accompanied by a proportional reduction in polyubiquitination, indicating that Ydj1 functions at or upstream of this step (Fig. 1, B, D, and F).

Because CytoQC is a dynamic process, we examined whether Ydj1 plays a role in substrate trafficking. For this question, we first performed indirect immunofluorescence to localize substrate constructs in WT and mutant cells. In WT cells, Δ2GFP and ΔssPrA were enriched in the nuclei as expected (Fig. 2, A and B). Ste6*C displayed a similar pattern, which was interesting because it was unclear where Ubr1-dependent substrates are degraded (Fig. 2 C). Intriguingly, all three substrates displayed a similar pattern, which was interesting because it was unclear where Ubr1-dependent substrates are degraded (Fig. 2 C). This likely reflects competition from endogenous substrates that increase under stress conditions or the formation of aggregates that are more difficult to degrade (Fang et al., 2014). In all cases, stabilization was accompanied by a proportional reduction in polyubiquitination, indicating that Ydj1 functions at or upstream of this step (Fig. 1, B, D, and F).

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nuclear proteins are unaffected (Fig. S1 D). Together, these data suggest a role for Ydj1 in substrate trafficking into the nucleus. To test the hypothesis, the superfolder (sf) GFP variant was appended to Δ2GFP to analyze substrate dynamics in live cells. sf-Δ2GFP behaves indistinguishably from Δ2GFP except that it more readily forms large intracellular foci at 37°C, reminiscent of structures termed insoluble protein deposits (IPODs) and juxtanuclear QC compartments (JUNQs; Figs. S1, E–G; Kaganovich et al., 2008). Live WT, ydj1-151, and Δsse1 cells expressing sf-Δ2GFP were given a high-intensity laser pulse to bleach fluorescent proteins in the nucleus. Our analysis was extended to cells lacking the Hsp110 factor Sse1 because these cells suggested a defect in substrate trafficking from immunofluorescence images (Prasad et al., 2010). The experiment was performed at 25°C to avoid the formation of the intracellular foci. Fluorescence intensities over photobleached areas were then measured at short intervals over a 100-s span and plotted (Fig. 3 A). Nuclear sf-Δ2GFP fluorescence recovered within a minute, revealing a significant lag between import and degradation. In both mutant strains, import was strongly impaired (Fig. 3, B and C). These data show that the...

Figure 1. Ydj1 is required for both San1 and Ubr1 pathways. (A, C, and E) All constructs were appended with a C-terminal HA epitope tag. The turnover of Δ2GFP, ΔssPrA, or Ste6*C in WT and ydj1-151 cells was determined by metabolic pulse chase. Cells were grown to log phase at indicated temperatures before labeling. Each strain was pulse labeled for 10 min and chased for the times indicated. After immunoprecipitation (IP), proteins were resolved by SDS-PAGE and quantified by phosphorimager analysis. Representative phosphor screen scans are shown. All data plotted were processed using Excel, reflecting three independent experiments with the means and SD indicated. (B, D, and F) Δ2GFP, ΔssPrA, or Ste6*C expressed in WT and ydj1-151 cells was immunoprecipitated, resolved by SDS-PAGE, and analyzed by immunoblotting (IB) to detect the ubiquitinated proteins. Protein amounts used for immunoprecipitation were normalized using an Odyssey infrared imaging system.
Figure 2. Localization of misfolded cytosolic proteins is defective in ydj1-151 cells. (A–C) WT and ydj1-151 cells expressing ΔssPrA, Δ2GFP, and Ste6°C were grown to log phase at room temperature and incubated at 37°C for 30 or 60 min. Cells were prepared for indirect immunofluorescence as described in Materials and methods. Substrates were detected using anti-HA antibodies in the green channel. ER and nuclear envelope were visualized in the red channel using anti-Kar2 antiserum. Nuclei were localized using DAPI staining. Cell imaging and acquisition were performed by confocal microscopy. Quantification of fluorescence intensity was done as described in Materials and methods. One-way ANOVA was used to test for significance (35 < n < 50; ****, P < 0.0001). The results shown are representative of two independent experiments. Bars, 2 µm.
Figure 3. **Misfolded protein nuclear trafficking is defective in YDJ1 and SSE1 mutants.** (A) Nuclear import of sf-Δ2GFP in WT, ydj1-151, and Δsse1 cells was analyzed by FRAP. Nuclei were photobleached (100% 488-nm laser transmission; 15–20 iterations) after three images, and recovery was monitored by acquiring images immediately after bleaching (postbleach). Images show cells before bleaching (prebleach), immediately after bleaching (postbleach), and 9, 30, 60, and 90 s after the initial bleach (postbleach). Bars, 2 µm. (B) The recovery of nuclear localized sf-Δ2GFP was plotted over the time. The fluorescence intensity (F. I.) of sf-Δ2GFP was quantified using ImageJ. Unpaired t test: *, P < 0.033; **, P < 0.002; ***, P < 0.0002; ****, P < 0.0001. n = 10. (C) Import rates were calculated using a pseudo–first order association kinetics curve using the model equation

\[ Y = Y_0 + (Y_{indef} - Y_0) \times [1 - e^{-K \times t}] \]

n = 10. The result shown is representative of three independent experiments. Error bars indicate means ± SEM.
import of misfolded cytosolic proteins into the nucleus is rapid and that the Ydj1 and Sse1 chaperones are required for this step.

**Ubr1 is a nuclear E3 ubiquitin ligase**

The dependence of both substrate classes on Ydj1 suggests a shared mechanism in the cytosol, which deviates only after nuclear import (Fig. 1). Although San1 is an established nuclear enzyme, Ubr1’s location was unclear because of its low abundance, but it is often described as a cytosolic protein in yeast (Amm et al., 2014). However, some fraction of Ubr1 must be in the nucleus because that is the location of some of its folded substrates. Applying two independent algorithms, Ubr1 is predicted to be a nuclear protein containing a strong nuclear localization signal (NLS; Fig. 4 A; Brameier et al., 2007; Kosugi et al., 2009). To test the prediction, we applied complementary experimental approaches. First, functional FLAG-tagged Ubr1 was expressed in WT cells using the strong TDH3 promoter to enhance detection (McAlister and Holland, 1985). For known nuclear proteins, expression from this promoter does not cause mislocalization (McAlister and Holland, 1985). Using indirect immunofluorescence, overexpressed Ubr1 and San1 was revealed by immunostaining. ER membranes were visualized using Kar2 antibodies and nuclei by DAPI staining. Arrowheads indicate the positions of nuclei. Bars, 2 µm.

**Hsp40/Hsp70/Hsp110 chaperones adapt nuclear QC for cytosolic proteins**

We next sought to better understand how misfolded proteins selectively traffic into the nucleus. For this, monopartite NLSs were appended to Δ2GFP and AssPrA to generate Δ2GFP-NLS and AssPrA-NLS, respectively (Fig. 2 D). As shown in Figs. 5 D and S2 E, Δ2GFP-NLS and AssPrA-NLS are exclusively nuclear localized in WT and mutant cells. Because the appended substrates no longer depend on CytoQC for import, their analyses can reveal factors specifically required for the import and/or degradation of misfolded cytosolic proteins. In pulse–chase experiments, NLS substrates require San1, Ubr1, and Ssa1/Ssa2, like their cytosolic counterparts (Figs. 5 A and S3 A). In contrast, the Sse1 chaperone, required for Δ2GFP and AssPrA (Prasad et al., 2010), is entirely dispensable for their NLS counterparts (Figs. 5 A and S3 A). This shows that the function of Sse1 in Δ2GFP and AssPrA CytoQC is exclusively for their nuclear import. Similarly, Δ2GFP-NLS bypasses the degradation block of Δ2GFP in ydj1-1S1 cells (Figs. 5 A and S3 A).
B and S3 B). As ubiquitination and degradation of Δ2GFP-NLS is only slightly impaired in ydj1-151 cells, the data indicate that like Sse1, the primary role of Ydj1 in CytoQC is in the nuclear import step (Fig. 5, B and C; and Fig. S3, B and C).

The efficient degradation of NLS-marked substrates in ydj1-151 background suggested a role for alternative Hsp40-like proteins. Of the 22 J domain proteins in budding yeast, we examined mutants of those localized in the nucleus or cytosol (Kampinga and Craig, 2010). Other than YDJ1, only SIS1 displayed a role in cytosolic/nuclear protein QC. Its role in CytoQC was already demonstrated previously in mediating the nuclear transport of the model protein CG* (ΔssCPY*-GFP fusion protein). It is also the target of polyglutamine aggregates, which helped explain the toxicity of such disease-related proteins (Park et al., 2013). Sis1 is also involved in localizing aberrant proteins to stress-inducible foci comprised of chaperones and misfolded proteins (Malinovska et al., 2012). In this study, analysis was performed in cells with SIS1 expression controlled by the tetracycline-responsive promoter (Tet-Off). Sis1 levels in Tet-Off SIS1 cells were 10–15% the level of WT even in the absence of doxycycline (Fig. 6, A and B; Summers et al., 2013). When misfolded proteins were expressed, these strains grew like WT (Fig. S5 D). At the same time, nuclear localization of San1 and Ubr1 was completely intact in these strains (Fig. S5, A and B). Thus, in this strain, reduced Sis1 impairs degradation

Figure 5. **Appending an NLS to Δ2GFP bypasses requirement for Ydj1 and Sse1.** (A and B) Turnover of Δ2GFP-NLS in WT, Δsan1, Δubr1, Δsan1Δubr1, Δssa1Δssa2, Δsse1, and ydj1-151 cells was determined by pulse chase at temperatures indicated. All data plotted were processed using Excel, reflecting three independent experiments with the means and SD indicated. (C) Ubiquitination of Δ2GFP-NLS was determined as described in Fig. 1 B. IB, immunoblot; IP, immunoprecipitation. (D) Cells expressing Δ2GFP-NLS were grown to log phase at room temperature and shifted to 37°C for 30 or 60 min. After fixation, Δ2GFP-NLS and Kar2 were visualized by immunostaining with respective primary and secondary antibodies. Nuclei were marked by DAPI staining. Arrowheads indicate positions of nuclei. Bars, 2 µm.
Sis1 is required for CytoQC and nuclear QC but dispensable for Δ2GFP nuclear import. (A and B) Cycloheximide decay experiments were performed in WT cells (R1158) and Tet-Off SIS1 cells expressing Δ2GFP or Δ2GFP-NLS in the absence and presence of doxycycline (DOX; 10 µg/ml, 20 h treatment before the experiment). Total protein extract was prepared as described above, and immunoblotting (IB) was performed to quantify the protein level of substrates. Sis1 and Pgk1 were used as loading controls. All data plotted were processed using Excel, reflecting three independent experiments with means and SD indicated. (C and D) The ubiquitination of Δ2GFP or Δ2GFP-NLS was determined in cells with or without SIS1 expression as described in Fig. 1 B. IP, immunoprecipitation. (E and F) The cellular localization of Δ2GFP or Δ2GFP-NLS in cells with or without SIS1 expression was examined by indirect immunostaining as described in Fig. 2. Arrowheads indicate positions of nuclei. Asterisks indicate the position of a nonspecific band. Bars, 2 µm.

Discussion

Of the broad classes of protein QC pathways, CytoQC is emerging as the most complex. In budding yeast, the numbers of E2, E3, and E4 enzymes of the UPS and chaperones involved in CytoQC are numerous and continue to grow (Park et al., 2007; Eisele and Wolf, 2008; Kohlmann et al., 2008; Lewis and Pelham, 2009; Heck et al., 2010; Prasad et al., 2010; Fang et al., 2011, 2014; Khosrow-Khavar et al., 2012). Some proteins traffic to the nucleus for processing, whereas others are degraded in the cytosol (Prasad et al., 2010; Park et al., 2013; Fang et al., 2014; Amm and Wolf, 2016). The reasons for the complexity remain unclear. The rationale to the complexity of CytoQC can only emerge through the coordinate analyses of the various pathways and their clients.
The discovery of San1 as a key CytoQC ubiquitin ligase upended the nascent field because San1 is a nuclear enzyme active in nuclear protein QC (Lewis and Pelham, 2009; Heck et al., 2010; Prasad et al., 2010). These studies led to the demonstration that misfolded cytosolic proteins traffic into the nucleus, where they are ubiquitinated and degraded (Heck et al., 2010; Prasad et al., 2010; Park et al., 2013). Such behavior is consistent with proteasomes being distributed mainly in the nucleus in dividing yeast cells (Russell et al., 1999). A nuclear import mechanism was confirmed and extended with additional San1–dependent substrates. The CG* substrate (ΔssCPY*–GFP fusion) traffics into the nucleus for ubiquitination and degradation (Heck et al., 2010; Stolz et al., 2013). Importance of this mechanism is disrupted in the presence of polyglutamine protein aggregates, providing a biochemical basis for the toxicity of polyQ disease proteins. This study further demonstrates a nuclear import–based CytoQC mechanism in mammalian cells, indicating that the basic pathway is conserved.

Although previously thought to be unique, we now know that the cytosol-to-nucleus mechanism is comprised of multiple pathways. This view was born out of the expansion of model CytoQC substrates. Indeed, an early hint came from the discovery that Ura3 models (Ura3-2/3 and Ura3-CL1) use the Doa10 E3 ubiquitin ligase (Metzger et al., 2008; Lewis and Pelham, 2009). This was notable because Doa10 also mediates the degradation of some integral membrane proteins by ERAD (Swanson et al., 2001; Huyer et al., 2004; Vashist and Ng, 2004). Although it is not yet determined where the Ura3 substrates are ubiquitinated and degraded, Doa10 is found in the inner nuclear envelope and the ER (Deng and Hochstrasser, 2006). Doa10 certainly functions in the nucleus for QC because a misfolded variant of the kinetochore protein Ndc10 (Ndc10-2) requires Doa10 for its degradation (Furth et al., 2011). Thus, Doa10 is involved in ERAD, CytoQC, and nuclear protein QC, making it the most versatile E3 in protein homeostasis. Whether its CytoQC mode also requires the trafficking of substrates into the nucleus remains to be determined.

Ubr1 is yet another CytoQC ubiquitin ligase that straddles pathways. The RING-type enzyme is best known as the E3 ligase of the well–characterized and conserved N–end rule degradation pathway (Bartel et al., 1990; Varshavsky, 2001). Here, the recognition mechanism is through substrate N-terminal sequences that directly bind pockets within Ubr1 (N degrons; Choi et al., 2010; Matta-Camacho et al., 2010). Ubr1 works with the HECT-type Ufd4 E3 enzyme, and together, the complex plays a role in QC (Hwang et al., 2010; Nillegoda et al., 2010). Although this recognition was first characterized for folded proteins, Ubr1’s QC mode also uses substrate N-terminal sequences for recognition (Kim et al., 2014). However, why it incorporates the N–end rule for the recognition of misfolded proteins was enigmatic. More recently, it was shown that some proteins bearing Ubr1 N degrons are normally stable (Shemorry et al., 2013). Their stability is caused by “shielding” of N degrons by interacting partners or ligands. Should such partners be limiting, the N degron is exposed and recognized by Ubr1, thus underlying a mechanism of QC for protein complexes.

Because the location of Ubr1 was previously unclear, misfolded Ubr1 substrates were thought to be ubiquitinated and degraded in the cytosol (Heck et al., 2010; Stolz et al., 2013). The data in this study support the view that Ubr1 is mainly a nuclear enzyme, and some of its substrates—misfolded proteins in particular—traffic into the nucleus for ubiquitination and degradation (Fig. 7). This explains the substrate overlap between San1 and Ubr1 (Lewis and Pelham, 2009; Heck et al., 2010; Khosrow-Khavar et al., 2012; Prasad et al., 2012; Guerriero et al., 2013; Amm and Wolf, 2016). Despite its preponderance as a nuclear enzyme, a fraction of active Ubr1 functions in the cytosol as some substrates are ubiquitinated there (Heck et al., 2010; Stolz et al., 2013). Perhaps diagnostic to identify such substrates is their exclusive dependence on Ubr1 because San1 has the intrinsic ability to recognize any unfolded protein in the nucleus (Rosenbaum et al., 2011).

Although the San1, Ubr1, and Doa10 pathways provide surveillance during normal conditions, stress conditions that increase protein misfolding require additional pathways. The Rsp5 and Hul5 ubiquitin ligases play specialized roles in targeting heat-damaged proteins for degradation (Fang et al., 2011, 2014). Although Rsp5 is involved in diverse cellular processes, during heat stress, it partners with the Ubp2 and Ubp3 deubiquitinases to efficiently degrade misfolded proteins (Fang et al., 2016). Interestingly, Rsp5 is also involved in the clearance of damaged membrane proteins that have already trafficked out of the ER (Haynes et al., 2002; Wang et al., 2011). Thus, it follows the theme of CytoQC ubiquitin ligases that participate in multiple pathways. Some damaged proteins localize to large intracellular complexes called JUNQs and IPODs under heat stress (Kaganovich et al., 2008), so it would be interesting to learn whether Rsp5 and Hul5 are constituents of these sites. San1, Ubr1, and Doa10 are less likely to function at cytosolic JUNQs and IPODs because they are localized primarily in the nucleus. However, it is possible they play a role in the processing nuclear inclusions (Miller et al., 2015). San1 and Ubr1 also play significant roles during heat stress (Khosrow-Khavar et al., 2012). Thus, the nuclear route is
active under broad conditions, whereas the Rsp5/Hul5 ligases may be better suited to process more aggregate-prone proteins.

There is broad agreement that cytosolic and nuclear chaperones are important for CytoQC in budding yeast (McClellan et al., 2005; Park et al., 2007, 2013; Metzger et al., 2008; Lewis and Pelham, 2009; Heck et al., 2010; Stolz et al., 2013; Comyn et al., 2016). Although it is known that the Hsp70 Ssa1/Ssa2 and the Hsp40 Sis1 proteins traffic some misfolded cytosolic proteins to the nucleus, how other chaperones act was less clear (Prasad et al., 2010; Park et al., 2013). A simple modification of established substrates with NLSs to bypass the CytoQC import step provided some clarity. These experiments showed that Ssa1/Ssa2 and Sis1 are also required for degradation after nuclear import (Figs. 5 A and 7). The Hsp110 protein Sse1, however, is only required for the import step (Figs. 5 A and 7). Similarly, Ydj1 is required for the import step, but its role in the nucleus is less clear (Fig. 7). Although the degradation of Δ2GFP-NLS is efficient in ydj1Δ1151 cells, it is delayed somewhat compared with WT (Fig. 5 B). Because ydj1 mutant cells grow poorly compared with WT, it’s possible that the small difference is attributable to one of Ydj1’s critical housekeeping functions.

Collectively, these data clarify the functional relationship between San1 and Ubr1, which were previously linked mostly through substrate overlaps. Their shared localization in the nucleus helps to explain much of the previous observations. In this study, we show that chaperones serve to traffic their cytosolic substrates to the nuclear protein QC system for processing (Fig. 7). This critical step suggests that chaperones make the first decision for which proteins should be degraded. However, the established mechanisms of substrate recognition by San1 and Ubr1 suggest that they comprise an additional step that improves stringency. San1 functioning through direct recognition of misfolded structures while Ubr1’s binding of N-terminal degrons (Rosenbaum et al., 2011 Kim et al., 2014) might also broaden the scope of potential substrates. San1 recognizes grossly misfolded proteins of all types, whereas Ubr1 recognizes orphan subunits whose N termini are normally shielded by partners or ligands (Shemorry et al., 2013). Thus, the N degron mechanism of Ubr1 would be particularly important for orphan subunits that do not grossly misfold. The purpose of segregating QC E3 enzymes to the nucleus could be as simple as to prevent promiscuous targeting of newly synthesized folding polypeptides, activities that occur in the cytosol. Thus, the complexity of the multistep process quickly can be seen as elegant and efficient if it is used also to protect newly synthesized proteins in their most vulnerable state. To accomplish this by making use of a standalone QC system for nuclear proteins makes the system even more efficient.

Materials and methods
Plasmids used in this study
Plasmids were constructed using standard cloning protocols (Sambrook et al., 1989). All genes encoding expression constructs were verified by DNA sequencing. All substrates contained an engineered single HA epitope tag at their C terminus. Unless otherwise noted, expression plasmids were constructed by placing coding sequences under the control of the constitutive TDH3 promoter in yeast centromeric vectors. Plasmid descriptions and oligonucleotide primers used in plasmid construction are listed in Tables S2 and S3, respectively.

pRP86, pRP88, and pRP120: Δ2GFP-HA, ΔssPrA-HA, and Ste6ΔC-HA fragments were released from pRP44, pRP42, and pRP22 (Prasad et al., 2012), respectively, by digesting the plasmids with BamHI and XbaI. Digested fragments were placed under the control of the TDH3 promoter in pRS316 vector to generate pRP86 and pRP88. pRP90: the SV40 NLS sequence SPKKKR KVEASGS was added to C terminus of Δ2GFP-HA by site-directed mutagenesis (Sawano and Miyawaki, 2000) using primers RP163 and pRP44. pRP91: an SV40 NLS sequence was added to the C terminus of ΔssPrA-HA by site-directed mutagenesis using primers RP164 and pRP42. pRP92 and pRP93: Δ2GFP-HA-NLS and ΔssPrA-HA-NLS fragments were released from pRP90 and pRP91, respectively, by digesting the plasmids with BamHI and XbaI. Digested fragments were ligated into pRP86 to generate pRP92 and pRP93, respectively. pRP96: an sfΔ2GFP-HA fragment was released from pSKI72 (Ng Laboratory plasmid collection) by digesting the plasmid with BamHI and XbaI. The digested fragment was ligated into pRP44 to generate pRP96. pRP97: an sfΔ2GFP-HA fragment along with ACT1 terminator was released from pRP96 by digesting the plasmid with BamHI and SphiI. The digested fragment was placed under the control of the TDH3 promoter in pRS314 vector to generate pRP97.

Strains and antibodies
S. cerevisiae strains used in this study are described in Table S1. Anti-HA monoclonal antibody (HA.11, raised in mouse) and antimyc monoclonal antibody (9E10 c-myc, raised in mouse) were purchased from Covance. Anti-Kar2 and anti-Sec61 antibodies (rabbit) were provided by P. Walter (University of California, San Francisco, San Francisco, CA). Anti-Sis1 antiserum (rabbit) was a gift from D. Cyr (University of North Carolina, Chapel Hill, NC). Monoclonal antiproteasome 20S α (mouse) and polyclonal anti–histone H3 (rabbit) were purchased from Abcam. Monoclonal anti–3-phosphoglycerate kinase antibody (mouse) and monoclonal anti-V5 antibody (mouse) were purchased from Invitrogen, monoclonal anti-Ydj1 antibody (mouse) was from StressMarq, monoclonal anti-GFP antibody (mouse) was purchased from Roche, and monoclonal anti–FLAG (mouse) was purchased from Sigma-Aldrich. Secondary antibodies labeled with Alexa Fluor 488 (anti–mouse) or Alexa Fluor 596 (anti–rabbit) were purchased from Molecular Probes. Anti–rabbit IRDye 680 and anti–mouse IRDye 800 secondary antibodies were purchased from LI-COR Biosciences. Anti-Ubr1 antiserum was raised in rabbit against a recombinant protein containing the 100 N-terminal amino acids of Ubr1. Anti-San1 antiserum was raised in rabbit against a protein containing the 100 C-terminal amino acids of San1.

Metabolic pulse–chase assay
Three OD600 units of log-phase cells were labeled with 82.5 µCi of [35S]methionine/cysteine (EasyTag EXPRESS [35S]; PerkinElmer) and chased with excess cold amino acids (final concentration, 2 mM methionine/cysteine) at the times indicated. Protein extract preparation, immunoprecipitation, and SDS-PAGE analyses were performed as described. Representative data are shown.
carried as described previously (Prasad et al., 2010). In brief, 100 μl of 100% TCA was directly added to 900 μl cell culture to terminate the metabolic labeling at each time point (e.g., 0, 30, and 60 min). Cells were disrupted with zirconium beads. The whole-cell lysate was transferred to a new tube and subjected to 10 min centrifugation at 16,000 g. The resulting pellet was resuspended in TCA resuspension buffer (100 mM Tris, pH 11.0, 3% SDS, and 1 mM PMSF; 50 μl/OD) and boiled at 100°C for 10 min. The insoluble fraction was removed after 10 min spin at 16,000 g. The volume of the soluble cell extract used for the following immunoprecipitation analysis was normalized based on the scintillation counting of total isotopically labeled proteins. Typically, we added ≤50 μl of total cell extract into solution containing 700 μl of IPS II buffer (13.3 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Triton X-100, and 0.02% sodium azide), 6 μl of 100 mM PMSF, 1 μl of protease inhibitor cocktail (1:200; Sigma-Aldrich), and 1 μl of specific antibody. After 1 h incubation at 4°C, the diluted extract was subject to 10 min centrifugation at 16,000 g. The soluble fraction was transferred a new tube, mixed with IgG agarose beads, and incubated at 4°C for an additional 2 h on a nutator plate. The precipitants were washed three times with IPS I buffer (0.2% SDS, 20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Triton X-100, and 0.02% sodium azide) and once with PBS. Lastly, the precipitants were eluted with protein loading buffer and analyzed by SDS-PAGE. The resulting gels were vacuum dried and exposed to phosphor screens for 24–48 h and then scanned and quantified using a Typhoon phosphorimager and ImageQuant TL software (GE Healthcare). All data plotted reflect three independent experiments with means and SD indicated. They were processed in Excel (Microsoft) with the AVE RAGE and STDEV functions.

**Cycloheximide chase assay and immunoblotting**

Cells were grown to mid-log phase in synthetic media. Cessation of protein synthesis was initiated by adding cycloheximide to 200 μg/ml to begin the chase. Aliquots of cells were collected at the times indicated, and total protein extract was prepared by TCA precipitation as described above. A portion of total protein extract was separated on a 4–15% gradient gel by SDS-PAGE and transferred to nitrocellulose membranes. After blocking, the membranes were incubated with appropriate primary antibodies and followed by fluorescence dye-labeled secondary antibodies (anti–mouse IRDye 800 and/or anti–rabbit IRDye 680). The protein level was quantified by the Odyssey infrared imaging system (LI-COR Biosciences). All data plotted reflect three independent experiments with means and SD indicated. They were processed in Excel with the AVERAGE and STDEV functions.

**Indirect immunofluorescence microscopy**

Indirect immunofluorescence was performed as described previously (Prasad et al., 2010) with minor modifications. In brief, log-phase cells were fixed with 3.7% formaldehyde at 30°C for 90 min. For temperature-sensitive mutants, cells were shifted to 37°C for 30 or 60 min and fixed at 37°C for 90 min by rotating in the incubator. Cells were spheroplasted by zymolysate digestion (1 mg/ml zymolase 20T [US Biological], 0.1 M potassium phosphate, pH 7.5, and 1.4 M sorbitol). Spheroplasts were applied to a poly-L-lysine–coated slide. Slides were sequentially immersed in methanol for 6 min and in acetone for 30 s at −20°C. After blocking with 5% BSA, cells were incubated with primary antibodies followed by secondary antibodies. HA.11 mAb (Covance) and polyclonal anti-Kar2 were diluted to 1:200 and 1:500, respectively. Secondary antibodies Alexa Fluor 488 goat anti–mouse and Alexa Fluor 596 goat anti–rabbit were diluted to 1:500. Nuclei were visualized with DAPI staining. Samples were examined at room temperature with a three-photon-multiplexer–tube detector using a 100× 1.4 NA oil Plan Apochromat objective in an upright confocal microscope (EXCITER or LSM510; ZEISS) controlled by LSM5 program software (ZEISS). Images were archived by LSM Image Browser and Photoshop (Adobe). For Fig. 2, an upright confocal LSM880 microscope using a 63× 1.4 NA oil Plan Apochromat (controlled by Zen 2012; ZEISS) with a tunable chameleon multiphoton laser, two conventional PMTs, and a high-sensitivity GaAsP detector (also with an airyscan detector) was used. For the analysis of images, background-subtracted images were quantified by drawing two regions of interest (ROIs) in the cytoplasm and one ROI in nucleus of each cell, and the ROIs’ mean fluorescence intensities were calculated using ImageJ (National Institutes of Health). Later for cytoplasm, means of two ROIs intensities were calculated. The reported relative nuclear enrichment was calculated as the ratio between mean nuclear and cytoplasmic fluorescence intensities. For drawing ROIs, the vacuole was avoided. Data of replicates were pooled before significance testing. One-way ANOVA was used to test for significance.

**Substrate ubiquitination assay**

The experiments were performed using mid-log-phase cells overexpressing ubiquitin. For temperature-sensitive mutants, cells were shifted to 37°C for 1 h before lysate preparation. 10 OD600 units of cells were collected, and total protein extract was prepared by TCA precipitation as described previously (Vashist and Ng, 2004). Protein levels were normalized before immunoprecipitation, and lysates were then mixed with IPS II (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, and 1% Triton X-100) supplemented with protease inhibitor cocktail (Roche), 100 mM PMSF, anti-HA affinity matrix, and 10 mM N-ethylmaleimide (Sigma-Aldrich). Immunoprecipitated samples were resolved by SDS-PAGE and transferred onto nitrocellulose. Total substrates and ubiquitinated substrates were detected by using anti-HA antibody and anti-myc antibody, respectively.

**GFP fluorescence microscopy**

Cells expressing sf-Δ2GFP were grown at room temperature and shifted to 37°C for 30 min. Cells were examined at room temperature using a Meta 510 confocal microscope under a 100× 1.4 NA oil-immersion objective lens (ZEISS). Images were archived by LSM Image Browser and Photoshop.

**FRAP**

For all FRAP experiments, fresh cells expressing sf-Δ2GFP were grown at room temperature, resuspended in fresh Synthetic Complete medium, and immobilized on a 2% agar pad containing Synthetic Complete medium. An LSM780 microscope (controlled by Zen 2012; ZEISS) with a multiframe 32PMT GaAsP detector and an Apochromat 63× 1.4 NA oil differential interference contrast...
Plan Apochromat objective was used for acquiring FRAP. Each nucleus was bleached with 15–20 iterations with 100% laser power of 488-nm light at 30°C. The FRAP protocol involved five prebleach images followed by bleaching and postbleach recording at 3-s intervals for 100 s. Imaging was typically performed by using a 488-nm argon laser at 3% laser intensity at 30°C. FRAP quantification was performed using ImageJ. The mean fluorescence recovery signal was quantified in the bleached nucleus. As a control, fluorescence of nonbleached nucleus of three neighboring cells was measured. After background subtraction, the fluorescence signals of nucleus were normalized to the means of the three control nuclei and set to 100% at beginning of the experiment. 10 different FRAP experiments were performed and transferred to Prism 6 (GraphPad Software) and plotted on an exponential FRAP curve. Nuclear import rates were calculated using nonlinear curve fitting of Prism software to a pseudo-first order association kinetics curve using the model equation 

$$Y = Y_0 + (Y_{\text{indef}} - Y_0) \times [1 - e^{-K \times t}]$$

to determine the apparent transport rate constant K for each strain. Images were also archived by ImageJ and Photoshop. All the error bars indicate ±SEM.

Statistics
A two-tailed unpaired Student’s test was used for significance of FRAP analysis. For Fig. 2, one-way ANOVA was used to test for significance. Unless otherwise indicated, all the analyzed mutants were always compared with WT.

Cytosol/membrane fractionation
Cells were grown to log phase (OD600 = 0.7–1) at 30°C. Cells (10 OD600 equivalent) were resuspended in lysis buffer (20 mM Hepes-KOH, pH 7.4, 250 mM sorbitol, 150 mM KOAc, and 5 mM MgOAc containing 1 mM PMSF and protease inhibitor cocktail). Glass beads (300 µl) were added, and cells were disrupted by agitation on a Vortex mixer (at full speed) 10 times for 30 s with 30-s intervals on ice between each cycle. The homogenate was collected and pooled after rinsing the beads with buffer 88 (20 mM Hepes, pH 7.4, 150 mM KOAc, 250 mM sorbitol, and 5 mM MgOAc). Unbroken cells were removed by centrifugation at 300 g for 5 min at 4°C twice. Subsequently, the supernatant was centrifuged at 18,000 g for 30 min at 4°C. The membrane pellet was resuspended in buffer 88, pH 7.4. The resulting supernatant (S18K) and the pellet (P18K) fractions were further processed for Western blotting.

Separation of cytosol and post-cytosolic fraction
10 OD600 units of log-phase cells were collected and resuspended in 1 ml Y-PER reagent (Thermo Fisher Scientific) containing 1 mM PMSF and protease inhibitor cocktail (Sigma-Aldrich) and then were incubated at room temperature for 30 min. Cells were centrifuged at 4°C for 30 min at 18,000 g to separate the cytosolic and post-cytosolic fractions. Each fraction was precipitated in 10% TCA and analyzed on a 4–15% gradient gel by SDS-PAGE, followed by immunoblotting.

Online supplemental material
The Supplemental material consists of three tables (Tables S1, S2, and S3) and five figures (Figs. S1, S2, S3, S4, and S5). Supplementary figures showed extended control experiments and data of additional examples to those shown in main figures. Tables list strains, plasmids, and oligonucleotide primers used in this study as well as their pertinent information.

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Author contributions: R. Prasad performed molecular, genetic, biochemical, and imaging analyses. R. Prasad and C. Xu performed quantitative analyses. D.T.W. Ng conceived and supervised the project. R. Prasad, C. Xu, and D.T.W. Ng wrote the paper.

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