The highly conserved 5′–3′ exonuclease Xrn1 regulates gene expression in eukaryotes by coupling nuclear DNA transcription to cytosolic mRNA decay. By integrating transcriptome-wide analyses of translation with biochemical and functional studies, we demonstrate an unanticipated regulatory role of Xrn1 in protein synthesis. Xrn1 promotes translation of a specific group of transcripts encoding membrane proteins. Xrn1-dependence for translation is linked to poor structural RNA contexts for translation initiation, is mediated by interactions with components of the translation initiation machinery and correlates with an Xrn1-dependence for mRNA localization at the endoplasmic reticulum, the translation compartment of membrane proteins. Importantly, for this group of mRNAs, Xrn1 stimulates transcription, mRNA translation and decay. Our results uncover a crosstalk between the three major stages of gene expression coordinated by Xrn1 to maintain appropriate levels of membrane proteins.
Proper tuning of protein levels under normal and perturbed conditions requires precise regulations at different stages of gene expression. These stages, classically considered isolated because of their different spatial and temporal incidence, are indeed interconnected. A major crosstalk between transcription and decay is mediated by Xrn1\textsuperscript{1–3}, a highly conserved exoribonuclease, which is the 5′-3′ messenger RNA (mRNA) degradation enzyme in the cytoplasm\textsuperscript{4,5}. Within the deadenylation-dependent decay pathway, Xrn1 forms a complex interaction network with the Dcp1/Dcp2 decapping enzyme and decapping activators, such as Lsm1–7, Pat1, and Dhh1/DDX6\textsuperscript{6,7}. At least one of the degradation activity of Xrn1 occurs co-translationally\textsuperscript{6}. Xrn1 further participates in the decay of mRNAs after internal cleavage and in the cytoplasmic mRNA surveillance system that degrades aberrant mRNAs\textsuperscript{7}. Moreover, Xrn1 directs degradation of long non-coding RNAs and hypomodified transfer RNA (tRNA), as well as maturation of ribosomal RNAs (rRNAs)\textsuperscript{8,9}. Remarkably, besides these exonucleolytic functions, Xrn1 acts as a transcriptional activator. Xrn1, together with other components of the deadenylation-dependent mRNA decay pathway, shuttles between the cytoplasm and the nucleus, where they bind to transcription start sites and directly stimulate transcription initiation and elongation of many yeast genes\textsuperscript{1}. The functions of Xrn1 in both cellular compartments are linked. Shuttling of Xrn1 and other decay factors to the nucleus depends on the proper exoribonucleolytic activity of Xrn1. By connecting mRNA synthesis to decay, Xrn1 maintains mRNA homeostasis, as defects in 5′-3′ mRNA decay are buffered by reductions in mRNA synthesis. Whether Xrn1 functions in other stages of gene expression has not been considered.

Here, we show that Xrn1 acts as a translational modulator. This unanticipated function is restricted to specific groups of genes enriched in distinct GO terms. Interestingly, Xrn1 activates both translation of mRNAs encoding membrane proteins and their localization at the endoplasmic reticulum, the translation compartment of membrane proteins. These mRNAs contain long and highly structured 5′UTRs. A physical and functional interaction of Xrn1 with the translation initiation factor eIF4G is required for translational activation, likely to overcome these unfavorable contexts for translation initiation. Remarkably, the group of mRNAs that depend on Xrn1 for translation highly depend on Xrn1 for transcription and decay. Moreover, these three functions of Xrn1 are linked. Our results show a coordinated control of the three main stages of gene expression by Xrn1 to maintain proper homeostasis of membrane proteins. This coordination may be important to prevent their toxic aggregation.

**Results**

**Xrn1 drives translation of Brome mosaic virus RNA2.** Given the multifunctional nature of Xrn1 and its association with translating mRNAs during co-translational decay, we examined whether Xrn1 in addition regulates translation. As a first approach, we used the Brome mosaic virus (BMV)/yeast system. The ability of the plant BMV RNA2 genome to translate in yeast is a useful tool to identify and characterize specialized translational control mechanisms of host mRNAs\textsuperscript{13,14}. The 5′capped BMV RNA2 contains a tRNA-like structure instead of a poly(A) tail at the 3′end. Its long and structured 5′UTR and coding sequence (CDS) contain cis-sequences involved in translational control\textsuperscript{13,15}, When expressed in yeast, RNA2 is recognized by the ribosomes to translate the viral 2a polymerase. The complete BMV lifecycle occurs within the cytoplasm. However, as we were interested in identifying translational control mechanisms affecting cellular mRNAs, we expressed the RNA2 from a cellular promoter. We transformed wild-type (WT) yeast and an isogenic XRN1 deletion strain (xrn1Δ) with a plasmid expressing BMV RNA2 by the GAL1 promoter, whose transcription is activated by Xrn1. Interestingly, whereas the steady-state level of the viral RNA2 was increased in xrn1Δ cells, expression of the 2a protein was substantially decreased (Fig. 1a). Translatability of RNA2 (change in 2a protein level divided by change in RNA2 level) in xrn1Δ was only 0.4% of that in the WT. In xrn1Δ cells the majority of mRNA molecules are capped\textsuperscript{16} and a major fraction of uncapped mRNAs is associated to polyribosomes. Similarly, we found that most of RNA2 molecules (77%) are capped (Supplementary Fig. 1). In our calculations, we used total RNA2 instead of capped RNA2 as the difference in the obtained values is minimal. Translatability of RNA2 in xrn1Δ cells when considering only capped RNA2 is 0.5% instead of 0.4%. To determine quantitatively whether the Xrn1 effect on 2a protein levels involves protein degradation, we fused 2a to Renilla luciferase (2a-Rluc). Turnover of 2a-Rluc was determined by blocking translation with cycloheximide and measuring luciferase activity thereafter. Whereas translatability of the chimeric mRNA was affected by deleting XRN1 (Supplementary Fig. 2a), the 2a-Rluc protein turnover was not (Supplementary Fig. 2b). We conclude that Xrn1 promotes translation of RNA2.

To assess which regions of BMV RNA2 confer Xrn1-dependence for translation, we replaced different RNA2 sequences and quantified 2a and RNA2 levels in the presence or absence of Xrn1. The 5′UTR was replaced by the GAL1 5′UTR and the non-polyadenylated 3′UTR of the ADH1 transcript\textsuperscript{14} (Fig. 1b). These changes affected both protein and mRNA levels. Replacement of RNA2 3′UTR with ADH1 3′UTR had no effect on translatability (Fig. 1b, lane 6). However, the 5′UTR played an important role on the capacity of Xrn1 to affect translatability (Fig. 1b, lane 8). Replacement of RNA2 CDS with GFP CDS had a modest effect on translatability (Fig. 1b, lane 10). Taken together, results in Fig. 1b indicate that the 5′ UTR is the most Xrn1-responsive region in RNA2.

To investigate which step of BMV RNA2 translation is stimulated by Xrn1, we performed polysome-profiling analyses in WT and xrn1Δ cells expressing RNA2. Consistently with previous studies\textsuperscript{17}, the global tRNA profile, indicative of the global translation, was only mildly affected in xrn1Δ (Fig. 2a). Northern blot analyses along the polysome profile showed that deletion of XRN1 shifted RNA2 toward monosomal, 60s and 40s fractions (Fig. 2b), suggesting a role of Xrn1 in translation initiation. In agreement with the known dependence of polysomes on the presence of Mg\textsuperscript{2+}, EDTA treatment shifted BMV RNA2 from heavy polysomes to lighter fractions (Supplementary Fig. 3). To further strengthen the link of Xrn1 to translational control, we examined whether Xrn1 co-sediments with ribosomes in polysome profiling. We observed an enrichment of Xrn1 in fractions corresponding to 40s subunits (Fig. 2c). Collectively, these results indicate that Xrn1 is required for efficient translation of RNA2, likely at early events of the translation initiation step.

The effect of Xrn1 on BMV RNA2 translation is specific. Stably deleting XRN1 can lead to selection of adaptive secondary mutations that might cause indirect effects. To overcome this potential limitation, we fused Xrn1 to an auxin-inducible degron (AID) that induces rapid degradation of Xrn1 and measured the immediate effects\textsuperscript{18}. AID-tagging of Xrn1 did not significantly affect its function in BMV RNA2 translation (Supplementary Fig. 4). WT cells carrying AID-tagged XRN1 in its natural genomic locus and a plasmid expressing RNA2-Rluc from a GAL1 promoter were grown in raffinose to logarithmic phase. Addition of galactose and auxin resulted in simultaneous induction of BMV RNA2-Rluc
transcription and Xrn1-AID depletion (Fig. 3a). Xrn1-AID protein levels decreased upon addition of auxin and were no longer detected after 35 min (Fig. 3b) while global translation was not affected at this time-point (Fig. 3c). Induction kinetics of RNA2-Rluc was comparable in WT and xrn1Δ (Fig. 3d), probably due to the compensatory effect of transcription and decay of this transcript. In contrast, the level of 2a-Rluc protein, which reflects mainly translation at these early time-points after galactose induction, was reduced in auxin-treated cells (Fig. 3e). The effect of Xrn1-AID depletion was observed already at early time-points after auxin addition, indicating that depletion of Xrn1 inhibits RNA2 translation immediately and that an indirect effect is unlikely.

### Table 1

| Protein          | WT     | xrn1Δ |
|------------------|--------|-------|
| 2a (kDa)         | 115.5  | 48.8  |
| m7G              |        |       |
| 2a Exp. (%)      | 100    | 1.8   |
| SEM              | 28     | 0.8   |
| 18S              |        |       |
| RNA2 acc. (%)    | 100    | 464   |
| SEM              | 14     | 35    |
| Translatability (%) | 100    | 0.4   |
| SEM              | 14     | 0.2   |

### Figure 1

**a** Xrn1 depletion inhibits BMV RNA2 translation. Simultaneously exposed western blot and northern blot panels showing steady-state levels of viral protein 2a and RNA2. **b** The BMV RNA2 5’UTR and CDS confer dependence on Xrn1 for translation. Black solid lines represent viral UTRs and orange lines GAL1 mRNA 5’- and ADH1 3’UTRs. The white and green boxes represent 2a and GFP CDSs, respectively. Throughout this study, BMV RNA2 was expressed from a plasmid by the GAL1 promoter, PGK protein, and 18 S RNA were used as loading controls for western and northern blots, respectively. Values denote expression relative to WT, taken arbitrarily as 100% and are calculated from n = 3 independent colonies and expressed as mean ± SEM. Dotted lines represent a separation of the shown samples in the same membrane. Open circles indicate the individual data points. Source data are provided as a Source Data file.
Next, by replacing Xrn1 with its nuclear paralog Rat1 we explored whether the positive role of Xrn1 in translation is specific. When forced to localize in the cytoplasm by deleting its nuclear localization signal (NLS), Rat1ΔNLS functionally replaces Xrn1ΔNLS. Accordingly, expression of Rat1ΔNLS in xrn1Δ cells fully rescued viral RNA2 degradation and cellular growth (Fig. 4a). In contrast, Rat1ΔNLS did not efficiently rescue translation of BMV RNA2, since, upon addition of Rat1ΔNLS to xrn1Δ cells, the expression of protein 2a was only marginally recovered (Fig. 4b). Hence, viral RNA2 translation requires an Xrn1-specific function, not simply its 5′ to 3′ exonuclease activity per se.

As our polysome-profiling results suggested a role of Xrn1 in translation initiation (Fig. 2b), we examined whether Xrn1 interacts with the eIF4F complex, a key component of the translation initiation machinery that binds to capped mRNA and mediates its interaction with the 43S pre-initiation complex. This complex consists of the RNA helicase eIF4A, the cap-binding proteins eIF4E and the large eIF4G scaffold protein. We carried out immunoprecipitation assays using yeast strains carrying functionally validated genomically tagged fusions of eIF4G, eIF4A, or eIF4E transformed with plasmids expressing FLAG-tag fusions of Xrn1 or Rat1ΔNLS (Fig. 4c and Supplementary Figs. 5 and 6). Remarkably, Xrn1, but not Rat1ΔNLS, co-immunoprecipitated with eIF4G in an RNase-resistant manner. Neither Xrn1 nor Rat1ΔNLS interacted with eIF4A while RNase-sensitive interactions with eIF4E were detected for both Xrn1 and Rat1ΔNLS. Thus, Xrn1, but not Rat1ΔNLS, interacts with eIF4G. To test whether the ability of Xrn1 to interact with eIF4G is functionally linked to its role in translation we designed a gain-of-function experiment. The sequence and folding of the N-terminal exonuclease domains of Rat1 and Xrn1 are very similar. However, Rat1 lacks an unstructured C-terminal domain present in Xrn1 that serves as an interaction platform in higher eukaryotes. We used structural modeling to generate a chimera between the Rat1ΔNLS N-terminal domain and the Xrn1 C-terminal tail (Rat1ΔNLS-XC, Supplementary Fig. 7a, b) and tested its expression (Supplementary Fig. 6) and functionality (Supplementary Fig. 7c). As found for Rat1ΔNLS, the chimeric protein rescued cellular growth in xrn1Δ cells indicating that the fusion of the Xrn1 C-terminal tail does not compromise the global function of Rat1ΔNLS. Expression of Rat1ΔNLS-XC resulted in a twofold increase of viral 2a expression when compared to Rat1ΔNLS (Fig. 4d) while the steady-state levels of RNA2 were similar. Notably, the Rat1ΔNLS-XC chimera concomitantly gained interaction with the translation initiation factor eIF4G (Fig. 4e). Consistently, in comparison to wild-type Xrn1, expression of Rat1ΔNLS shifted RNA2 from light polysomes to 40s and 60s fractions in polysome-profiling experiments. This shift was partially abrogated when Rat1ΔNLS-XC was expressed (Supplementary Fig. 8a). Overall our data suggest a specific function of Xrn1 in translation mechanistically linked to its C-terminal domain and the ability to interact with eIF4G.

**Xrn1 drives translation and localization of secretome mRNAs.** Our results using the BMV RNA2 model in yeast prompted us to investigate whether Xrn1 also regulates translation of cellular mRNAs. We used the Xrn1-AID degron system to avoid adaptive effects and studied genome-wide translational changes using polysome-profiling. This method is based on the isolation and deep-sequencing of ribosome-protected fragments (RPFs) and

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**Fig. 2** Xrn1 depletion shifts RNA2 toward single ribosomal subunits fractions. a ultraviolet (UV) absorbance rRNA profile at 260 nm of an extract from WT and xrn1Δ cells expressing RNA2 after sedimentation on a 10 to 50% sucrose gradient. b Depletion of Xrn1 shifts BMV RNA2 toward monosomal fractions. Upper panel: distribution of normalized BMV RNA2 accumulation across the polysome profile. Fractions were grouped into free (1-5), single ribosome subunits (6-11), monosomes (12-15), light polysomes (16-21), and heavy polysomes (22-26). RNA was quantified by northern blot. Results represent averages of n = 3 biological replicates. Error bars represent SEM. Open circles indicate the individual data points. Lower panel: representative northern blots. c Xrn1 cofractionates with free 40s subunits in polysome-profiling analysis. Top panel: UV absorbance rRNA profile at 260 nm of an extract from WT cells. Lower panel: Fractions were TCA-precipitated and analyzed by western blot. Specific antibodies detecting Xrn1p, S8 protein (small ribosomal subunit) and L1 protein (large ribosomal subunit) were used. S lane corresponds to soluble proteins not associated to ribosome subunits and P lane to a pool of three fractions corresponding to polysomes. Source data are provided as a Source Data file.
parallel transcriptome analysis. Ribosome-profiling was performed on samples before and after 30 min of auxin treatment (Fig. 5a). Replicates of the RPF and RNAseq libraries clustered in principal-component analyses (Supplementary Fig. 9). To identify genes showing changes of translational efficiency upon degron-mediated Xrn1 knock-down (Xrn1-KD), we used the Riborex R-package, which assesses whether changes of ribosome occupancy could be explained by changes of the corresponding mRNA. Genes were plotted according to their log2-fold changes in mRNA abundance and ribosome occupancy (RPF) (Fig. 5b). A majority of genes showed no significant changes in translational efficiency. Strikingly, we identified a specific set of genes translationally activated (445) or repressed (597) by Xrn1. Genes showing significant alterations of translational efficiency (FDR < 0.05) were grouped according to their relative mRNA and RPF efficiency (Fig. 5c and Supplementary Data 2). These results revealed that Xrn1-dependent regulation characterizes at least three different groups of genes, which are defined by different behaviors in RNA steady-state levels and translation, and are enriched for different cellular functions.

The absence of Xrn1 can lead to accumulation of uncapped mRNAs that are normally degraded by Xrn1 (see Supplementary Fig. 1), which might affect our results. However, as we observe for RNA2, most of the mRNAs are capped in xrn1Δ cells. The minor fraction of deadenylated and uncapped mRNAs is mainly associated to polyribosomes and thus are captured by the ribosome-profiling libraries but are depleted from the oligo(dT)-selected RNAseq libraries. Thus, the activating effects of Xrn1 on the translationally activated mRNAs would be slightly higher than our estimates, while the repressing effects on the translationally repressed mRNAs would be slightly lower.

The cytosol and the ER represent distinct biological environments for translation with different regulatory factors. As membrane proteins are translated and glycosylated at the ER, we questioned whether Xrn1-dependence for translation is linked to a possible role of Xrn1 in localizing the affected mRNAs to the ER. Based on our Ribosome-profiling data, we selected three groups of transcripts: (i) transcripts encoding membrane proteins that depend on Xrn1 for translation and, as controls, (ii) transcripts encoding membrane proteins that do not depend on Xrn1 for translation, and (iii) transcripts encoding cytosolic proteins. Next, we isolated ER membranes from the cytosol and quantified the relative amounts of the selected transcripts associated to ER and cytosol in WT as compared with that in Xrn1-KD cells (Fig. 5d). Transcripts that depend on Xrn1 for

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**Fig. 3** Xrn1 depletion leads to immediate defects on viral RNA2 translation. **a** Scheme of the experimental set-up to simultaneously deplete Xrn1-AID and express viral 2a-Rluc. **b** Expression of Xrn1-AID upon addition of auxin to the media. c UV absorbance profile at 260 nm of an extract from the Xrn1-AID strain before and after 35 min of auxin addition. **d** Relative RNA2-Rluc RNA accumulation, obtained by quantifying RNA level by RT-qPCR (arbitrary units (a.u)) and **e** Relative 2a-Rluc protein expression (R.U.) before and after galactose and auxin addition. Results represent averages obtained from n = 3 biological replicates. Error bars represent SEM. Source data are provided as a Source Data file.
Fig. 4 Xrn1-dependence for translation is specific and mediated by the interaction with eIF4G. **a** Rat1ΔNLS fully replaces Xrn1 in mRNA decay and cell proliferation. xrn1Δ cells expressing BMV RNA2 under GAL1 promoter and either WT Xrn1, Rat1ΔNLS or an empty plasmid were grown in galactose. Left, transcription of RNA2 was shut-off upon glucose addition and BMV RNA2 stability was determined by monitoring RNA2 levels by northern blot analysis at various time-points post-glucose addition. A representative example out of three replicates is shown. Right, growth curves in galactose media. Results represent averages obtained from three replicates. **b** Rat1ΔNLS does not replace Xrn1 in RNA2 translation. Western blot (upper panel) and northern blot (lower panel) analysis showing steady-state levels of viral protein 2a and viral RNA2 in xrn1Δ cells expressing Xrn1 or Rat1ΔNLS. Asterisk points at a non-specific band. Quantifications are relative to xrn1Δ transformed with WT Xrn1 plasmid. Results represent averages of n = 3 biological replicates. **c** Xrn1 interacts with eIF4G. Western blot analysis of immunoprecipitation assays. Xrn1-FLAG and Rat1ΔNLS-FLAG proteins were expressed in yeast strains expressing either eIF4G-GFP, eIF4A-GFP, or eIF4E-GFP fusion proteins. As a control, the functionality of GFP-fused strains was assessed (Supplementary Fig. 5). Immunoprecipitations were carried out with GFP-trap beads with extracts treated (+) or not treated (−) with RNase A. Expression levels of eIF4G, eIF4A, and eIF4E were detected by anti-GFP antibody. **d** Expression of Rat1ΔNLS-XC rescues BMV RNA2 translation. Western blot (upper panel) and northern blot (lower panel) analysis. Results represent averages of three replicates. Expression levels of flag-tagged Xrn1, Rat1ΔNLS, and Rat1ΔNLS-XC were analyzed by western blot (Supplementary Fig. 6). **e** Interaction with eIF4G studied by co-immunoprecipitation analyses. Source data are provided as a Source Data file.
translational activation exhibited a fourfold increase in ER localization when Xrn1 was present while transcripts that do not depend on Xrn1 for translation exhibited a twofold increase. No such effect was observed for cytosolic transcripts. Membrane proteins (as defined in ref. 24) were enriched among the Xrn1-dependent transcripts of our ribosome-proliling data as Xrn1-activated transcripts were 2.09 times more likely to be part of the membrane compared to transcripts that do not depend on Xrn1 for translation. In contrast, transcripts that were translationally repressed upon Xrn1-KD were devoid of membrane genes (Supplementary Data 3). Considering that our current subset of Xrn1-activated genes represents a stringent selection of
Xrn1-dependent events, our analysis might miss transcripts affected by Xrn1 depletion (false negatives). Based on the observed behavior of membrane genes among transcripts that are translationally controlled by Xrn1, many of these false negatives can be anticipated to be membrane genes as well. This might explain the twofold increase in ER localization observed in mRNAs encoding membrane proteins classified as not dependent on Xrn1.

We conclude that Xrn1-KD cells are defective in recruiting mRNAs to the ER while the extent of this defect correlates with the capacity of Xrn1 to activate translation. As this defect was observed shortly after Xrn1 depletion, it is unlikely an indirect effect of the absence of Xrn1. The described routes of targeting mRNAs to the ER for translation include signal recognition particle (SRP)-dependent and -independent pathways. Based on previous studies, we found that both SRP-dependent and -independent transcripts are similarly represented in the transcripts encoding membrane proteins whose translation is activated by Xrn1. This argues in favor of Xrn1 functioning along both routes.

Xrn1-activated mRNAs have long and structured 5’UTRs. Next, we investigated whether mRNAs regulated by Xrn1 share common physical properties. First, we calculated the average length of the 5’UTRs, CDSs and 3’UTRs for the three groups and compared them to all genes that were not significantly altered (Fig. 6a and Supplementary Data 4a–d). Xrn1-activated mRNAs had significantly longer 5’UTRs (80 nt) and CDS (1555.5 nt) when compared to transcripts not significantly altered (52 and 1113 nt). In contrast, Xrn1-repressed mRNAs had shorter 5’UTRs (44 nt) and CDS (35.5 nt) and CDS (mRNA buffered: 753; mRNA decreased: 600 nt) compared to not significantly altered transcripts, whereas only repressed genes with buffered mRNA levels showed an increased 3’UTR length (128 nt) compared to background genes (105 nt). Interestingly, as for the cellular Xrn1-activated mRNAs, BMV RNA2 contains long 5’UTRs and CDS (92 nt and 2468 nt). Second, as BMV RNA2 5’UTR contains highly structured sequences, we examined whether this feature also characterizes the cellular Xrn1-activated mRNAs and extended this analysis to the CDS. We used previously published datasets of genome-wide RNA secondary structure obtained by PARS (Parallel Analysis of RNA structure) to analyze the RNA structure profile in the 5’UTRs, CDSs and 3’UTRs. PARS scores are based on deep-sequencing of RNA fragments obtained by RNA digestion with enzymes that exhibit structural preferences. While there were no substantial differences in the 3’UTR, all groups of transcripts regulated by Xrn1 had a significantly higher PARS score in the CDS (0.32–0.35) compared to transcripts not significantly affected by Xrn1 (0.25) (Fig. 6b). Importantly, only activated transcripts had a higher structured 5’UTR (0.16) compared to transcripts not significantly affected (0.06). Repressed genes, whose mRNA levels decreased due to depletion of Xrn1, exhibited the lowest structure of the 5’-UTRs (0.03). These differences are more visible when plotting the PARS score distribution (Fig. 6c). Note, in particular, that the PARS score drops around the translation initiation sites (TIS) (Fig. 6c). This relatively unstructured region has probably evolved to permit an easy access to the ribosome at the canonical translation initiation site, and may contribute to the recognition of true start sites. Interestingly, Xrn1-activated mRNAs have a substantially higher PARS score at the TIS and their upstream regions, suggesting that one of the Xrn1 functions is to override the structural barrier at the TIS. Notably, we found that long 5’UTRs and CDSs and highly structured 5’UTRs with unfavorable contexts for translation initiation are indeed common features for mRNAs encoding membrane proteins (as defined in ref. 24). Remarkably, these mRNAs identified to depend on Xrn1 for translation show the highest PARS scores (Supplementary Fig. 10).

To further understand the role of Xrn1 in translational regulation, we questioned whether the changes observed in ribosome occupancy in activated transcripts were due to changes in initiation and/or elongation. Ribosome-profiling can detect defects in specific translation steps because discrete ribosome pausing increases the likelihood of capturing footprints in the pausing site by deep-sequencing. Therefore, defects in elongation caused by ribosome stalling result in a peak in ribosome density and an accumulation of ribosomes upstream of the stalling site. This would be visible as a change in slope in a metagene analysis, with the 5’ end showing an increased footprint density and the 3’ end showing a decrease. In contrast, differences in translation initiation result in a shift of ribosome occupancy along the entire CDS. To distinguish between these two possibilities, we examined the relative RPF distribution along the CDS of the three mRNA groups defined previously and compared them to genes that did not show any significant alterations (Fig. 6d). In agreement with a role of Xrn1 in translation initiation, activated genes exhibited a general reduction of footprint density while repressed genes exhibited a general increase. Collectively, we conclude that long 5’UTRs and CDSs and highly structured 5’UTRs are common features of both BMV RNA2 and cellular transcripts that depend on Xrn1 for translation. This suggests a common role for Xrn1 in regulating translation of cellular mRNAs and BMV RNA2, likely at the translation initiation step.
The distinct functions of Xrn1 in gene expression are linked. To examine whether the function of Xrn1 in translation is linked to its known roles in transcription and decay, we determined the effect of Xrn1 on the transcription and decay of those mRNAs that are translationally activated by Xrn1. To obtain a whole-genome view of transcription rates (TRs), we performed genomic run-on experiments (GRO) in the Xrn1-AID degron system 30 min after auxin addition. In parallel with GRO analysis, we determined the mRNA steady-state levels, which allowed us to calculate mRNA half-lives. Reassuringly, we found that depletion of Xrn1 for 30 min had a similar effect on transcription rates to that previously observed in xrn1Δ strain. The GO categories enriched in genes transcriptionally activated or repressed by Xrn1 were similar to those observed when Xrn1 is permanently depleted although quantitatively

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**Fig. 6** Features of mRNAs translationally controlled by Xrn1. **a, b** Box-plot depicting the mean length (a) and the mean PARS score (b) of the 5'UTRs, CDSs and 3'UTRs for the three subgroups studied and the control group (gray). For boxplots, box boundaries represent the 1st and 3rd quartile of the distribution, while the center line represents the 2nd quartile (median). Whiskers indicate either the most extreme values or extend to 1.5 times the interquartile range starting from the respective box boundary. Black dots indicate outliers (default R parameters). Statistical significance was calculated using a Wilcoxon-test (p-values available in Supplementary Data 4). **c** Metagene analysis of the PARS scores. Vertical dashed line corresponds to the translation initiation site (TIS). The x-axis represents the nucleotide position relative to the TIS. **d** Metagene analysis of the average mRNA-normalized RPF coverage along the CDS for activated (red, mRNA = ), repressed (green, mRNA = and orange, mRNA ↓), and not affected mRNAs (gray). A light red line corresponds to untreated cells, whereas turquoise corresponds to treatment with auxin for 30 min.

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The distinct functions of Xrn1 in gene expression are linked. To examine whether the function of Xrn1 in translation is linked to its known roles in transcription and decay, we determined the effect of Xrn1 on the transcription and decay of those mRNAs that are translationally activated by Xrn1. To obtain a whole-genome view of transcription rates (TRs), we performed genomic run-on experiments (GRO) in the Xrn1-AID degron system 30 min after auxin addition. In parallel with GRO analysis, we determined the mRNA steady-state levels, which allowed us to calculate mRNA half-lives. Reassuringly, we found that depletion of Xrn1 for 30 min had a similar effect on transcription rates to that previously observed in xrn1Δ strain. The GO categories enriched in genes transcriptionally activated (ribosome biogenesis, translation) or repressed (mitochondria, respiration) by Xrn1 were similar to those observed when Xrn1 is permanently depleted although quantitatively
lower, suggesting that permanent depletion of Xrn1 intensifies this phenotype. Likewise, the distinction between the Xrn1 Synthegradon group, which is highly responsive to XRN1 disruption, and the anti-Synthegradon group, which contains the least responsive genes, was clearly visible upon Xrn1-AID depletion (Supplementary Fig. 11). Importantly, within the groups of mRNAs translationally regulated by Xrn1, the translationally activated one showed significantly decreased transcription rates and increased half-lives upon Xrn1 depletion (Fig. 7a, b = mRNA, red). Thus, Xrn1 stimulates mRNA synthesis, translation, and decay of this group of mRNAs.

Next, we investigated whether these three Xrn1-driven functions are linked. If this is the case, defects in one function should affect the others. To address this key issue, we made use of the catalytically inactive xrn1D208A mutant. This mutant binds decapped mRNAs but is unable to degrade them and becomes

![Graph of transcription rate and half-life](image)

**Fig. 7** Xrn1 functions in mRNA transcription, decay, and translation are linked. **a** Transcription rates and **b** half-lives of the studied mRNA groups. Statistical significance was calculated using a Wilcoxon-test (*represents p-value < 0.001) The results represent an average of n = 3 biological replicates. For boxplots, box boundaries represent the 1st and 3rd quartile of the distribution, while the center line represents the 2nd quartile (median). Whiskers indicate either the most extreme values or extend to 1.5 times the interquartile range starting from the respective box boundary. **c** Xrn1 D208A effect on BMV RNA2 translation. **d** When introduced directly into the cytosol by electroporation, RNA2 does not depend on Xrn1 for translation. RNA2-Rluc mRNA and a control mRNA expressing Fluc were directly electroporated into the cytosol. 2a-Rluc protein values were normalized by those of Fluc and represented in comparison to the WT. Quality controls for RNA integrity and capping were performed (Supplementary Fig. 12). The results represent an average of n = 3 biological replicates ± SEM. Open circles indicate the individual dot plots. Source data are provided as a Source Data file.
trapped in an RNA-bound state. Consequently, the nuclear import of Xrn1, and other mRNA decay factors whose import depend on Xrn1, is impaired. Replacing XRN1 at its genomic locus by XRN1D208A results in transcription and decay defects comparable to those in xrn1Δ cells. We reasoned that if the function of Xrn1 in translation, transcription and decay are linked, xrn1D208A mutation should also impair Xrn1-dependent translation. Ribosome-profiling analyses of the xrn1D208A mutant identified a group of cellular mRNAs that depend on Xrn1 for translation. These mRNAs were enriched for cellular functions related to protein glycosylation and membrane localization (Supplementary Data 6) and significantly overlapped with those identified with the Xrn1-AID system (p-value = 1.4e-29, Hypergeometric test). Moreover, xrn1D208A mutation inhibited BMV RNA2 translation like XRN1 deletion (Fig. 7c). Similarly, a catalytically inactive version of Rat1ΔNLS-XC (Rat1ΔNLS-XC-D235A) inhibited RNA2 translation (Supplementary Fig. 8a, b). To obtain further insights into the linkage between the three functions of Xrn1, we uncoupled RNA2 translation from transcription. We reasoned that if Xrn1-dependent translation requires a previous function of Xrn1 in transcription, introducing RNA2 directly into the cytosol would result in an Xrn1-independent translation of RNA2. To test this, we electroporated in vitro transcribed RNA2-Rluc into WT and xrn1Δ cells together with an mRNA control expressing Firefly-luc for normalization (Fig. 7d). Remarkably, in contrast to the inhibition of 2a protein expression observed in xrn1Δ when RNA2 was expressed from a plasmid (Fig. 1a), deletion of XRN1 did not inhibit 2a protein expression (Fig. 7d). Instead, we observed an increase in the expression of 2a protein consistent with an extended RNA2 half-life (Fig. 4a), indicating that the linkage between transcription and post-transcriptional stages was broken by circumventing transcription and introducing mRNAs by electroporation. Altogether, we conclude that the functions of Xrn1 in synthesis, translation, and decay of the translationally activated mRNAs are linked.

**Discussion**

In this study, we reveal an unanticipated role for the major 5′–3′ exonuclease Xrn1 in both activating translation of mRNAs encoding membrane proteins and in directing these mRNAs to the ER, their translation site. Remarkably, for this group of mRNAs Xrn1 activates translation, transcription and decay, and these functions are linked. Our results uncover a crosstalk between the three major processes of gene expression coordinated by Xrn1 to express membrane proteins.

A number of observations are consistent with a specific role of Xrn1 in translation initiation. First, transient depletion of Xrn1 has an immediate inhibitory effect on RNA2 translation. Consequently, a selection of adaptive mutations that may account for the observed translation effects seems unlikely (Fig. 3d, e). Second, Rat1ΔNLS, the nuclear paralog of Xrn1 retained in the cytoplasm due to the deletion of its NLS, fully rescues RNA2 stability (Fig. 4a), but not RNA2 translation (Fig. 4b). These data argue against the possibility that changes in mRNA abundance driven by Xrn1 depletion are responsible for the observed translation defects. It has been reported that Xrn1 depletion causes changes in the formation of RNA duplexes that result from transcription of convergent genes and affect their protein expression levels. Our ribosome-profiling data show, however, no enrichment for convergent genes. Third, polysome-profiling analyses demonstrate that, upon Xrn1 depletion, BMV RNA2 shifts from polysomes to monosomes (Fig. 2b) and ribosome-profiling analyses show a decrease in the footprint density along the CDSs of cellular mRNAs that are translationally activated by Xrn1 (Fig. 6d). These features are consistent with effects in translation initiation. The co-fractionation of Xrn1 with 40s ribosomal subunits would point to a role of Xrn1 in early translation initiation events (Fig. 2c). Fourth, Xrn1 but not Rat1ΔNLS interacts genetically, physically, and functionally (this study) with the translation initiation factor elf4G. This was shown by immunoprecipitation analyses (Fig. 4c, d) and gain-of-function studies in which the C-terminal domain of Xrn1, the only domain not present in Rat1, was fused to Rat1ΔNLS (Fig. 4d). This domain is highly disordered and includes short linear motifs (SLiMs), features typical of protein sequences with capacity to bind multiple interacting partners. Such motifs also characterize the Xrn1 homologs in Homo sapiens and Drosophila melanogaster. Last, the mRNAs whose translation is activated by Xrn1 have an unfavorable translation initiation context. They contain long and highly structured 5′UTRs as demonstrated by PARS scoring for host mRNAs and by functional studies for RNA2. Typically, PARS scores drop immediately upstream of the translation initiation site to favour translation initiation. In contrast, Xrn1 activates translation of mRNAs that are translationally activated by Xrn1 and exhibit an unusual high PARS score in this region, suggesting that one function of Xrn1 is to overcome this impediment. Reassuringly, replacement of the highly structured 5′UTR in RNA2 resulted in a 35-fold increase of translatability in WT cells and a 13-fold decrease in Xrn1-dependence (Fig. 1b).

Xrn1 coordinates a linkage between transcription, translation, and decay for a specific group of mRNAs enriched in transcripts encoding membrane proteins. We surmise that this linkage has evolved to control proper gene expression of membrane proteins. These proteins contain hydrophobic domains with strong tendencies to aggregate. Consequently, their expression levels and localization must be finely tuned to avoid aggregations that might be toxic. Indeed, mRNAs related to vacuole transport in yeast and the endomembrane system in Arabidopsis thaliana are among the most frequently co-translationally degraded by Xrn1. This feedback mechanism would ensure that Xrn1-dependent mRNAs are efficiently translated only when decay is working properly. The decision to express a gene, or a family of genes, is obtained by crosstalks between all major stages of the mRNA lifecycle. For example, it is counterproductive to transcribe a gene if the translation apparatus is not capable of translating it. One mechanism to permit these crosstalks is by factors that function in all these stages. Xrn1 seems to carry this function. Another example is Rpba7, that also acts in transcription, mRNA export, translation, and decay. Interestingly, XRN1 and RPBA7 interact genetically, raising the possibility that they function in a coordinated manner. One advantage for the cell of using factors that function in all stages of the mRNA lifecycle is the ability to regulate the synthesis and functionality of mRNAs by regulating these factors.

In our study, we also identified two subsets of mRNAs translationally repressed by Xrn1. One of them is enriched for functions related to proteasomal degradation and protein folding (Fig. 5c). This hints to a crosstalk of mRNA biogenesis and turnover with protein degradation. The other subset of mRNAs is enriched for functions related to translation and ribosomes. Interestingly, these mRNAs are the most dependent on Xrn1 for transcription and decay at optimal growth conditions. This would allow a precise control of the global translation, the most energy consuming cellular process. Although the mechanistic details of Xrn1 function in translation repression are unclear, our data indicate that it is context dependent (Fig. 6a–c) and, in the case of repressed mRNAs, not linked to transcription and decay activation (Fig. 7a, b).

Based on our observations, we propose a model wherein Xrn1 promotes translation initiation of mRNAs with unfavorable
transcription initiation contexts by interacting with eIF4G and stabilizing the interaction of the scanning 40s subunit with the mRNA. As eIF4G shuttles to the nucleus, where it is proposed to function in splicing, an interesting possibility is that the two shuttling proteins interact in the nucleus or even shuttle together. By promoting translation, Xrn1 will then favour co-translational targeting of mRNAs to the ER. Alternatively, Xrn1 may assist mRNA localization at the ER by a translation-independent mechanism. Once at the ER and after several rounds of translation, the mRNAs would then be co-translationally degraded by Xrn1.

We propose that the Xrn1-mediated crosstalk between transcription, translation, and decay involves the decaysome complex. The decapping activators Lsm1–7, Pat1, and Dhh1 shuttle to the nucleus in an Xrn1-dependent manner, associate to chromatin and stimulate transcription1. Likewise, these decapping activators are required for the function of Xrn1 in translation. Lsm1–7, Pat1, and Dhh1 are required for BMV RNA2 translation and the DEAD-box RNA helicase Dhh1 activates translation of cellular RNAs13 that significantly overlap with those translationally activated by Xrn1 (p-value = 1 × 10−5, hypergeometric test). However, our data support that each factor carries a distinct non-overlapping function that complements each other. For example, Lsm1–7/Pat1-requirements for BMV RNA translation are related to sequences within the 3′ UTR10,41. Dhh1-requirement to sequences within the 5′ and 3′ UTRs and CDS13 and Xrn1-requirement primarily to sequences within the 5′ UTR (this study).

Collectively, our data provide mechanistic details of the function of Xrn1 in translation activation and uncover an Xrn1-mediated crosstalk between the three major stages in gene expression. A key area for future research will be in elucidating how the transitions between the subsequent Xrn1 functions are regulated and responsive to the environment.

**Methods**

**Yeast cultures.** Yeast cells were grown in synthetic complete medium (SC) at 30°C. Galactose (2%) was used as carbon source and it also served as inductor for GAL1-directed viral RNA expression. After transformation, three colonies for each condition were selected and streaked on a selective media plate. Cells were grown over-day in selective liquid media and diluted to grow overnight. Next day, they were diluted and grown until the doubling time between triplicates was similar and an OD600 of ~0.6 was reached. Yeast strains and plasmids are listed in Supplementary Data 7.

**Cloning of Xrn1 derivative strains.** Xrn1-AID strain was generated by amplification of the cassette present in BYP7427 plasmid with Xrn1-specific primers. The PCR product was purified and transformed in BY4741 and plated in minimal media with corresponding selection markers (+His, +Ade, +Leu, +Tryp). Positive colonies were selected by checking the correct integration of the AID cassette in Xrn1 and were subsequently sequenced. Xrn1 D208A strain was generated by the following strategy. First, XRN1 gene was amplified and cloned in pS229 using Fw-Xrn1–250-Agell and Rev-Xrn1+110-AcI primers. Second, this plasmid was transformed in BY4741 Xrn1Δ strain and plated in −G418 +His selective plates for homologous recombination to occur. Third, colonies were tested by PCR to ensure recombination had occurred in the right locus and subsequently sequenced. All primers and strains are listed in Supplementary Data 7 and Supplementary Data 8.

**Cloning of Xrn1 derivative plasmids.** The plasmids generated for this study were cloned following conventional molecular cloning strategies. In order to be able to select for all plasmids used in this study, we changed the selection markers of plasmids pAJ37 and pAJ2281 as URA3 marker, and we named the plasmids pBBM1 and pBBM3, respectively. Xrn1 D208A mutant plasmid (pBBM2) was generated using the KOD Hot Start DNA polymerase from Millipore. pLCM2 was generated by inserting one single copy of a FLAG-tag (GATTACAAGGATGACGAGTAAAG) in the 3′ end of pBBM2. To increase the signal, we inserted two more copies in pLCM2, generating a 3FLAG (GATTACCAAGGATGACGAGTAAAGGAGGATGACGAGTAAAGGAGGATGACGAGTAAAG) bait plasmid (pLCM7). All the experiments done along this study have been performed with pLCM6 and we refer to it as Rat1ΔNLS-FLAG plasmid. pLCM7 was generated by taking out the 3FLAG from pLCM6 and introducing it in the 3′ end of pBBM1 plasmid.

**BMV RNA translation assay.** To evaluate BMV protein translation, yeast cells were transformed with the corresponding plasmids and grown as specified in the previous section. Two OD units (Optical Density units) were harvested for protein extraction and total RNA extraction. BMV mRNA was generated by fusing the N-terminal domain (1–884 aminoacids) of Xrn1 in pBBM3 to the C-terminal domain (733–1528 aminoacids) of Rat1ΔNLS in pLCM7. Also, we depleted a loop in the CDS of Rat1ΔNLS (Δ24–42 aminoacids) because it was hampering the interaction between the N-terminal and the C-terminal domain of the chimera.

**Protein turnover assay.** Yeast cells transformed with a plasmid encoding for RNA2A-Bluc reporter (pJL-16) were grown as described in Yeast Cultures section. When they reached an OD600 of 0.5, protein synthesis was stopped with 0.5 mg/ml cycloheximide. Renilla luciferase assay (Dual-Glo, Promega) was used following the protocol provided by the manufacturer. Samples were collected at different intervals during 3 h by directly transferring 10 µl of culture to 100 µl of 1x Passive Lysis Buffer. Only 10 µl of the lysate were used (the rest was stored at −80°C and 200 µl of LARII-StopGlo solution (1:1) was subsequently added. FB12 Luminometer was employed to read Luciferase activity, with 5 s of equilibration time and 5 s of measurement time. The values obtained were corrected by the corresponding OD600 and were represented relatively to the first time-point (t = 0). This protocol was adapted from ref. 46.

**RNA ligase-mediated (RLM)-RAPID amplification of cDNA ends (RACE).** To characterize the 5′ terminus of BMV RNA2A, total RNA obtained by phenol extraction was used to perform 5′ RLMPARCaMediated (RLM) rapid amplification of cDNA ends (RLM-RACE) assay using FirstChoice RLM- RACE Kit (Thermo Fisher Scientific) following manufacturer’s instructions. For the outer 5′ RLM-RACE PCR, 5′ RACE outer primer and gene-specific primer (5′-CATTGTG TGGACGGTTGCGCAAA-3′) were used. One thousand dilution of the above-mentioned PCR was used to amplify a nested PCR fragment with a 5′ RACE inner primer and a gene-specific primer (5′-CTCCTACTTCCAAGGCGGCCATGTA3′).

**Polysonome profiling.** Cultures were grown from OD600 = 0.02 to an OD600 = 0.5 in YPD media (Formedium) at 30°C. In order to stabilize elongating ribosomes, cells were treated with cycloheximide (CHX, 100 µg/ml final concentration) during 1 min with manual shaking at room temperature. Cells were quickly harvested with a vacuum filtration system, scraped out of the filter and immediately frozen in liquid nitrogen with 500 µl of lysis buffer (20 mM Tris-HCl (pH = 7.5), 100 mM NaCl, 5 mM MgCl2, 1% Triton X100, 100 µg/ml CHX). Cells were lysed with the Evans Lysis Buffer (SPEX SamplePrep) with two cycles of 2 min at 5000 rpm with a 2 min cooling-down step in between. Cell lysates were thawed at 30°C for 1 min and centrifuged at 3000 × g for 4 min and 3 min. The soluble fraction was transferred to new tubes and centrifuged at 10,000 × g for 4 min for 5 min. After quantification, aliquots of 12 µgRNA were made and stored at −80°C. Linear gradients of 10–50% sucrose were prepared in 50 mM Tris-HCl (pH = 7.5), 50 mM NaCl,12 mM MgCl2, 0.5 mM DTT, 100 µg/ml CHX. The Gradient Master (Biocomp) was used for making the gradients in 14 × 89 mm polyallomer tubes (331372, Beckman Coulter). One aliquot of 12 µgRNA was loaded on each gradient and centrifuged in a Beckman SW41 rotor at 209,490 g × 30°C for 4 h. Gradients were fractionated in 2 ml fractions and 10 µl of each fraction was loaded on gel to analyze RNA43.
Co-immunoprecipitation. Yeast cells carrying genomic GFP-tag fusions of either eIF2G, eIF4E, or eIF4A were transformed with Xrn1, Rta1ΔNLS, and Rta1ΔNLS-XC FLU plasmids (and were grown on 30 μM cycloheximide) in 100 OD units (1 OD unit was similar and an OD600 of ~0.5 was reached. Ten ODs of the culture were harvested by centrifugation. To analyze the RNA distribution in cellular fractions between 0 and 30 min after auxin addition, we obtained mRNA synthesis rates (SR). Whole RNA polymerase II transcribe SR was obtained by summing up all individual genes SR data. Transcriptome data (mRNA levels, RA) were obtained from the hybridization of labeled CDNA onto nylon filters. Total mRNA concentration in yeast cells was determined by quantifying poly-A+ in total RNA samples by oligo-dT hybridization of a dot-blot following the protocol described59 and dividing by average cell volume. mRNA half-lives (HL) in arbitrary units for every mRNA were divided individual RA values by SR ones.

Ribosome-profiling analysis. Ribosome-profiling reads were aligned to the sac-cerevisiae transcriptome (SGD annotation) with bowtie60 using the following settings: “-v 0 -m 1 -i 1.25 -n 1 -t 0.005”. The transcriptome consisted of all CDS flanked by 18 nt of genomic sequence on either side (representing the UTR). Reads mapping to the first 63 nucleotides (18 in 5' UTR + 15 codons) were discarded to remove cycloheximide-induced artefacts. After codon assignment and quantification (analogous to ref.13), raw per gene counts were supplied to the Riborex R-package v2.3.261 to identify significant differences of translational regulation upon Xrn1 knock-down (minMeanCount = 0; FDR < 0.05). As Riborex only provides a fold change estimate for the interaction between condition (t0 vs. t30) and technique (mRNA vs. RPF), all significant genes were grouped according to their behavior relative to the diagonal (=no change of translational control) by obtaining the corresponding moderated fold changes for mRNA and RPF samples separately (as displayed in Fig. 5b). To simplify subsequent analyses, we considered genes whose mRNA log, fold change was smaller than ±0.433 (analogous to ref.13) to have buffered/stable levels of mRNA.

Gene ontology enrichment analysis for the resulting groups was performed using a correction method262 with settings “-moderate” and “-Thier_fdr_correcting = "moderate"”. For visualization, we used REVIGO to define redundant GO terms (http://revigo.irb.hr/) and only considered the top 5 non-dispersable terms.

For the metagenome analysis, per codon RPF counts were normalized by gene length and library size (observed/expected) before being averaged per condition. The averaged normalized footprints were then corrected by the corresponding mRNA fold change relative to t0. Visualization was done using the ggplot2 R-package33.

Analysis of PARs scores. Nucleotide resolution PARs scores for yeast RNAs were obtained from Github (https://github.com/ablewe/prfdb/tree/master/pars/scene_Score.tab) and further processed using the statistical programming language R. For a high resolution analysis of the region surrounding the translation initiation site (TIS) we extracted up to 100 nt of the 5' UTR (if available) as well as the first 300 nt of the coding sequence (CDS) and plotted the mean (na.rm = TRUE) PARs profiles for all the groups studied. PARs scores were also averaged across the 5' UTR, CDS and 3' UTR in order to compare increased or decreased RNA structure in different regions between the different groups; otherwise, a Wilcoxon-Mann–Whitney test was used to detect significant differences in all analyses.

Structure modeling of the chimera Rta1ΔNLS-XC. The sequences of Rta1ΔNLS and Xrn1 were aligned with the sequences of template structures of S. pombe and K. lactis taken from the Protein Data Bank (PDB)63, with codes 3FQD and 3PIF, respectively. We superposed both structures with MATCHMENDER, using CH-MERA64, and detected that the structure of Rta1 deviates from Xrn1 at the position of Arg 653 in chain A of 3FQD, which corresponds to position 884 in the sequence of Rta1, and this was selected for merging both the sequences. The N-terminal domains of both structures are very similar. However, we detected a protruding loop on the structure of Rta1 that collided with the C-terminal domain of Xrn1 (between Ile19 and Gln25 positions in chain A of 3FQD, corresponding to positions Val21 and Pro45 in chain A of 3FQD, aligned with the sequence LEEQPQIVDGVIL of Rat165. The sequence LEEQPQIVDGVIL of Rat1 was modeled with MODELLER57 and optimized with Rosetta58. The structure was modeled with MODELLER57 and optimized with Rosetta58.

Cellular fractionation for mRNA recruitment analysis. Xrn1-AID strain was grown in YPD with 2% glucose at 30 °C until the doubling time between triplets was similar and an OD600 of ~0.5 was reached. Ten ODs of the culture were harvested (−1 WT). In the remaining culture the degradation of Xrn1 was induced for 30 min by adding auxin to a final concentration of 0.5 mM. After 30 min, cells (10 ODs) were harvested by centrifugation. To analyze the RNA distribution between cytoplasm and membrane fractions 240 μl of lysate buffer (50 mM MOPS, 275 mM potassium glutamate, 5 mM Mg Acetate and freshly added 100 μg/ml CHX, 1 mM DTT, 20 U/ml Superscript and Roche protease inhibitors) were added to ODs of RNAs treated samples and centrifuged in a Retsch MM400 at 30 Hz 2 min (four times). The lysate was thawed at 25 °C and centrifuged 10 min at 13,000 x g. The supernatant, corresponding to the cytoplasm,
was transferred to a new eppendorf tube and triton was added to 0.1%. The pellet was resuspended in lysis buffer containing 1% triton and homogenized in a dounce homogenizer. After centrifugation (10 min 13,400 × g) the supernatant contains the membrane fraction. RNA extraction was carried out using the hot-phenol method. 

mRNA distribution between cytosol and membrane fraction was analyzed by reverse transcription quantitative (q)PCR using TaqMan probes and qScript XLT One-Step RT-qPCR Though Mix from Quanta Biosciences. Twelve nanograms of total RNA were loaded and amplified using specific primers. The list of primers is available in Supplementary Data 8.

Electroporation of in vitro transcribed RNA2-Rluc. The protocol used is an adaptation of the one previously described. Reporter RNAs (RNA2-Rluc and pLucA) were in vitro transcribed from a plasmid using the MAXScript T7 In Vitro Transcription Kit (Ambion) and the MAXscript T3 In Vitro Transcription Kit (Ambion), respectively. The transcripts were subsequently capped with ScriptCap™ m7G Gapping System (CELLSCRIPT). Their integrity was evaluated with a denaturing formaldehyde gel. After quantification with a NanoDrop device, RNAs were aliquoted and stored at −80 °C.

Yeast cells (wt and Δxrn1Δ) were treated with lyticase to remove the cell wall and incubated with Sorbitol 1 M. Next, the spheroplasts were incubated with YAPD-Sorbitol 1 M for 90 min at 30 °C to allow cell recovery. Finally, cells were pelleted, resuspended with Sorbitol 1 M, and kept on ice. Electroporation cuvettes (0.2 cm electrode gap, BioRad, Hercules) were kept on ice and RNA2-Rluc and pLucA RNA were added (8 and 0.5 μg, respectively). Yeast spheroplasts were pipetted (180 μl) into the cuvette and pulsed immediately (BioRad gene pulser II: 80 V, 25 μFaraday and 1000 μl, 20–25 ms). Spheroplasts were transferred to ice-cold 2 ml tubes and placed at 30 °C with gentle swirling. After 30 min, cells were pelleted (3 min, 1500 × g), the supernatant was discarded and the pellets frozen immediately in liquid nitrogen (−80 °C). For measurement, cells were resuspended in 50 μl of 1X reporter lysis buffer (Renilla Luciferase activity assay, Dual-Glo® Promega) and vortexed vigorously for 30 s. Cell lysate (20 μl) and 50 μl of luciferase assay substrate were mixed and measured immediately.

Statistical information. All the statistical information is detailed throughout the Figure legends and the methods section.

Reporting summary. Further information on experimental design is available in the Nature Research Reporting Summary linked to this article.

Data availability A reporting summary for this Article is available as a Supplementary Information file. 

Ribosome Profiling and Genomic Run-On (GRO) raw data are available under accession GSE109734 and GSE123326 at Gene Expression Omnibus (GEO). The source data underlying Figs. 1–5, 7, and Supplementary Figs. 1–3, 8, 12 are provided as a Source Data file.

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38. Lotan, R., Goler-Baron, V., Duek, L., Haimovich, G. & Choder, M. The Rpb7p subunit of yeast RNA polymerase II plays roles in the two major cytoplasmic mRNA decay mechanisms Rona. J. Cell Biol. 178, 1133–1143 (2007).

39. Das, S., Saha, U. & Das, B. Cbc2p, Upf3p and eIF4G are components of the DRN (Degradation of mRNA in the Nucleus) in Saccharomyces cerevisiae. FEBS Yeast Res. 14, 922–932 (2014).

40. Galán, R. P. et al. Lsm1-7 complexes bind to specific sites in viral RNA genomes and regulate their translation and replication. RNA 16, 817–827 (2010).

41. Jung, T., Cudmore, K. & Darnell, J. E. Nuclear RNA turnover during stress responses. Mol. Cell. 71, 894–907 (2018).

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Author contributions

B.B.-M., L.D.C.-M., J.D., M.C., S.A.L., D.D.N. and J.E.P.-O. were responsible for the conceptualization of the project. B.B.-M., D.D.N., J.J., L.D.C.-M., J.G.-M., J.E.P.-O., M.C., S.C., S.A.L. and M.E.-G. designed and adapted the methodology. R.B., J.G.-M. and B.O. implemented computer code and algorithms. B.B.-M., L.D.C.-M., J.E.P.-O., J.G.-M., R.B. and B.O. applied statistical or computational techniques to analyze the data. B.B.-M., L.D.C.-M., J.J., J.G.-M., M.E.-G. and S.C. performed the experiments. B.B.-M. performed experiments regarding the BMV RNA system, the Xrn1-AID system, poly-some profiling, Ribosome Profiling, electroporation, and the generation of the respective plasmids and strains. L.D.C.-M. performed experiments regarding the BMV RNA system, co-immunoprecipitation experiments with Xrn1 and Rat1 constructs, polychrome-profiling experiments, functional validation of GFP-fused strains and the generation of the respective plasmids. J.J. performed ER recruitment analysis and RLM-RACE experiments, functional validation of GFP-fused strains and the generation of the respective plasmids. L.D.C.-M. performed experiments regarding the BMV RNA system, co-immunoprecipitation experiments with Xrn1 and Rat1 constructs, polychrome-profiling experiments, functional validation of GFP-fused strains and the generation of the respective plasmids. J.J. performed ER recruitment analysis and RLM-RACE experiments. J.G.-M. performed genome-wide TR and HL experiments. M.E.-G. performed RLM-RACE experiments. S.C. performed initial co-immunoprecipitation assays. R.B. and J.G.-M. were responsible for data curation. B.B.-M., J.D. and M.C. wrote the original draft. All authors reviewed and edited the manuscript.

Additional information

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