Role for ribosome-associated complex and stress-seventy subfamily B (RAC-Ssb) in integral membrane protein translation

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Targeting of most integral membrane proteins to the endoplasmic reticulum is controlled by the signal recognition particle, which recognizes a hydrophobic signal sequence near the protein N terminus. Proper folding of these proteins is monitored by the unfolded protein response and involves protein degradation pathways to ensure quality control. Here, we identify a new pathway for quality control of major facilitator superfamily transporters that occurs before the first transmembrane helix, the signal sequence recognized by the signal recognition particle, is made by the ribosome. Increased rates of translation elongation of the N-terminal sequence of these integral membrane proteins can divert the nascent protein chains to the ribosome-associated complex and stress-seventy subfamily B chaperones. We also show that quality control of integral membrane proteins by ribosome-associated complex–stress-seventy subfamily B couples translation rate to the unfolded protein response, which has implications for understanding mechanisms underlying human disease and protein production in biotechnology.

Integral membrane protein ORFs make up ~20–30% of eukaryotic genomes (1). Membrane protein (MP)3 biogenesis is a highly coordinated, multiphase process in which many of the targeting steps are thought to be kinetically controlled. Early in membrane protein synthesis, the ribosome–nascent chain complex is identified by the signal recognition particle (SRP) and targeted to the endoplasmic reticulum (ER), for insertion of the MP into the ER membrane through the Sec translocon (2, 3). For membrane proteins, exposure of the first transmembrane helix (TM1) at the ribosome peptide exit site serves as the signal for SRP binding, although SRP can be recruited to the ribosome while TM1 is still inside the ribosome exit tunnel (4–8). SRP-target binding is also mediated by the nascent polypeptide–associated complex (NAC), which prevents nonspecific binding of SRP to non-secretory substrates and deters antagonistic interactions between SRP and N terminus–modifying enzymes during scanning of nascent polypeptides (9–11). These early targeting steps are crucial for proper membrane protein insertion into the ER and to prevent proteostatic stress due to the aggregation of the highly hydrophobic transmembrane helices of membrane proteins in the cytosol (5).

The contributions of kinetics in controlling nascent chain targeting have remained unclear. SRP-mediated attenuation of translation elongation has been proposed to extend the time window for SRP targeting in eukaryotes and is necessary for efficient translocation in human cells (12–14). However, a recent study using SRP-mediated selective ribosome profiling in yeast did not find evidence for general SRP-mediated pausing on secretory nascent chain substrates (7). Instead, computational analyses of mRNAs encoding membrane proteins and secreted proteins have identified clusters of rare codons that modulate their translation elongation rates to facilitate constructive interactions with SRP (15, 16). Tuller et al. (17) found that transcripts from several gene ontology categories, including those composed of secreted and membrane proteins, had a 5′-end slow translational ramp of 30–50 codons composed of non-optimal codons that optimizes ribosome spacing and reduces the chances of ribosomal “traffic jams.” Moreover, a genome-based analysis of N-terminal signal sequences in Sec-dependent secreted proteins from Escherichia coli showed that these sequences had a high frequency of non-optimal codons (18). Downstream from the N terminus, Pechmann et al. (16) showed through in silico sequence analysis and ribosome profiling data that secreted and membrane proteins had mRNA-
encoded slowdown of translation (REST) elements of non-optimal codons 30–40 codons downstream of the signal sequence/TM1 that assisted SRP interactions through slowing translation elongation. All of these studies relied on computational methods to discern patterns of codon usage bias. The earliest metric designed to detect this bias was the codon adaptation index (CAI), which is based on the codon usage frequencies of highly expressed genes in a given species (19). The CAI adequately predicts gene expression levels in *E. coli* and *Saccharomyces cerevisiae*, whose codon usage frequencies are positively correlated with isoaccepting tRNA abundances (20). Subsequently, a wide-ranging scale, the tRNA adaptation index (tAI), was introduced that better described the relationship between tRNA pools and codon usage patterns for unicellular and multicellular organisms. The tAI integrates the tRNA gene copy number and Crick’s wobble rules for codon–anticodon pairing (21). Global tAI has a strong correlation with high expression levels for cytosolic proteins, but this correlation is weak for membrane and secretory proteins in *S. cerevisiae* (15).

At the protein level, codon usage preferences also play an important role in regulating membrane protein function, folding, and structural stability. Several pathogenic synonymous polymorphisms in the human *MDR1* gene, which encodes the P-glycoprotein drug efflux transporter, have been identified that alter the structure and efflux activity of the transporter (22, 23). Misfolding and aggregation of secreted and membrane proteins trigger the ER quality control system characterized by two separate processes, the unfolded protein response (UPR), a signaling pathway that monitors ER homeostasis and responds to problems during membrane protein biogenesis at the ER, and the ER-associated degradation pathway (ERAD), which targets misfolded proteins in the ER lumen and membrane for proteasomal degradation (24–26).

Among the different classes of MPs, the major facilitator superfamily (MFS) is a ubiquitous family of secondary transporters that employ electrochemical gradients to drive substrate translocation through the membrane (Transporter Classification Database number 2.A.1 (27)) (28). It is the largest family of secondary transporters, including up to 25% of membrane transporters in prokaryotes (29). Most members range in size from 400 to 600 amino acids and are organized into 12 transmembrane α-helices. MFS transporters are important drug and therapeutic targets because they transport a variety of substrates, including simple carbohydrates, peptides, oligosaccharides, and lipids (30). These transporters can be divided based on their use of three distinct transport mechanisms: uniporters, symporters, and antiporters (28). The sugar porter subfamily is one of the largest and most widely studied subfamilies of MFS transporters and includes the human *SLC2* (GLUT) family of membrane transporters, the *HXT* family of hexose transporters from *S. cerevisiae*, and the oligosaccharide family of transporters in *Neurospora crassa*. The transcriptional regulation of these transporters has been extensively studied in relation to extracellular substrate concentrations and tissue specificities. However, codon preferences in mRNAs encoding these transporters and regulation of their proper targeting to the ER have not been studied to date (31–34). Here, we present experimental evidence for a new quality control mechanism that couples translation rates of the N terminus of MFS transporters to the chaperone machinery at the ribosome exit tunnel, which may contribute to proper SRP interactions with these ribosome–nascent chains.

**Results**

**Codon optimization of the *N. crassa* CDT-1 transporter**

The production of biofuels from plant cell walls requires the breakdown of cellulose into soluble glucose. However, the cellulases that cleave cellulose are inhibited by the product cellobiose, a β-1,4-linked dimer of glucose (35). We previously showed that heterologous co-expression of the *N. crassa* CDT-1 MFS transporter and intracellular β-glucosidase GH1-1 (NCU00801 and NCU00130, respectively) enabled *S. cerevisiae* cells to consume cellobiose directly, which could circumvent cellulase inhibition during biofuel production (36) (Fig. 1A). However, the CDT-1 transporter is slow relative to glucose transporters in yeast and remains a bottleneck in the cellobiose consumption pathway (37). To improve cellobiose consumption rates, we compared codon usage bias in the endogenous *cdt-1* sequence derived from *N. crassa versus* either the *N. crassa* genomic background or the *S. cerevisiae* genomic background. Using the %MinMax algorithm (38), which uses a sliding window to compare a coding sequence to the most common (%Max) or the most rare codons (%Min) from the genomic codon usage tables of a particular organism, revealed that the *Neurospora* coding sequence is optimal in *N. crassa* but suboptimal in *S. cerevisiae*, with most of the open reading frame composed of codons used less frequently in yeast (Fig. 1B). Other codon usage metrics, such as the CAI and the tAI, yielded similar results (supplemental Fig. S1) (19, 21, 39).

We therefore optimized the codons in the *cdt-1* sequence to better match the codon preferences of *S. cerevisiae*. After codon optimization of *cdt-1*, the CAI metric showed that the optimized version consists of the most common codons, whereas the *Neurospora*-derived sequence had much lower codon preferences (supplemental Fig. S1A) (19, 39). The tAI was also higher for the optimized *cdt-1* sequence (supplemental Fig. S1B) (21). Notably, improvement in the CAI and tAI still resulted in sequence identity at the nucleotide level for the *Neurospora*-derived and codon-optimized open reading frames of 76.2% (Clustal Omega alignment, supplemental Fig. S2).

**Codon-optimized CDT-1 causes a slow-growth phenotype**

We previously found that codon optimization of GH1-1 improved fermentation rates under aerobic and anaerobic conditions (40). However, we found that the codon-optimized version of the CDT-1 transporter (here denoted OPT for the mRNA and Opt for the expressed transporter) resulted in reduced cellobiose uptake and slower growth in cellobiose by *S. cerevisiae* relative to the transporter with *Neurospora*-derived coding sequence (hereafter NC for the mRNA, Nc for the expressed transporter) (Fig. 1C and supplemental Fig. S3). Additionally, in the course of preparing cultures for cellobiose growth assays, we noticed that single-colony seed cultures of *S. cerevisiae* cells constitutively expressing OPT grew significantly slower in glucose media than did their counterparts expressing NC, although both genes were expressed using the...
same vector with identical promoter and terminator (pRS316-PTHDS-CYC1tt), which also result in identical 5‘- and 3‘-UTRs. We therefore set up a growth assay in optimized minimal medium plus glucose and lacking uracil (oMM-Ura), which revealed that yeast expressing OPT grew 0.6 times the rate of cells expressing NC and 0.3 times the rate of cells carrying only the empty vector (Fig. 1D). Given these results, we suspected that codon-optimizing cdt-1 led to improper localization of the transporter or to protein aggregation. However, fluorescent microscopy of cells expressing either Opt or Nc transporter with eGFP fused to the C terminus revealed identical localization patterns for both proteins. The GFP fluorescence signal was present predominantly at the plasma membrane with very little to no intracellular fluorescence (Fig. 1E).

Figure 1. Codon-optimized CDT-1 and growth phenotypes. A, schematic of the cellobiose utilization pathway in S. cerevisiae. The CDT-1 transporter imports cellobiose into the cell, and the GH1-1 β-glucosidase hydrolyzes cellobiose into two glucose monomers, which then enter the glycolytic pathway. B, %MinMax profiles for NC using S. cerevisiae and N. crassa codon usage tables and %MinMax profile for OPT using the S. cerevisiae codon usage table. C, growth curves and growth rates for D452-2 cells expressing NC or OPT and GH1-1 in medium containing cellobiose as the sole carbon source, under aerobic conditions. Growth rates were calculated from the exponential growth phase and were 0.07 h⁻¹ for cells expressing NC and 0.05 h⁻¹ for cells expressing OPT. D, growth curves and growth rates for D452-2 cells expressing NC or OPT in glucose-containing medium. Growth rates were 0.16 h⁻¹ for cells transformed with empty plasmid, 0.1 h⁻¹ for cells with pRS316-NC, and 0.065 h⁻¹ for cells with pRS316-OPT. All growth rates shown represent the mean growth rate from three biological replicates, with the corresponding S.E. value (error bars). E, cellular localization of Nc and Opt. Epifluorescent microscope images of S. cerevisiae strain D452-2 cells with Nc-GFP (Nc) or Opt-GFP (Opt) transporters.
We wondered whether the transporter expressed from \( OPT \) was misfolded, but not severely enough to trigger ERAD and prevent trafficking to the plasma membrane. For example, \( MDR1 \) mutants with synonymous polymorphisms that led to conformational changes and impaired drug-efflux activity in the P-glycoprotein transporter had proper plasma membrane localization (23). The presence of a partially misfolded population could explain the lower cellobiose uptake rates and slower growth in cellobiose (Fig. 1C and supplemental Fig. S3).

To test for more subtle conformational destabilization of the CDT-1 transporter, we compared the thermal stability (\( T_m \)) of the transporter encoded by \( NC \) or \( OPT \) via fluorescence size-exclusion chromatography (FSEC) (41). In FSEC experiments, eGFP is fused to the C terminus of the target membrane protein to use GFP fluorescence as a proxy for absorbance at 280 nm during size-exclusion chromatography. Importantly, whereas the peak heights for Nc-GFP or Opt-GFP decreased with temperature, the height of the GFP-only peak decreased marginally, indicating that GFP did not unfold during these experiments (41) (Fig. 2A). Plotting the experiments as a thermal melt curve revealed the \( T_m \) value for both Nc and Opt transporters to be \( 38 \pm 0.1 \) °C (Fig. 2B). These results indicate that codon optimization did not affect the stability of the CDT-1 transporter significantly.
RAC-Ssb and integral membrane protein translation

We also tested whether codon optimization of cdt-1 may have resulted in substantially increased transporter protein expression levels that would tax cell resources, resulting in stress and slower growth. We measured Nc-GFP or Opt-GFP protein concentration in whole-cell lysates using GFP fluorescence and found that Opt protein levels were 1.5-fold higher than Nc levels (Fig. 2C). Protein concentrations for both transporters did not change between mid- and late-exponential phases ($A_{600}$ = 3 versus $A_{600}$ = 6; Fig. 2C).

Codon-optimized CDT-1 activates the unfolded protein response

To assess whether a specific proteostatic stress response pathway was associated with the slow-growth phenotype of cells expressing OPT, we examined the ubiquitous UPR (24). We constructed a UPR reporter strain using the yeast background strain used for cellbiose fermentations (D452-UPR), based on the single-cell fluorescence assay developed previously (42). Briefly, a bipartite UPR reporter cassette with GFP is expressed under the control of UPR elements, and RFP is expressed constitutively. The levels of GFP fluorescence reveal the level of UPR activation, and the levels of RFP fluorescence serve as internal controls for cell size and translation activity to normalize for cell-to-cell variation. UPR activation is represented as the median of log$_2$ GFP/RFP ratio for the entire population of cells sampled during flow cytometry.

We transformed plasmids pRS316-NC and pRS316-OPT, along with an empty pRS316 control plasmid, into strain D452-UPR. We then grew cultures to the same final $A_{600}$ of 3 (mid-exponential phase) and measured GFP and RFP levels using flow cytometry. Compared with controls with D542-UPR alone or treated with DTT to fully induce the UPR (42), both NC and OPT expression activated the UPR response, but the level of activation was significantly higher for OPT (Fig. 2D). UPR activation has been observed during expression of membrane protein mutants that destabilize the native structure or during overproduction of specific membrane proteins for crystallographic purposes (43, 44). Although Opt was not misfolded (Fig. 2B) and its concentration was not significantly higher than that of NC (Fig. 2C), activation of the UPR stress response suggests that post-transcriptional or translational events during expression of OPT may be responsible for the slow-growth phenotype.

The first 30 optimized codons of CDT-1 are responsible for the slow-growth phenotype

Nascent membrane protein insertion and folding in the ER membrane, as well as subsequent steps, such as TM insertion by the Sec61 complex, is under kinetic control (45, 46). Thus, synonymous codon changes that could lead to variation in translation elongation speed could affect this highly synchronized process (16, 47). SRP misrecognition or TM misinsertion could lead to activation of quality control pathways that result in nascent chain degradation and cellular stress (5). We therefore dissected the region of the OPT sequence responsible for the slow-growth phenotype to help pinpoint the step in membrane protein biogenesis that was affected by codon optimization.

We exchanged coding regions from the NC sequence into the corresponding regions in OPT, using the topology of MFS transporters, which have a basic structural unit consisting of four triple-helix bundles (48), as a guide. We divided the sequence into four regions, each encompassing one of these triple-helix motifs (Fig. 3A), using the Phobius TMH predictor to demarcate the sequence boundaries for each triple-helix bundle (supplemental Table S1) (49). We constructed four chimeras in which we sequentially switched the NC sequence for each triple-helix bundle into the OPT mRNA and used growth assays in glucose to detect whether switching any of these regions would rescue the slow-growth phenotype observed in cells expressing OPT. We first confirmed that all four transporters expressed from the chimeric mRNAs localized to the plasma membrane (supplemental Fig. S4). Remarkably, switching the first triple-helix bundle sequence in OPT for the NC counterpart entirely rescued the slow-growth phenotype (Fig. 3B). We next switched secondary structural elements individually in the first three-helix bundle region to yield seven different chimeras: one for the N-terminal tail region, three for the TMs, and three for the loops connecting TMs (supplemental Table S1). Only the N-terminal tail chimera (NC codons 1–72 fused to OPT codons 73–579) showed the same growth rate as cells expressing NC (Fig. 3C).

We further divided the first 72 codons of OPT into 30-codon overlapping windows and swapped in the corresponding NC sequences to make three chimeras. The region 1 chimera had NC codons 1–30; the region 2 chimera had NC codons 21–50; and the region 3 chimera had NC codons 41–72 (Fig. 4A). Growth assays in glucose showed that only cells expressing the region 1 chimera, with NC sequence at the extreme N terminus, had the same growth rate as cells expressing the intact NC sequence (Fig. 4A). To further test whether the extreme N-terminal sequence of OPT was responsible for the slow-growth phenotype, we carried out a reciprocal swap, switching the first 30 codons of the NC sequence for the coding sequence of OPT. The reciprocal swap of the OPT N-terminal coding sequence onto the remaining coding sequence of NC conferred the same slow-growth phenotype as OPT on cells expressing this chimera, as anticipated (Fig. 4B). The slow-growth phenotype of cells expressing OPT and the region 1 OPT/NC chimera also correlates with UPR activation in cells expressing these mRNAs (Fig. 5A). Thus, the first 30 codons, and possibly the first 20 unique to the region 1 chimeras, were sufficient to establish the slow-growth phenotype and UPR activation induced by codon optimization of CDT-1.

To test whether similar high-level expression of other transporters would induce the same growth and UPR effects, we chose two endogenous hexose transporters from S. cerevisiae, HXT1 and HXT5, that, like CDT-1, are members of the MFS sugar porter subfamily (Transporter Classification Database number 2.A.1.1) and are not aggressively degraded in our growth conditions (supplemental Fig. S5) (50, 51). Hxt1 is a low-affinity hexose transporter expressed under high-glucose conditions and is one of the major hexasaccharide transporters in S. cerevisiae (31). When expressed from a plasmid using the same expression cassette as NC and OPT, the growth rates for cells expressing HXT1 and HXT5 are quite different (Fig. 5B).
Cells expressing HXT1 from the plasmid had similar growth rates as OPT, whereas cells expressing HXT5 from the plasmid had higher growth rates, similar to NC. Importantly, these growth assays were performed using yeast strain BY4741, not D452-2, indicating that the slow-growth phenotype for OPT is strain-independent. The growth rates for the endogenous transporters were correlated with activation of the UPR response, as observed for NC and OPT expression (Fig. 5A). The protein concentration levels for Hxt1 and Hxt5, which were fused with eGFP at their C termini, were 2.3 and 1.6 times higher than the levels for Opt (Fig. 5C). Thus, although some endogenous transporters can induce the effects seen with OPT, there does not seem to be a correlation between transporter protein levels and slower growth rates.

The N terminus of CDT-1 with optimized codons does not impact growth when fused to a cytoplasmic protein

We also investigated whether the slow-growth phenotype caused by the first 20–30 codons of OPT was an inherent property of this particular nucleotide sequence or whether the phenotype was due to other properties, such as codon usage in this region. We fused the first 30 codons of NC or OPT to the N terminus of eGFP (NC30-GFP and OPT30-GFP) and expressed them with the same transcriptional regulation as NC and OPT. The growth rate of cells expressing these GFP fusions was identical (Fig. 6A), despite the protein concentration for the Opt30-GFP fusion being 2.3-fold higher than Nc30-GFP and 7-fold higher than Opt (Fig. 6B). These results indicate that the synonymous codon changes in OPT region 1 were only...
**Figure 4. Impact of the N-terminal codons of OPT on growth phenotypes.**

A, growth rates of cells expressing chimeras within the first 72 codons. The first 72 codons of OPT were divided into 30-codon overlapping windows, which were replaced with the corresponding NC sequences. REG1 corresponds to codons 1–30 from NC, REG2 corresponds to NC codons 21–50, and REG3 corresponds to NC codons 41–72 (black brackets). B, growth rate of cells expressing OPT codons 1–30 replacing the NC coding sequence (REG1 NC). Mean growth rates and S.E. values (error bars) from biological triplicates are shown.

**Figure 5. Impact of transporter expression on UPR activation and growth.**

A, UPR activation levels in cells expressing NC, OPT, and the REG1 OPT and REG1 NC chimeras. The level of UPR activation is higher in the presence of region 1 (codons 1–30) from the OPT sequence (REG1 NC) compared with region 1 of NC fused to the remainder of OPT (REG1 OPT). Mean levels are shown for five biological replicates, with corresponding S.E. values (error bars). B, growth rates of BY4741 cells expressing HXT1 and HXT5 from plasmids. Growth rates represent the mean growth rate and S.E. from three biological replicates. C, protein concentration of plasmid-expressed Hxt1, Hxt5, and Opt. The Hxt1 levels were 3.55 ± 0.16, Hxt5 levels were 2.45 ± 0.03, and Opt levels were 1.47 ± 0.04 μg of GFP/mg of total protein. HXT1 and HXT5 were expressed from the same expression cassette as NC and OPT.
The most striking feature of the footprint distribution is the high footprint count at codons 113–114 in both NC and OPT transcripts (Fig. 7C, blue box). This peak is 40–41 codons downstream of the predicted beginning of TM1 in CDT-1 and is possibly an encoded REST element (16). However, the coding sequences around the pause region for both NC and OPT (\(110\text{DTGPKVSV}^{117}\)) have high CAI values (0.761 for NC and 0.908 for OPT) (supplemental Fig. S6), i.e. both regions are composed of preferred codons (19), whereas REST elements have been predicted to be encoded by less-preferred codons (16). A more likely explanation for the peak is SRP-induced elongation attenuation after binding TM1 (6, 12, 14). Interestingly, this peak is attenuated in the OPT sequence, which suggests that not all of the ribosome–nascent chain complexes were recognized by SRP.

**The ribosome-associated complex interacts with Opt and triggers proteostatic stress**

Because the OPT phenotype was only present in the context of membrane proteins and did not affect cellular fitness during expression of cytosolic GFP chimeras, we wondered whether early membrane protein–targeting interactions at the ribosome could be affected by altered elongation rates through the N-terminal sequence. We noticed that the ribosome footprint density for the first 72 nucleotides of the OPT open reading frame was much lower than that on NC through this region (Fig. 7D and E) and supplemental Fig. S7). Because the mRNA levels were the same for both versions, the changes in footprint counts probably reflect faster translation through this region of the OPT transcript. For example, Yu et al. (56) observed the same depletion of ribosome footprints with synthetic luciferase constructs that contained regions with the most commonly used codons from *N. crassa*.

Notably in CDT-1, the first 30 amino acids precede the signal sequence recognized by the SRP by more than one length of the ribosome exit tunnel (TM1 in CDT-1 begins at codon 73). We therefore focused our attention on complexes upstream of the SRP recognition step (Fig. 8A). NAC modulates early interactions of SRP with nascent chains by preventing low-affinity binding to non-secreted substrates and facilitating nascent chain sampling by SRP (9–11, 57). NAC is a heterodimer of \(\alpha\)NAC (encoded by *EGD2*) and \(\beta\)NAC (encoded by *EGD1* or *BTT1*) (58). Egd1 and Egd2 form the predominant dimer species of NAC and interact with ribosomes translating secreted and membrane proteins (11). Due to the increased elongation speed in the first 24 codons of OPT (supplemental Fig. S7), we suspected that NAC could be misidentifying the Opt polypep-
RAC-Ssb and integral membrane protein translation

tide in the exit tunnel and blocking SRP from binding. To address this possibility, we expressed NC or OPT or an empty vector plasmid control in NAC knock-out strains from the Saccharomyces Gene Deletion Project (∆egd1 or ∆egd2) (Fig. 8A). We observed the same pattern of growth phenotypes on glucose as in WT strains, with cells expressing the empty plasmid

Figure 7. Ribosome footprint distributions for NC and OPT mRNAs. A, mRNA RPKM levels for NC and OPT transcripts. The transcript levels for NC were 530.4 ± 18.9 RPKM; for OPT, they were 502.3 ± 41.7 RPKM, determined from biological triplicates. B, rpM counts at the A site in the ribosome for both NC and OPT transcripts (these transcripts include the eGFP fusion). The x axis represents the nt position in the coding sequence. C, rpM counts for nucleotides 1–411 in NC and OPT. Shaded boxes represent the locations of TM1 and TM2. The blue box marks the pause before TM2. D, ribosome footprint distribution for the first 90 nucleotides of the NC and OPT open reading frames. Shown are the mean rpM counts from biological triplicates with their corresponding S.E. values. E, mean OPT rpM counts subtracted from the mean NC rpM counts (ΔrpM) and corrected for the S.E. (error bars).
RAC-Ssb and integral membrane protein translation

growing fastest, followed by cells expressing NC, and cells expressing OPT growing the slowest (Fig. 8B). These results indicate that NAC is not involved in the slow-growth phenotype induced by translation of the N terminus of Opt.

The other complex that is present at the ribosomal exit tunnel is the ribosome-associated complex (RAC). RAC and its cognate chaperones, the Hsp70 heat shock proteins Ssb1 and Ssb2, together form a chaperone triad and bind concurrently to the ribosome but are thought to have a different subset of client nascent chains compared with NAC, mostly nuclear and cytoplasmic proteins (11, 59), as well as ribosomal subunit assembly (60, 61). RAC is formed by the Hsp70 homolog Ssz1 and the J domain Hsp40 protein Zuo1 in S. cerevisiae (62, 63). Surprisingly, although RAC has not been reported to interact with SRP substrates (59), Δzuo1 and Δssz1 cells (Fig. 8A) expressing NC and OPT and grown on glucose had nearly identical growth rates and eliminated the slow-growth phenotype (Fig. 8C). This phenotypic rescue is specific to expression of NC and OPT, because the Δzuo1 and Δssz1 cells expressing the empty plasmid control grew slightly slower than WT (Fig. 8C). To confirm that this rescue was due to loss of the nascent peptide–Hsp70 chaperoning interaction, we assessed phenotypic changes in cells in which SSB1 or SSB2 or both were deleted (Δssb1, Δssb2, Δssb1/Δssb2), because RAC modulates cotranslational binding of Ssb1 and Ssb2 to client proteins (59) (59). We measured growth rates in glucose for Δssb1 and Δssb2 cells expressing NC or OPT and found that single deletion of either Hsp70 did not rescue the slow-growth phenotype in cells expressing OPT (Fig. 8D). This was not unexpected, as SSB1 and SSB2 are functionally redundant in yeast. However, deletion of both SSB1 and SSB2 (Δssb1/Δssb2), which had no impact on growth of NC-expressing cells, rescued the slow-growth defect in cells expressing OPT (Fig. 8D). As with the Δzuo1 and Δssz1 cells, the rescue is specific to expression of OPT, because the Δssb1/Δssb2 double deletion slowed growth of cells expressing the empty vector. The rescue observed in Δzuo1, Δssz1, and Δssb1/Δssb2 cells is also the opposite of that seen in RAC/Ssb deletion studies (64, 65), including those that identified RAC/Ssb involvement in ribosomal subunit assembly (60, 61). Taken together, these results indicate that recognition and binding of a subset of OPT-encoded proteins by the RAC-Ssb chaperone system led to proteostatic stress and decreased cell fitness. These results are consistent with the observed attenuation of the putative SRP pause in OPT transcripts after TM1 (Fig. 7C).

To test whether RAC binding to a subset of Opt polypeptides possibly leads to their degradation, we measured Opt levels in WT, Δdeg1, and Δzuo1 cells. Notably, Opt levels were highest in lysates from the Δzuo1 strain, whereas the Δdeg1 and WT Opt levels were similar (Fig. 8E). These results suggest that RAC recognition targets a portion of the Opt polypeptides for degradation, as deletion of one of the components of RAC increased productive protein levels, and the C-terminal GFP fused to Opt would only be made after translation of Opt. As noted previously, overexpression of HXT1 also led to a slow-growth phenotype and UPR activation, whereas HXT5 expression under similar conditions did not (Fig. 5). To test whether RAC may also interact with Hxt1, we transformed WT, Δegd1, and Δzuo1 cells with pRS316-HXT1, pRS316-HXT5, or an empty plasmid as control and measured their growth rates in glucose. Whereas the growth rate remained the same for all strains expressing HXT5 from the plasmid, WT and Δegd1 cells expressing HXT1 from the plasmid had much lower growth rates (Fig. 8F). Notably, as observed in Δzuo1 cells expressing OPT, Δzuo1 cells expressing HXT1 from the plasmid grew at nearly the rate of cells expressing HXT5 from the plasmid (Fig. 8F). Consistent with the observations with OPT-expressing cells, the phenotypic rescue in Δzuo1 cells expressing HXT1 is specific to expression of the transporter, as Δzuo1 cells expressing the empty plasmid exhibited slower growth. Furthermore, Hxt1 and Hxt5 levels were higher in the Δzuo1 strain and were similar in WT and Δegd1, whereas Nc levels remained the same in all three strains (Fig. 8G). RAC therefore interacts with Hxt1 in the same manner as it does with Opt ribosome–nascent-chain complexes.

*Ssb1 interacts with MFS transporter nascent chains at the ribosome before the emergence of TM1*

We performed Ssb1-mediated selective ribosome profiling to identify substrates and analyze how RAC–Ssb interacts with nascent chains at the ribosome (66). For this analysis, we compared ribosome profiling data from two samples generated from the same culture: the translatome set, which includes footprints for all ribosome–nascent chain complexes; and the Ssb1 interactome set, which includes footprints from ribosome nascent chains interacting with Ssb1 isolated after immunoprecipitation. We focused on the mRNAs for endogenous MFS transporters and were able to reliably identify eight transporters that interacted with Ssb1 during their translation (Fig. 9 and supplemental Fig. S8 and Table S4). Figure 9 shows the enrichment of HXT1 footprints in interactome samples versus the translatome samples. There are two regions that are enriched early in translation where Ssb1 would interact with the nascent polypeptide at codon positions 74 and 93, before TM1 exits the tunnel. These results indicate that Ssb1 interacted with nascent chains of Hxt1 before the substrate binding site of SRP emerged (TM1). Further, nascent chains from the other seven transporters that were identified (Arn3/Sit1, Arn4/Enb1, Hxt2, Hxt3, Itr1, Pho84, and Tna1; supplemental Fig. S8) had similar interactions with Ssb1; the chaperone bound very early in translation before or as TM1 was emerging from the peptide tunnel. We tried to survey other MFS transporters for interactions with Ssb1 (supplemental Table S5), but due to their inherent low expression, we could not obtain a reliable signal for these transcripts.

*Figure 8. Contributions of chaperone activities to transporter expression and growth phenotypes.* A, schematic of RAC and NAC interacting with the ribosome. NAC modulates SRP substrate recognition and chaperones a subset of nascent chains. RAC recruits Ssb to interact with nascent chain substrates and links aberrant proteins at the ribosome with the cytosolic network of chaperones (91). B, growth rates for WT, Δdeg1, and Δzuo1 strains expressing NC and OPT. C, growth rates for WT, Δssz1, and Δzuo1 strains expressing NC and OPT. D, growth rates for WT, Δssb1, Δssb2, and Δssb1/Δssb2 expressing NC and OPT. E, Opt protein concentration in WT, Δdeg1, and Δzuo1 whole-cell lysates. Protein levels were quantified via GFP fluorescence. F, growth rates for WT, Δdeg1, and Δzuo1 strains expressing plasmid-borne HXT1 and HXT5. All assays represent the mean and corresponding S.E. (error bars) from three biological replicates. G, Nc, Hxt1, and Hxt5 protein concentrations in WT, Δdeg1, and Δzuo1 whole-cell lysates. Protein levels were quantified via GFP fluorescence.
**Discussion**

Although ERAD serves as a quality control mechanism for membrane proteins post-translationally, less is known about quality control of membrane proteins before they are targeted to the translocon in the ER (25). Our results suggest that RAC-Ssb acts in the quality control of integral membrane proteins by actively sensing translation elongation rates of nascent membrane proteins on the ribosome. Indeed, cryo-EM reconstructions of RAC-80S ribosome complexes show that Zuo1 bridges the two subunits and extends from the ribosome peptide tunnel exit to the 40S subunit, where it interacts with the ES12 RNA helix (67).

RAC and Ssb are far more abundant than SRP in yeast cells, although both complexes interact dynamically with the ribosome (45, 68). For RAC and Ssb, there are ~27 and 89 complexes per 100 ribosomes in exponentially growing *S. cerevisiae* cells, respectively. By contrast, there are only about 1–3 SRP complexes per 100 ribosomes under the same conditions (68). Furthermore, the large hydrophobic sequences in TM1 would present ideal binding regions for Ssb (59). Thus, the RAC-Ssb complex could easily outcompete SRP for binding a subset of Opt and Hxt1 nascent polypeptides. Consistent with this model, we observed binding of Ssb1 to nascent Hxt1 and seven other MFS transporters polyepitides at the ribosome very early in translation (before or as TM1 emerges from the exit tunnel; Fig. 9 and supplemental Fig. S8). Our results indicate that, for some integral membrane protein substrates, RAC-Ssb may compete with SRP at the ribosome. Indeed, in cryo-EM reconstructions of SRP-80S and RAC-80S complexes, SRP and RAC bind to proximal regions of the 60S subunit, which would prevent their simultaneous binding to the ribosome (67).

Several studies have shown that SRP’s interactions with substrates and the SRP receptor are kinetically constrained (7, 45, 46, 69, 70). For example, SRP interactions with substrates containing a signal sequence fall off dramatically after 140 amino acids have been translated (70). Further, SRP targeting to substrates with weak signal sequences is sensitive to translation elongation rates (69). In the context of integral membrane protein biogenesis, translation elongation rates may affect the sensitivity of SRP binding to the first transmembrane segment, TM1. Taking into account the substoichiometric cellular concentration of SRP compared with ribosomes, higher translation elongation rates before TM1 emergence may favor RAC-Ssb binding and preclude later SRP interrogation of the emerging nascent polypeptide. Our results suggest that another function of the ubiquitous slow translation ramp in the first 30–50 codons of membrane proteins (17) would be to minimize RAC recognition of membrane proteins and subsequent sorting to Ssb. Although we did not measure translation elongation rates on *HXT1* and *HXT5* mRNAs, we predict that the N-terminal codon usage in *HXT1* has not been under evolutionary pressure to reduce translation speed. It will be interesting in the future to determine whether there is a negative correlation between endogenous MFS transporter expression level and translation speeds through the N-terminal sequence.

RAC-Ssb-bound Opt and Hxt1 nascent chains are probably targeted for degradation cotranslationally, as we observe increases in Opt and Hxt1 levels in Δzuo1 strains (Fig. 8, E and G). Wang et al. (71) observed cotranslational ubiquitination of nascent chains in actively translating ribosomes in human cells; however, the mechanism whereby these ubiquitinated nascent chains are delivered to the proteasome has not yet been elucidated to date. Our results suggest that the subset of Opt nascent chains targeted by RAC are likely to be cotranslationally ubiquitinated, because we observed an increase in the concentration of Opt transporters in strains where a component of RAC (Δzuo1) was deleted (Fig. 8E). Furthermore, systems-level analysis of *S. cerevisiae* nascent proteins targeted for cotranslational proteasomal degradation identified that these polyubiquitinated nascent chains generally had higher mRNA abundance and tAI, predictive of higher translation elongation rates (56, 72), than nascent polypeptides that were less susceptible to cotranslational ubiquitination (73). The cell may therefore use a quality control mechanism present at the ribosome to distin-

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**Figure 9. Ribosome footprint enrichment for *HXT1* from Ssb1-mediated selective ribosome profiling.** A, footprint distribution over the entire *HXT1* transcript. The plot shows the enrichment per codon position in footprint counts in the interactome samples over the total translatome samples for *HXT1*. The top schematic shows primary structure of Hxt1 with the predicted positions for the transmembrane domains (red boxes). The bottom schematic shows the length of the nascent chain in the ribosome at the peaks denoted by dashed lines. B, same plot enlarged. Light blue shading indicates variance between biological duplicates. AU, arbitrary units.
RAC-Ssb and integral membrane protein translation

guish nascent polypeptides headed toward non-native folding species even after chaperone intervention. Our results suggest that RAC-Ssb may be responsible for this role in the context of at least the MFS class of membrane proteins. When rapidly translated membrane proteins, such as Opt, are highly expressed, RAC-Ssb interactions with these nascent chains and the subsequent degradation of a subset of these polypeptides could overwhelm the basal level of protein quality control in cells, which would increase the proteasomal load and trigger cell stress and possibly the UPR (44, 71). Notably, transporters were highly enriched in the subset of proteins identified to be toxic in a genome-wide analysis of protein overexpression in yeast (74), consistent with our model for RAC-Ssb function and the relationship between OPT expression and cell growth (supplemental Fig. S9).

Our observations reveal another layer of complexity to expressing membrane proteins in heterologous systems to engineer novel metabolic pathways, especially when the host organism and the donor organism are phylogenetically distant. Even under autologous expression conditions, codon optimization of the human wild-type cystic fibrosis transmembrane conductance regulator resulted in lower protein yields (75). Current theoretical metrics that rely on codon and tRNA frequencies (CAI and tAI) may not effectively predict whether a codon-optimized membrane protein sequence will have higher translational elongation rates. Pechmann and Frydman (76) proposed a more comprehensive metric, which they termed the normalized translation efficiency (nTE) scale that takes into account the tAI for a specific sequence and the composition of the supply pool of tRNAs in the cell under a given growth condition. This scale may present a better method to predict translational efficiency under non-stressed conditions, and indeed, in a subsequent study, the authors showed that regions of low nTE after signal sequences in secreted proteins correlated with higher densities of ribosomal footprints (16). Nevertheless, under the stress conditions we observed in our experiments, the tRNA pool may vary drastically from normal conditions, and these changes must be taken into account to increase the predictive power of the nTE scale under stress conditions (77, 78). Previous reports have highlighted how silent polymorphisms in membrane protein-coding sequences linked to human disease affect protein folding and function (22, 23, 79). Our observations provide a connection between these observations, translational efficiency, and RAC-mediated quality control. The role of RAC-Ssb in membrane protein biogenesis also is likely to impact membrane protein expression in biotechnological applications.

Experimental procedures

Strains, plasmids, and media

The N. crassa and the codon-optimized versions of cdt-1 and the HXT transporters were cloned into the pRS316 plasmid (CEN URA) using the In-Fusion HD cloning kit (Clontech, Mountain View, CA). These vectors are pRS316-NC, -OPT, -HXTI, or -HXT5, respectively. The NC version of cdt-1 was PCR-amplified from cDNA synthesized from mRNA isolated from N. crassa (FGSC 2489) grown on minimal medium plus Avicel (microcrystalline cellulose) as the sole carbon source (36). Lifetech Geneart (ThermoFisher, Waltham, MA) performed gene synthesis and codon optimization for the optimized version of cdt-1. S. cerevisiae HXT transporters were cloned from gDNA isolated from strain D452-2.

For all chimeras described, the region to be swapped was PCR-amplified from the donor sequence and cloned into a plasmid containing the flanking sequences from the recipient gene using the In-Fusion HD cloning kit. All genes were expressed under the control of the S. cerevisiae TDH3 promoter and the CYCI terminator; all MFS transporters were fused with eGFP at the C terminus. Chimera 1 consisted of NC codons 1–170 and OPT codons 171–579; chimera 2 consisted of OPT codons 1–170 and 329–579 and NC codons 171–328; chimera 3 consisted of OPT codons 1–328 and 447–579 and NC codons 329–446; chimera 4 consisted of OPT codons 1–446 and NC codons 447–579.

S. cerevisiae strains BY4741, BY4741 deletion strains (from the Saccharomyces Genome Deletion Project at Stanford University (80)), D452-2 (81), and D452-2 UPR reporter were used in this study (supplemental Table S2). BY4741 Δssb1, Δssb2, and Δssb1Δssb2 were constructed for this study. In the case of the Δssb1 and Δssb2 strains, deletion of the respective gene was carried out following the same deletion design used to create the Saccharomyces genome deletion library. Briefly, the KanMX cassette was cloned from the gDNA of the Δzwo1 strain with primers that had 50-bp homology to the promoter and terminator regions of SSB1 and SSB2 and transformed into WT BY4741 cells. Δssb1 was then used as the parental strain to create the Δssb1Δssb2 deletion strain; SSB2 was deleted via CRISPR-Cas9 using a protocol developed previously (37). The D452-UPR reporter strain was constructed as follows. The UPR reporter was cloned from genomic DNA from strain YJW1200 provided by Jonathan Weissman’s laboratory at the University of California (San Francisco, CA) (42, 82). Details of reporter construction are provided in Ref. 42. The 4XUPRe-GFP-TEF2pr-RFP reporter cassette was inserted into the HO1 endonuclease locus of D452-2 using CRISPR-Cas9 (37). A list of primers for cloning and Cas9 guides is provided in supplemental Table S3. Yeast cells were transformed using the Frozen-EZ yeast transformation kit (Zymo Research Corp., Irvine, CA).

For yeast growth experiments, oMM–Ura (40) was used. oMM contained 10 g/liter (NH4)2SO4, 1 g/liter MgSO4·7H2O, 6 g/liter KH2PO4, 100 mg/liter adenine hemisulfate, 30 g/liter glucose or 20 g/liter cellobiose, 1.7 g/liter YNB (Sigma-Aldrich, Y1251), 2× recommended CSM–Ura dropout mix (MP Biomedicals, Santa Ana, CA), 10 mg/liter of inositol, 100 mg/liter glutamic acid, 20 mg/liter lysine, 375 mg/liter serine, 100 mM MES, pH 5.7.

Codon usage analysis

Codon usage tables for N. crassa and S. cerevisiae were obtained from the Codon Usage Database (http://www.kazusa.or.jp/codon/) (83). The NC sequence was analyzed using the %MinMax web server (http://www.codons.org/) (38). CAI calculations were carried out using the CAIcal server (http://genomes.urv.es/)

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CAIcal) (39). Per codon tAI was calculated from the nucleotide sequence using the tAI calculator (http://web.stanford.edu/group/frydman/codons/codons.html) (76).

**Growth assays**

All growth assays were performed using Bioscreen C (Oy Growth Curves Ab Ltd., Helsinki, Finland) with biological triplicates or quadruplicates. Single colonies of *S. cerevisiae* strains transformed with pRS316 containing the MFS transporter of interest were grown in oMM−Ura to late-exponential phase at 30 °C in 24-well plates. Cultures were pelleted at 4000 rpm, the spent medium supernatant was discarded, and cells were resuspended in H2O. The OD was measured to calculate the inoculum volume needed for 200-µl cultures at an initial OD of ~0.1 in Bioscreen plates. The OD at 600 nm was measured in 10-min intervals for 48–72 h at 30 °C. Growth curves were fit, and growth parameters were calculated using the grofit package in R (84).

**Microscopy**

D452-2 cells expressing NC or OPT were grown in oMM−Ura medium to mid-exponential phase at 30 °C. The cultures were centrifuged, spotted onto glass slides, and examined on a Leica DM 5000B epifluorescence microscope at ×100 differential interference contrast (Leica, Germany). Transporters were visualized using the L5 filter cube; images were captured using the Leica DFC 490 camera and analyzed with the accompanying microscope software.

**Yeast cell–based cellobiose uptake assay**

Yeast strain D452-2 cells transformed with pRS316-NC or pRS316-OPT were grown to mid-exponential phase. Cells were harvested and washed twice with Transport Assay Buffer (5 mM MES, 100 mM NaCl, pH 6.0) and resuspended to a final A590 of 40. 500 µl of cell resuspension was quickly mixed with an equal volume of Transport Assay Buffer containing 400 µM cellobiose. For the initial time point (t = 0 s), an aliquot was removed and centrifuged for 1 min at 4 °C at high speed to pellet the cells, and the supernatant was removed. The remaining cell resuspension was incubated at 30 °C for 30 min with constant shaking. After incubation, samples were centrifuged for 5 min at 4 °C at 14,000 rpm, and supernatant was removed. For analysis, 400 µl of supernatant was mixed with 100 µl of 0.5 M NaOH, and cellobiose concentrations remaining in the supernatant were measured on a Dionex ICS-3000 HPLC (ThermoFisher) using a CarboPac PA200 analytical column (150 × 3 mm) and a CarboPac PA200 guard column (3 × 30 mm) at room temperature. 25 µl of sample was injected and run at 0.4 ml/min with a mobile phase using 0.1 M NaOH.

**Protein concentration determination using fluorescence emission spectroscopy**

50-ml cultures were grown at 30 °C until they reached an A600 of 3, at which point they were harvested by centrifugation, resuspended in ~4–5 ml of growth medium, and aliquoted into microcentrifuge tubes to yield a total A600 of 30. Samples were spun down at 14,000 rpm for 1 min, and the supernatant was aspirated. Cell pellets were quickly flash-frozen in liquid N2. Frozen cell pellets were thawed on ice, and 400 µl of Buffer A (25 mM Heps, 150 mM NaCl, 10 mM cellobiose, 5% glycerol, 1 mM EDTA, 0.2× HALT protease inhibitor mixture (ThermoFisher), pH 7.5) was added for resuspension. Cells were lysed with zirconia/silica beads in a Mini-Beadbeater-96 (Biospec Products, Bartlesville, OK). Cell debris was pelleted at 10,000 × g for 10 min at 4 °C, lysates were diluted 3-fold with Buffer A, and their GFP fluorescence was measured using a Horiba Jobin Yvon Fluorolog fluorometer (Horiba Scientific, Edison, NJ). The λex was 485 nm, and the emission wavelength was recorded from 495 to 570 nm, with both excitation and emission slit widths set to 3 nm. A fluorescence calibration curve was prepared with eGFP purified from *E. coli* (>95% purity). The settings and the eGFP protein concentration range were chosen to yield a linear correlation between the fluorescence intensity at the maximum λem (510 nm) and the protein concentration of the standard. The maximum fluorescence intensity of the samples fell within this range. Target protein concentrations represent the mean from three biological replicates. Total protein concentration of the lysate was determined using the Pierce BCA protein assay kit (ThermoFisher).

**In-gel fluorescence detection**

BY4741 cells expressing pRS316-HXT1 and pRS316-HXT5 were grown to an A600 of 3, pelleted, and flash-frozen in liquid N2. Cells were resuspended in 400 µl of Buffer A and lysed using zirconia/silica beads in a Mini-Beadbeater-96. Cell debris was pelleted at 10,000 × g for 10 min at 4 °C, and total lysate protein concentration was measured using the Pierce BCA protein assay kit. Lysates were diluted to 2 mg/ml, 4× SDS-PAGE loading buffer was added, and 30 µl of samples was loaded immediately after loading buffer addition without boiling into a 4–20% SDS-polyacrylamide gel. A standard consisting of purified recombinant GFP was used as a size control. Gels were run at 100 V for ~1.5 h. After electrophoresis, the gel was rinsed with nanopure H2O and imaged using an ImageQuant LAS-4000 imager (GE Healthcare). The imager was set to the SYBR-SAFE fluorescence setting, with the light source set to 460 nm Epi, the excitation filter set to 515 nm, and the iris set to F 0.85.

**FSEC**

Three separate 0.8-liter cultures of oMM−Ura were inoculated with 10 ml of seed cultures of D452-2 cells transformed with either pRS316-NC or OPT. These cultures were grown to late-exponential phase at 30 °C and harvested at 4000 × g for 10 min at 4 °C. Cells were resuspended in 50 ml of Buffer A, flash-frozen in liquid N2, and stored at −80 °C. Cells were thawed on ice and lysed using 0.5-mm zirconia/silica beads in a BeadBeater (Biospec Products) at 4 °C. Cell debris was pelleted at 6000 × g, and the supernatant was centrifuged at 125,000 × g for 1 h at 4 °C. The supernatant was discarded, and the membrane pellet was resuspended via Dounce homogenization in Buffer A. Resuspended membranes were stored at −80 °C.

Hattori et al. (41) developed and optimized the thermostability FSEC analysis for membrane proteins fused with a C-terminal GFP fusion. The protocol used here is similar, with the following modifications. Membrane protein fractions (1600 µl) were solubilized by adding 10% n-dodecyl-β-D-maltoside to a final concentration of 2% and incubating at 4 °C for 1.5 h in a
rotating mixer. 90 μl was aliquoted into 15 thin-walled PCR tubes, and samples were heated to the target temperature for 10 min in an Applied Biosystems thermal cycler (ThermoFisher) and then cooled down to 4 °C. The thermal cycler lid temperature was kept at 100 °C. Five aliquots were then pooled and centrifuged at 100,000 × g for 30 min at 4 °C; 300 μl of clarified supernatant was collected (resulting in technical triplicates for each temperature). From the remaining non-heated sample, a 400-μl control sample incubated only at 4 °C was collected; this sample was also centrifuged at 100,000 × g after solubilization.

For FSEC, a Superose 6 10/300 GL SEC column (GE Healthcare) was attached to an Agilent 1200 HPLC system with a fluorescence detector module (Agilent Technologies, Santa Clara, CA). GFP fluorescence chromatograms were recorded at an excitation wavelength λ ex of 485 nm and an emission wavelength λ em of 512 nm, with the output set to 1 V and the photomultiplier tube gain to 12. 100-μl fractions were loaded onto the Superose 6 column; the flow rate was 0.4 ml/min, and the chromatography runs were carried out at 22 °C. Chromatograms were analyzed using the “ChemoStation for LC 3D Systems” software from Agilent Technologies. The peak area for the peak corresponding to CDT-1 was calculated; the fraction folded was calculated from the ratio of the peak area at the target temperature to the peak area of the corresponding control incubated at 4 °C. The FSEC-Tm values were determined as the temperature at which half the CDT-1 protein remained folded. All samples for a temperature set were run consecutively on the same day. Tm values were calculated via non-linear regression analysis after fitting the Boltzmann sigmoidal equation to the data.

Flow cytometry and UPR reporter measurements

All UPR reporter experiments were carried out using the D452-UPR reporter strain as described previously (42) with minor modifications. The UPR reporter strain was transformed with a pRS316 plasmid carrying the specified protein with a C-terminal GFP tag (P THO3 and CYC1 terminator). In these plasmids, an R97S mutation was introduced into eGFP to eliminate fluorescence (85). We confirmed that this mutation abolished GFP fluorescence by expressing transporter-(R97S)eGFP in D452-2 cells and then carrying out flow cytometry measurements.

Transformed colonies were grown overnight in oMM–Ura in 24-well plates at 30 °C. The following day, fresh oMM–Ura cultures were inoculated with these overnight seeds and grown at 30 °C to a final A600 of 3–4 (mid-exponential phase). Cultures were then centrifuged at 2500 × g, medium was discarded, and cells were resuspended in 1× PBS buffer. Flow cytometry measurements were carried out subsequently, and samples were analyzed in a BD Biosciences LSR Fortessa X20 (Flow Cytometry Facility, University of California, Berkeley, CA). The number of events was stopped at 20,000 counts. Data were collected using FITC parameter settings for GFP (488-nm laser, 505-nm LP filter) and PE-Texas Red for RFP (561-nm laser, 595-nm LP filter), FCS files were converted to CSV files using the “FcStoCsv” module in the GenePattern Server from the Broad Institute at MIT (86). These files provided the raw data for further analysis, which consisted of the pulse heights in the GFP and RFP channels for each event. The median for the log2 ratio of GFP/RFP for all events in a sample was used as a proxy for UPR reporter activation levels. All experiments were carried out using four or five biological replicates.

Cell harvesting for ribosome profiling and RNA sequencing

For both ribosome profiling and RNA sequencing procedures, the same cell cultures were used. 500-ml cultures of oMM–Ura were inoculated with D452-2 cells transformed with pRS316-NC or -OPT at 30 °C. Biological triplicates were concurrently grown for each condition (NC or OPT). Cultures were harvested when they reached an A600 of 3 via filtration using a Millipore filter device and a 0.8-μm filter (GE Healthcare). Cell pellets were quickly scraped off the filter and submerged in liquid N2; no cycloheximide was added to the cultures before harvesting. Polysome lysis buffer (20 mM Tris, 140 mM KCl, 1.5 mM MgCl2, 200 μg of cycloheximide, 1% Triton X-100, pH 8) was slowly dripped into liquid N2 tubes containing cells. Samples were stored at −80 °C until future use. Cycloheximide was added to the lysis buffer, and a portion of harvested cells was used for total mRNA sequencing.

RNA sequencing

The RiboPure yeast kit (Ambion, Austin, TX) was used to extract total RNA from 40 OD of cells following the manufacturer’s instructions. Four μg of total RNA was used to prepare multiplexed libraries with barcodes using the TruSeq RNA sample preparation kit (Illumina). The final cDNA libraries were quantified using an Agilent Bioanalyzer 2000 (Fuctional Genomics Laboratory, University of California, Berkeley, CA) and sequenced with the Illumina Genome Analyzer-II using standard Illumina operating procedures (Vincent J. Coates Genomic Sequencing Laboratory, University of California, Berkeley, CA). Sequence reads were assembled and analyzed in CLC Genomics Workbench version 6.5 (CLC Bio, Aarhus, Denmark). The genome for S. cerevisiae S288C (version R64.2.1), the parent strain of D452-2, was downloaded from Refseq at the NCBI (http://www.ncbi.nlm.nih.gov/refseq/); the mitochondrial genome was included. The sequences for pRS316-NC and -OPT were manually annotated and added to this file. Expression values were normalized by calculating the reads per kb of mRNA per million mapped reads (RPKM) and normalized further by using the option “By totals.” The reproducibility of RNA-Seq experiments is shown in supplemental Fig. S10.

Ribosome profiling

Preparation of ribosome profiling libraries followed protocols similar to those described previously (53, 54). Frozen cell pellets were lysed by low-frequency cryogenic mixer milling. Lysate was clarified of cell debris, and 25 A260 units of extract were treated with 450 units of RNase I (Ambion) for 1 h at room temperature with gentle rotation; digestion was stopped by the addition of 120 units of Superase-15. Ribosomes were pelleted using high-speed centrifugation through a 1 M sucrose cushion. The miRNeasy minikit (Qiagen) was used to purify ribosome-protected mRNA fragments following the manufacturer’s instructions. After size selection and dephosphorylation, a Universal miRNA cloning linker (New England Biolabs) was ligated to the 3′-end of footprints, followed by reverse transcription.
and circular ligation. Subtraction of rRNA fragments was performed using antisense biotinylated oligonucleotides for the most common rRNA fragment contaminants. After rRNA subtraction, the cDNA library was PCR-amplified. The final DNA libraries from biological triplicates for each condition (NC and OPT) were quantified using an Agilent Bioanalyzer 2000 (Fuc-

cional Genomics Laboratory, University of California, Berkeley, CA) and sequenced with the Illumina Genome Analyzer-II using standard Illumina operating procedures (Vincent J. Coates Genomic Sequencing Laboratory, University of California, Berkeley, CA).

For footprint analysis, trimmed sequencing reads without the linker were aligned to the S. cerevisiae ribosomal sequences using Bowtie (87). These reads were removed, and the non-rRNA reads were then mapped to the S. cerevisiae genome using Tophat (88). The sequences for the pRS316-NC/OPT plasmids were manually annotated and added to separate genome fasta files created for each condition and designated as new chromosomes. Only uniquely aligned reads were used for subsequent analyses. Most of the reads were between 27 and 32 nt long, and these reads were mapped onto their respective coding sequences as described previously (53, 54). The reproducibility of the ribosome profiling experiments is shown in supplemental Fig. S10. For rpM normalization, reads were normalized to the total number of reads mapped. For ΔrpM, the mean rpM for each position in OPT was subtracted from the mean rpM for the corresponding position in NC.

SSB-mediated selective ribosome profiling

Selective ribosome profiling library preparation and analysis have been described in detail (66). The total mapped reads per library were as follows: translatome 1, 10,504,232; translatome 2, 9,249,833; interactome 1, 13,161,962; interactome 2, 20,801,425. Genes with fewer than 64 raw reads in the translatome or interactome were excluded from the analysis. MFS transporters in the yeast genome are included in supplemental Tables S4 and S5. Ratios of RPM-normalized interactome and translatome data were built over a window of 15 nucleotides for both replicates. The average of both replicates was indicated using standard Illumina operating procedures (Vincent J. Coates Genomic Sequencing Laboratory, University of California, Berkeley, CA) and sequenced with the Illumina Genome Analyzer-II using standard Illumina operating procedures (Vincent J. Coates Genomic Sequencing Laboratory, University of California, Berkeley, CA).

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RAC-Ssb and integral membrane protein translation

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