Vms1p is a release factor for the ribosome-associated quality control complex

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Eukaryotic cells employ the ribosome-associated quality control complex (RQC) to maintain homeostasis despite defects that cause ribosomes to stall. The RQC comprises the E3 ubiquitin ligase Ltn1p, the ATPase Cdc48p, Rqc1p, and Rqc2p. Upon ribosome stalling and splitting, the RQC assembles on the 60S species containing unreleased peptidyl-tRNA (60S:peptidyl-tRNA). Ltn1p and Rqc1p facilitate ubiquitination of the incomplete nascent chain, marking it for degradation. Rqc2p stabilizes Ltn1p on the 60S and recruits charged tRNAs to the 60S to catalyze elongation of the nascent protein with carboxy-terminal alanine and threonine extensions (CAT tails). By mobilizing the nascent chain, CAT tailing can expose lysine residues that are hidden in the exit tunnel, thereby supporting efficient ubiquitination. If the ubiquitin-proteasome system is overwhelmed or unavailable, CAT-tailed nascent chains can aggregate in the cytosol or within organelles like mitochondria. Here we identify Vms1p as a tRNA hydrolase that releases stalled polypeptides engaged by the RQC.
effects that impair the decoding of a messenger RNA (mRNA) can cause ribosomes to stall and distort gene expression. To prevent the accumulation of potentially toxic aberrant polypeptides, and to maintain ribosome availability, eukaryotic cells employ surveillance and clearance mechanisms, including the evolutionarily conserved ribosome-associated quality control complex (RQC). The RQC is composed of the E3 ubiquitin ligase Ltn1p, the ATPase Cdc48p and its ubiquitin-binding cofactors, and the poorly characterized proteins Rqc1p and Rqc2p—whose homologs are Listerin, VCP/p97, TCF25, and NEMF, respectively, in mammalians.

Upon recognition of certain types of stalls, the 60S and 40S ribosomal subunits are split by the eRF1 homolog Dom34p, the GTPase Hbs1p, and the ATPase Rli1p. The RQC then recognizes and assembles upon the 60S ribosomal subunit that contains an incomplete polypeptide linked to a transfer RNA (tRNA) molecule (60S:peptidyl-tRNA). Ltn1p ubiquitinates the nascent chain and marks it for proteasomal degradation, a process facilitated by Rqc1p through unclear mechanisms. Rqc2p recognizes and binds to the exposed tRNA and stabilizes Ltn1p on the 60S. Rqc2p also recruits charged tRNAs to the 60S subunit to catalyze elongation of the nascent protein with Carboxy-terminal Alanine and mobilize potential ubiquitination sites, including lysine residues that are hidden in the ribosome exit tunnel, enhancing the efficiency of ubiquitination and the capacity of the RQC to protect cells from stochastic translation failures.

Nascent chain ubiquitination promotes extraction and proteasomal degradation of the aberrant polypeptide in a Cdc48-dependent process. If the ubiquitin–proteasome system is overwhelmed or unavailable, CAT tailing enables nascent chains to form aggregates in the cytosol or within organelles like the mitochondria. This non-templated elongation of the nascent chain can mobilize potential ubiquitination sites, including lysine residues that are hidden in the ribosome exit tunnel, enhancing the efficiency of ubiquitination and the capacity of the RQC to protect cells from stochastic translation failures.

Like the canonical RQC components, Vms1p is conserved throughout Eukarya and promotes protein quality control in diverse settings through its interaction with Cdc48p/p97. In Saccharomyces cerevisiae, Vms1p localizes to mitochondria in response to mitochondrial stress or damage. Mutants lacking Vms1p are sensitive to rapamycin, which impairs ribosomal efficiency of ubiquitination and the capacity of the RQC to protect cells from stochastic translation failures.

In light of the association between Vms1 activity and mitochondrial stress, we examined the ability of these single and double mutant strains to grow in glycerol medium, which requires mitochondrial respiration. Interestingly, deletion of LTN1—but not RQC1, RQC2, or DOM34—strongly impaired glycerol growth of vms1Δ cells (Supplementary Fig. 1b). Similarly, ski7Δ vms1Δ double mutant cells also exhibited partially impaired glycerol growth (Supplementary Fig. 1b). These data indicate a specific relationship between Vms1p, RQC function, and mitochondrial homeostasis, which is consistent with a recent report that stalled polypeptides that cannot be ubiquitinated by Ltn1p accumulate within and compromise mitochondria—underscoring the reversal of the genetic interaction we observed between LTN1 and VMS1 on glucose vs. glycerol.

These genetic interactions prompted us to determine whether Vms1p physically interacts with members of the RQC, as has been reported previously. As expected, isolation of Rqc2p-HA co-immunoprecipitated Vms1p-V5. In contrast, Rqc1p-HA or Ltn1p-HA showed minimal Vms1p interaction. Overexpression of Vms1p from the strong SKI7 promoter in galactose medium had no effect on the co-migration of Rqc2p with the 60S ribosomal subunit (Supplementary Fig. 1e, f). Vms1p is required for resolving RQC substrates. The genetic and physical interactions described above motivated us to evaluate whether RQC substrates accumulate in VMS1 mutant cells. We utilized a well-characterized mRNA that encodes FLAG-tagged green fluorescent protein (GFP) followed by a hammerhead ribozyme that self-cleaves in vivo (FLAG-GFPΔz, Fig. 2a). To generate a truncated mRNA encoding FLAG-GFP without a stop codon or poly-A tail. Translation of this mRNA triggers ribosome recycling by 60S:peptidyl-tRNA species expands upon the previously characterized function of Vms1p as a Cdc48p/p97 binding protein and raises interesting new questions regarding the potential interaction of protein and organelle homeostasis.

**Results**

**Vms1p physically and genetically interacts with the RQC.** Because vms1Δ cells are sensitive to rapamycin, we tested whether other perturbations to translation elicit a similar phenotype. We found that the vms1Δ strain is sensitive to the protein synthesis inhibitor cycloheximide (CHX), as are other RQC mutants in a sensitizing SKI mutant background. Surprisingly, deletion of any one of the RQC system components RQC1, RQC2, or LTN1, was sufficient to reverse the lethality of the vms1Δ mutant in CHX (Fig. 1a). By contrast, deletion of the no-go decay components DOM34 or SKI7 had no effect. These data suggest that CHX causes accumulation of an RQC product that becomes toxic in the absence of Vms1p.

The RQC assembles on a failed 60S subunit to both ubiquitinate and elongate nascent polypeptides with CAT tails. To determine whether one or both of these activities generate the putative toxic RQC product, we eliminated them separately by expressing a CAT-tailing-defective Rqc2p mutant (RQC2Δz, Supplementary Fig. 1f) or a variant of Ltn1p deficient in ubiquitination (LTN1Δz/LTN1Δz, Supplementary Fig. 1f). CAT tailing by Rqc2p was dispensable, while ubiquitination by Ltn1p was essential for conferring CHX sensitivity on a vms1Δ mutant strain—indicating that the toxic species in this context depends on ubiquitination rather than CAT-tailing (Fig. 1b). Presumably because of its proposed role in nascent chain ubiquitination, and similar to RQC2 and LTN1 (Fig. 1b), plasmid expression of wild-type (WT) RQC1 was sufficient to reverse the rescue of vms1Δ cycloheximide sensitivity conferred by RQC1 deletion (Supplementary Fig. 1f).

Vms1p is required for resolving RQC substrates. The genetic and physical interactions described above motivated us to evaluate whether RQC substrates accumulate in VMS1 mutant cells. We utilized a well-characterized mRNA that encodes FLAG-tagged green fluorescent protein (GFP) followed by a hammerhead ribozyme that self-cleaves in vivo (FLAG-GFPΔz, Fig. 2a). To generate a truncated mRNA encoding FLAG-GFP without a stop codon or poly-A tail. Translation of this mRNA triggers ribosome stalling and targeting to the RQC. Deletion of SKI7 enhances expression of GFP by inhibiting degradation of the cleaved mRNA. We confirmed that deletion of RQC1, RQC2, and LTN1 each lead to accumulation of FLAG-GFPΔz, whereas the nascent chain failed to accumulate in the ski7Δ single mutant (Fig. 2a–c, Supplementary Fig. 2a). Loss of Vms1p also led to accumulation of FLAG-GFPΔz to a level similar to that observed.
for the core RQC components (Fig. 2a–c, Supplementary Fig. 2b). Combination of VMS1 deletion with the deletion of RQC1, RQC2, and LTN1 had no additive effect on GFP accumulation (Fig. 2b). Immunoblot analysis showed similar results for the single, double, and triple mutant strains, in which RQC2-dependent, high-molecular-weight aggregates are also apparent (Fig. 2c, Supplementary Fig. 2b)19–21. Loss of DOM34 led to decreased accumulation of GFP fluorescence, even in the vms1Δ strain, consistent with the upstream role of Dom34p in ribosome splitting and suggestive of alternative pathways for degrading nascent chains when the Dom34p/Hbs1p subunit splitting activity is unavailable (Fig. 2a–c). Interestingly, accumulation of FLAG-GFP<sup>Rz</sup> in vms1Δ mutant cells occurs despite lower mRNA abundance (Supplementary Fig. 2c).

In addition to the FLAG-GFP<sup>Rz</sup> construct, which generates a cytosolic RQC substrate, we also examined RQC activity on fumarase, which is encoded by the FUM1 promoter, was also maintained at a low level in the ski7Δ mutant strain (Fig. 2d). Deletion of VMS1, RQC1, RQC2, or LTN1 each led to profound accumulation of GFP fluorescence, almost all of which colocalized with mitochondria-targeted red fluorescence protein (mtRFP). The vms1Δ, rqc1Δ, and ltn1Δ mutants, which retain Rqc2p and CAT-tailing activity, all exhibited Fum1-GFP aggregates within or near mitochondria, comparable to recent observations of other mitochondria-derived nascent chains28 (Fig. 2d). The rqc2Δ mutant exhibited a more uniformly mitochondrial localization pattern (Fig. 2d), consistent with the model that CAT tailing mediates intra-mitochondrial aggregation of polypeptides that stall during co-translational import28. Together, these data demonstrate that Vms1p is required for the degradation of substrates derived from truncated mRNAs, whether they are destined for a membranous organelle or the cytosol.

**Vms1p is structurally homologous to tRNA hydrolases.**

Understanding of how Vms1p facilitates the clearance of stalled translation products was guided by our recent crystal structure determination of a portion of S. cerevisiae Vms1p (Fig. 3a, b)31. This structure includes the highly conserved central region of Vms1p, which we named the mitochondrial targeting domain (MTD) because it is necessary and sufficient for mitochondrial localization32. This localization activity requires a hydrophobic groove along the region of the MTD where the LRS interacts and direct binding to ergosterol peroxide31. Intriguingly, the Vms1p MTD structure resembles structures of the catalytic domain of eukaryotic peptide chain release factor subunit 1 (eRF1), as well as Dom34p and RNaseE, which both resemble tRNA hydrolases32–35 (Fig. 3b, Supplementary Fig. 3a). The only region of the Vms1p MTD structure resembles structures of the catalytic domain of eukaryotic peptide chain release factor subunit 1 (eRF1), as well as Dom34p and RNaseE, which both resemble tRNA hydrolases32–35 (Fig. 3b, Supplementary Fig. 3a). The only region of the Vms1p MTD that diverges substantially from the release factor fold is the face of the MTD that mediates ergosterol peroxide binding and mitochondrial localization31 (Fig. 3b).

Sequence alignment showed that although Vms1p lacks a strict GGQ motif characteristic of eRF1p, it does possess an invariant glutamine that can align with the catalytic glutamine of eRF1 (Fig. 3c). In yeast Vms1p, this glutamine residue is
embedded within a GGSQ motif that is reminiscent of the eRF1 catalytic GGQ, while in other species the conservation other than the initial glycine and glutamine is less apparent (Fig. 3c, d).

These observations inspired us to determine whether Vms1p is required for resolving RQC substrates. We asked which residues and regions are required for the stalled 60S by hydrolyzing the ester bond anchoring them to tRNA. We enabled the extraction of failed translation products from the ribosome. This is consistent with Vms1p functioning in stop codon-independent tRNA hydrolysis within a 60S, rather than 80S, ribosome.

The observations inspired us to determine whether Vms1p enables the extraction of failed translation products from the stalled 60S by hydrolyzing the ester bond anchoring them to tRNA. We asked which residues and regions are required for the genetic functions of VMS1. We first tested an Hb8T motif just N terminal to the conserved ‘GxxQ’ motif that mediates ribosome interactions of eRF1 (Fig. 3c, d). Vms1p mutants of these residues, H279A, H283A, R284A, and T286A, were indistinguishable from WT, while the Y285A mutant exhibited a partial CHX sensitivity (Fig. 3f). In contrast, mutation of the ‘GxxQ’ residues G292 and Q295 and the highly conserved R288 residue conferred strong loss-of-function phenotypes (Fig. 3e). Deletion of S294 to convert the GGSQ of S. cerevisiae Vms1p into a GGQ motif, as in eRF1, also abrogated VMS1 function (Fig. 3e). While all of these ‘GxxQ’ mutants failed to confer resistance to 200 ng ml⁻¹ CHX, only the R288A and G292A/G293A mutants were inactive at the lowest (100 ng ml⁻¹) concentration of CHX tested (Fig. 3e).

Interestingly, both of these mutants also failed to rescue glycerol growth in an ltn1Δ vms1Δ double mutant, whereas wild-type VMS1 and the other mutants did rescue growth (Supplementary Fig. 3b). The R288A, G292A/G293A, and Q295L mutants also exhibited enhanced accumulation of FLAG-GFP² in the ski7Δ background similar to the vms1Δ mutant (Supplementary Fig. 3d).

Importantly, the Vms1p mutants interact normally, if not more strongly, with Rqc2p based on co-immunoprecipitation experiments (Supplementary Fig. 3e). In light of these observations, we
Fig. 3 Vms1p is structurally homologous to tRNA hydrolases. **a** Domain structure of Vms1p. LRS leucine-rich sequence, ZnF zinc finger, MTD/eRFL mitochondrial targeting domain/eRF1-like, AnkR ankyrin repeat, CC coil-coil, VIM VCP-interacting motif. Residues 188–417 represent the MTD/eRFL boundaries. **b** Structural alignment of Vms1p (left, 5WHG31) and eRF1p (middle, 3JAHii35, residues 144–280). Dashed lines indicate connections made by residues that are not resolved in the Vms1p crystal structure. The GGQ (red) loop of eRF1p is ordered in the ribosome-bound structure shown here. **c** Sequence alignment of Vms1p and eRF1p. **d** Sequence alignment of Vms1p orthologs across the GxxQ region. Coloring as in **c, e, f**. Serial dilutions of indicated strains were spotted on media containing glucose or glucose supplemented with cycloheximide (CHX). EV empty vector.
hereafter refer to the MTD as the MTD/eRFL domain, where eRFL refers to eRF1-like.

In addition to these loop residues, the ability of Vms1p to confer complete CHX resistance in both vms1Δ and ski7Δ vms1Δ also required the p97/valosin-containing protein (VCP)-interacting motif (VIM), which mediates interaction with Cdc48p/VCP/p97\(^2\) (Supplementary Fig. 3c). Interestingly, the VIM is not required for growth of the ltn1Δ vms1Δ double mutant on glycerol (Supplementary Fig. 3c), which indicates that mitochondrial homeostasis can be maintained even without Cdc48p binding.

**Vms1p exhibits tRNA hydrolase activity towards RQC substrates.** To directly test whether Vms1p catalyzes peptidyl-tRNA hydrolysis, we utilized our recently described *S. cerevisiae* in vitro translation (ScIVT) system to monitor the synthesis and fate of a robust stalling reporter and its peptidyl-tRNA intermediate\(^4\). RQC-intact extracts translate this reporter, split the stalled 80S ribosome into constituent 60S and 40S subunits, elongate the nascent chain with a CAT tail, and ubiquitinate exposed lysine residues. These extracts also hydrolyze the peptidyl-tRNA ester bond to generate the released polypeptide\(^5\) (Fig. 4a). We observed that extracts prepared from vms1Δ mutant cells also produced peptidyl-tRNA conjugates, but loss of the peptidyl-tRNA species and appearance of the released translation product were slower than in WT extracts (Fig. 4a, b). This is somewhat obscured by the fact that the vms1Δ mutant has lower overall translation, which leads to a decreased amount of the free nascent chain and peptidyl-tRNA conjugates. We performed a similar experiment in the ski7Δ and ski7Δ vms1Δ mutant strains and found that in the ski7Δ background the deletion of VMS1 conferred a much more obvious stabilization of the peptidyl-tRNA species and qualitatively delayed release of the polypeptides (Fig. 4c, d). In this ski7Δ background, deletion of RQC2 conferred a modest stabilization of the peptidyl-tRNA conjugate and had little effect on the vms1Δ mutant (Supplementary Fig. 4a). We next purified full-length and C-terminally truncated (1–417) *S. cerevisiae* Vms1p and found that each of these proteins dramatically accelerated the production of the released polypeptide in a dose-dependent manner in WT, rqc2Δ and vms1Δ extracts (Fig. 4e, Supplementary Fig. 4b). Importantly, the 1–417 truncation mutant lacks the C-terminal VIM domain and is unable to interact with Cdc48p (Supplementary Fig. 3e). Therefore, we conclude that while the Vms1p–Cdc48p interaction is important for CHX resistance and other RQC-related functions that are relevant to the VMS1 genetic interactions, Vms1p association with Cdc48p is dispensable for peptidyl-tRNA hydrolysis.

We next tested release factor activity of the MTD/eRFL domain structure-based mutants described above. The Q295L, G292A, and ΔS294 mutants also exhibited strongly impaired release factor activity (Fig. 4f and Supplementary Fig. 4c). Consistent with the stronger growth phenotypes we observed on different concentrations of CHX and on glycerol, R288A and G292A/G293A mutants had no hydrolysis activity, even at 10-fold higher concentration than the concentration at which WT Vms1p catalyzed complete tRNA release (Fig. 4f).

![Fig. 4](image-url) Vms1p exhibits tRNA hydrolase activity towards RQC substrates. a Time courses of *S. cerevisiae* in vitro translation (ScIVT) reactions prepared with a truncated mRNA (lacking a stop codon). Extract genotypes are indicated above. Peptides that have been CAT-tailed and released are denoted with a cat label. b Quantification of peptidyl-tRNA species in a. Mean ± s.e.m.; n = 6. \(**P < 0.0001\). The p value was calculated using a two-way ANOVA. c Time courses of ScIVT reactions prepared as in a. d Quantification of peptidyl-tRNA species in c. Mean ± s.e.m.; n = 8. \(****P < 0.0001\). The p value was calculated using a two-way ANOVA. e ScIVT reactions prepared as in a with a vms1Δ extract. At t = 15, buffer (–) or pure protein was added. Slopes indicate a titration series of decreasing protein concentrations (see Methods). FL full length Vms1, 1–417 N terminus through eRFL domain. f ScIVT reactions prepared as in a with a vms1Δ extract. At t = 15, buffer, WT (1–417) protein, or mutant (1–417) protein was added.
Discussion
Our data have identified a key constituent of the RQC pathway in Eukarya: a tRNA hydrolase that liberates failed polypeptides from the aberrant 60S:peptidyl-tRNA species that accumulate when ribosomes stall. Without this activity, translation products remain anchored in 60S ribosomes, which therefore cannot be recycled for future use. The dual functions of the Vms1p MTD/eRFL identified here as an RQC release factor and previously as a mediator of the resolution of peptidyl-tRNA conjugates of stalled nascent cause disease. We therefore propose that Vms1p is required for the resolution of ribosome subunits for future use and defends proteostasis from inadequate Ltn1p activity by releasing CAT tailed nascent.

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The diseases associated with even subtle distortions in protein quality control underscore the importance of the activity of Vms1p as a release factor for the RQC. Among many other examples, hypomorphic mutations in the RQC-associated ubiquitin E3 ligase, Listerin (Ltn1p in yeast), lead to profound neurodegeneration in mice. As we show, Vms1p protects cells from inadequate Ltn1p activity by releasing CAT tailed nascent chains from stalled 60S ribosomes. In so doing, Vms1p rescues these ribosome subunits for future use and defends proteostasis from translation products whose accumulation could otherwise cause disease. We therefore propose that Vms1p is required for the resolution of peptidyl-tRNA conjugates of stalled nascent chains in the cytosol as well as those destined for organelles like the mitochondria, where it might mediate a particular role in protecting mitochondria from proteostasis challenges.

Methods
Yeast strains and growth conditions. S. cerevisiae BY4743 (MATa, his3 leu2 met15 ura3) was used as the wild-type strain. Each mutant was generated in diploid cells using a standard PCR-based homologous recombination method. The genotypes of all strains used in this study are listed in Supplementary Table 1. Yeast transformations were performed by the standard TE/LiAc method and transformed cells were recovered and grown in synthetic complete glucose medium lacking the appropriate amino acid(s) for selection. The medium used included YPA and synthetic minimal medium supplemented with 2% glucose or 3% glycerol. Cycloheximide was added at a final concentration of 100 ng ml⁻¹ or 200 ng ml⁻¹ when indicated. All plasmid constructs were generated by PCR and cloned into the yeast expression vectors pRS413, pRS14 or pRS416 as indicated in Supplementary.

Fluorescence microscopy. WT (BY4741) or derived mutant strains were transformed with a plasmid expressing mitochondria-targeted (ATPase subunit, Su9) RFP, mRFP, and plasmids expressing FlagHiss-GFP-Rz or FUM1-FlagHiss-GFP-Rz under the GPD or native FUM1 promoter, respectively. The cells were grown to mid-log phase and imaged using the Axio Observer Z1 imaging system (Carl Zeiss). Digital fluorescence and differential interference contrast images were acquired using a monochromic digital camera (AxioCam MRm) and analyzed using the Zen software (Carl Zeiss).

Fluorescence-assisted cell sorting. GFP-expressing strains and untransformed control were grown to OD600~1 and pelleted by centrifugation at 100 × g for 5 min. Cell pellets were washed once in 1× phosphate-buffered saline (PBS) buffer, resuspended in 1 ml of 1× PBS, and analyzed using the BD FACSAria Canto Analyzer (488 laser and optical filter FITC). A total of 30,000 events were measured and the mean fluorescence of three independent biological replicates was analyzed by one-way analysis of variance (ANOVA; Bonferroni correction analysis) with a confidence interval of <0.05 using the statistics software: Graphpad Prism 6. Additionally, sk7Δ, sk7Δvms1Δ, and strains in Supplementary Fig. 3 were compared via unpaired Student’s t-test (two-tailed) confidence interval value set to p < 0.05. Error bars represent standard error of the mean.

Immunoprecipitations. The VMS1-VMS1-V5 (or VMS1 mutant) was co-expressed with an endogenous promoter-HisR, HA tagged RQC component (RCQ1, RCQ2, LTN1) in the cognate double mutant strain. Approximately 50 ODs were harvested in log phase and resuspended in IP buffer (20 mM Tris pH 7.4, 50 mM NaCl, 0.2% Triton X-100), vortexed 10 x 1 min, clarified via centrifugation, and added to anti-HA magnetic beads (Thermo Scientific R8836). After 4 h of incubation, beads were pelleted via magnet and washed 4x with 1 ml of IP buffer. Proteins were eluted with 50 µl of 2x Laemmli buffer (20% glycerol, 125 mM Tris-HCl pH 6.8, 4% SDS, 0.02% bromophenol blue). Source data can be found in Supplementary Fig. 5a.

False-coloring. Yeast cultures were grown to OD600~1, cycloheximide was added to a final concentration of 0.05 mg ml⁻¹, and cells were harvested by centrifugation 5 min later. Cell pellets were washed in buffer A (20 mM Tris-HCl pH 7.4, 50 mM KCl, 10 mM MgCl₂, 1 mM dithiothreitol (DTT), 100 µg ml⁻¹ cycloheximide, 1 × RNaseAcc (Ambion), and 1× yeast protease inhibitor (Sigma)). Pellets were weighed and resuspended in 1.3 volumes of Buffer A. An equal volume of glass beads was added and suspensions were vortexed for 30 s, followed by 8 times interspersed with 8 min of incubation on ice. Following centrifugation at 3000 × g for 5 min, supernatant was centrifuged at 11,300 × g for 2 min at 4°C, after which supernatant was centrifuged at 11,300 × g for 10 min. Protein extracts were overlaid onto a linear sucrose gradient of 15–50% and centrifuged at 234,600 × g for 3 h at 4°C. The gradients were pelleted through a continuous-flow chamber and monitored at 254 nm with an ultraviolet absorbance detector (ISCO UA-6) to obtain ribosomal profiles. Fractions (16) were collected, resuspended in 2× Laemmli sample buffer supplemented with 2.5% beta-mercaptoethanol, and analyzed by western blotting. Source data can be found in Supplementary Fig. 3b.

SDS-PAGE. Whole-cell extracts were prepared from 3 to 5 ODs of cells at OD600~1.5 by solubilization in 250 µl of 2 M LiAc, incubated for 8 min on ice followed by centrifugation at 0.9 × g for 5 min at 4°C. The pellet was resuspended in 250 µl of 0.4 M NaNO₃ and incubated on ice for 8 min followed by centrifugation at 16,000 × g for 3 min. Next, the pellet was resuspended in 1× Laemmli buffer with 2.5% beta-mercaptoethanol, boiled for 5 min, and centrifuged at 0.9 × g for 1 min. Supernatants were collected and loaded onto acrylamide/bisacrylamide (37.5:1) gels. Subsequent immunoblotting was done with the indicated antibodies: HA (PRB-101C-200), V5 (ab9116), FLAG (F7425), PK1: (ab113687), and Rpl3 (sc3RPL3). Source data can be found in Supplementary Fig. 6.

Protein expression and purification. For the His3 tagged proteins, constructs were transformed into JRY1734 (pep4::HIS3 prb1::LEU2 bar1::HISG lys2::GAL1/3), and strains in Supplementary Fig. 3 were compared via unpaired Student’s t-test (two-tailed) confidence interval value set to p < 0.05. Error bars represent standard error of the mean.
supplemented with 480 ng mRNA to initiate translation. Reaction aliquots were sampled at indicated time points by quenching in 2× Laemml Sample Buffer. Proteins were separated by SDS-PAGE, and hemagglutinin (HA)-tagged translation products were visualized by immunoblotting (Roche 3F10). To quantify release, the abundance of peptidyl-tRNA was measured with Fiji (https://imagej.net/Fiji) and normalized as percent of the initial 15 min time point. Mean values of at least 6 technical replicates were analyzed and plotted in Prism (GraphPad software). Values were calculated using a two-way ANOVA. Error bars represent standard error of the mean. The ramps of Fig. 4e represent a decreasing titration series of 4.2 μM, 0.42 μM, 0.21 μM, and 0.105 μM final protein concentrations. In Fig. 4f and Supplementary Fig. 4c, 1x and 1/10 refer to final protein concentrations of 4.2 μM and 0.42 μM, respectively. Source data can be found in Supplementary Fig. 7.

Quantitative RT-PCR. RNA was purified from 40 ml of yeast cultures grown to OD600 ~1. Pelleted cells were washed once with water and resuspended in 700 μl of Trizol reagent (Ambion). An equal volume of glass beads was added and suspensions were vortexed for 30 s with 1 min rest intervals. Next, the manufacture’s protocol of the Direct-zol kit (Zymo research: R2850-11-330) was followed. Complementary DNA (cDNA) was obtained from 0.5 μg of purified RNA using the High-capacity cDNA Reverse Transcription kit (4368841) from Applied Biosystems. Quantitative PCR was performed using the LightCycler 480 SYBR Green Master (4070516001) from Roche and using a FLA-HS and Actin primer pair. Quantitative PCR analysis was done by Absolute Quantification/2nd derivative of three independent biological replicates, each performed in triplicate. The statistical analysis of mRNA transcript abundance was done after normalization with Actin. The statistics software Graphpad Prism 6 was used to perform Student’s t-test (unpaired two-tail) with a confidence interval value of p < 0.05. Error bars represent standard error of the mean.

Data availability. The authors declare that the data supporting the findings of this study are available within the paper and its supplementary information files. Further relevant data on the genes studied in this manuscript (VMS1: YDR049W, RQC1: YDR333C, RQC2: YPL099C, LTN1: YMR246, SK7: YOR076C, DOM34: YNL01W) can be found at: https://yeastgenome.org.

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Author contributions
O.Z.R., E.K.F., C.J.H., C.P.H., J.R, and A.F. designed the study and wrote the manuscript. N.D.T. and J.V.V. ran the polysome assays. O.Z.R., E.K.F., and C.J.H. collected the data. J. V.V. and S.F. generated plasmid constructs and yeast strains. C.P.H. helped with structural analyses. R.K. performed structural homology modeling and alignments. B.A. O. helped with the IVT assays. P.S.S. helped with the co-immunoprecipitation experiments. All authors commented and approved of the final manuscript.

Additional information
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