Transcription feedback dynamics in the wake of cytoplasmic mRNA degradation shutdown

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ABSTRACT

In the last decade, multiple studies demonstrated that cells maintain a balance of mRNA production and degradation, but the mechanisms by which cells implement this balance remain unknown. Here, we monitored cells' total and recently-transcribed mRNA profiles immediately following an acute depletion of Xrn1—the main 5'-3' mRNA exonuclease—which was previously implicated in balancing mRNA levels. We captured the detailed dynamics of the adaptation to rapid degradation of Xrn1 and observed a significant accumulation of mRNA, followed by a delayed global reduction in transcription and a gradual return to baseline mRNA levels. We found that this transcriptional response is not unique to Xrn1 depletion; rather, it is induced earlier when upstream factors in the 5'-3' degradation pathway are perturbed. Our data suggest that the mRNA feedback mechanism monitors the accumulation of inputs to the 5'-3' exonucleolytic pathway rather than its outputs.

INTRODUCTION

Gene expression is a multistep process that starts at the nucleus, where structural (e.g. histones) and regulatory factors interact to facilitate transcription by the general transcription machinery and RNA polymerase II (PolII). During transcription, nascent mRNA molecules are capped at their 5' end with a nucleotidylic-resistant nucleotide (m7G), spliced, cleaved, and polyadenylated. Protein-mRNA complexes are then exported from the nucleus to undergo translation in the cytoplasm by ribosomes. To allow for dynamic gene expression, most mRNA species are actively degraded by cells within a short time frame (minutes in yeast to hours in mammalian cells (1,2)). mRNA degradation can be triggered by various quality control mechanisms (e.g. nonsense-mediated decay), but degradation is also thought to be coupled to the translation of the mRNA by ribosomes (3,4). Seminal work established the main degradation pathway in eukaryotes: mRNA is deadenylated by the Ccr4–Not complex, decapped by the Dcp1-Dcp2 complex, and degraded by the highly processive 5'-3' exoribonuclease Xrn1 (5–10). An important alternate route involves the exosome, which degrades mRNA from its 3' end following deadenylation by the Ccr4–Not complex. Additional less common endonucleolytic degradation pathways were also described (9,11).

As presented above, this process is largely unidirectional, namely, messages are generated in the nucleus, exported, translated, and degraded with no information flow back to the nucleus. However, mounting evidence from the past decade suggests that transcription in the nucleus is coupled to degradation in the cytoplasm. This coupling was demonstrated along two main branches of evidence. The first is gene-specific regulation of transcript fate by nuclear signals, e.g. replacing the promoter of a gene can alter its transcript half-life or cytoplasmic localization (12–16). In these cases, the functional implications are generally thought to be carried out by different mRNA-binding proteins that are exported with the transcript (17). Another line of evidence linking nuclear transcription and cytosolic degradation is a global phenomenon (termed mRNA buffering or mRNA homeostasis). It was demonstrated that large-scale perturbations to the degradation machinery are compensated by the transcription machinery (18–20) and vice versa (15,16,21–27). In these cases, the underlying mechanisms for the observed global transcription-degradation coupling remain contested and speculative. Proposed mechanisms involve several main components, including Rpb4/7 (POLR2D/G in humans) (27–29), Pab1/Nab2 (30–32), the Ccr4–Not complex (16,33), Snf1 (AMPK) (34,35), and Xrn1 (18,19).

Xrn1 is the main cytosolic 5'-3' exonuclease in eukaryotic cells (6,8,36–38). Xrn1 knockout causes developmental and fertility defects in multicellular organisms (39,40), and its knockout in yeast was shown to affect cell size and growth rate, hinder growth in certain stress conditions, and cause spindle-pole separation defects (41–43). In

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two important works published in 2013 Xrn1 was implicated in the coupling between degradation and transcription (18,19). However, these studies arrived at opposite conclusions about the role of Xrn1. Haimovich and colleagues reported that Xrn1 knockout maintains global mRNA levels. They could explain the observed buffering by attributing to Xrn1 (and related RNA binding factors) a role as an transcriptional activator that acts directly on chromatin. Conversely, in a systemic screen of RNA processing factors, Sun and colleagues reported that Xrn1 knockout results in the most significant increase in total mRNA levels, which is the result of a significant decrease in degradation rates and a slight increase in global transcription rates. Mechanistically, they linked Xrn1 levels to the transcript levels of a negative transcriptional regulator - Nrg1. Since then, other studies with Xrn1 knockout/knockdown exhibited various transcriptional effects (44–50). Similarly, the Ccr4–Not complex which is crucial for mRNA deadenylation and degradation was also implicated in transcription regulation (16,33,51,52). Interestingly, in the systemic screen by Sun et al. some components of the Ccr4–Not complex incur pronounced deviations from the wildtype strain (18).

To summarize, there is ample evidence that mRNA buffering takes place in perturbed yeast cells, and probably in higher eukaryotes, but there is little understanding of the mechanisms underlying various contradicting observations. We reasoned that if a feedback process is at play, the dynamics of the process can shed light on its mechanism. However, we found little information about the dynamics of the process, as most studies were performed in knockout strains at steady-state growth conditions. In several notable exceptions the time scale of the feedback was determined to be in the order of several minutes (23) and up to an hour (18,21), but these were observed in different settings, and their generality is unclear.

Several of the works mentioned here applied comparative Dynamic Transcriptome Analysis (cDTA (22)) to samples, which measures mRNA levels and monitors recently-transcribed molecules by pulse labeling RNA with uracil analogs. We adapted the cDTA protocol to a high-throughput, quantitative, and sequencing-based version that we termed cDTA-seq. The cDTA-seq technique allowed us to monitor Xrn1 depletion from cells in high resolution, and observe the dynamics of mRNA accumulation and cells’ adaptation to this perturbation. Utilizing metabolic labeling data we identify a delayed global reduction in transcription, which results in a return to wildtype mRNA levels after several hours. We further expanded our data to multiple other RNA processing factors and found that the transcriptional response was not unique to Xrn1, and it also occurred upon depletion of Dcp2 and Not1. Interestingly, we find that the transcriptional response initiates earlier when upstream components in the 5′-3′ pathway are perturbed, suggesting that the trigger for the transcriptional response is sensed upstream of the degradation pathway. These results provide a rich resource for studying the cellular response to perturbations in the general mRNA degradation machinery, and our analysis provides insights into the basic properties of the mechanism underpinning the mRNA buffering phenomenon.

**MATERIALS AND METHODS**

### cDTA-Seq

**Sample preparation.** For each experimental batch, *Kluyveromyces lactis* (KL) yeast cells were grown to log-phase, fixed in frozen methanol and the KL-spiked methanol was equally pre-distributed in deep well plates in large volumes to minimize the effect of pipetting errors. The plate with pre-spiked methanol was kept at −80°C until the fixation of *Saccharomyces cerevisiae* (SC) samples. Immediately before SC fixation, sample biomass was measured by optical density at 600 nm and the culture was metabolically labeled by 4-thiouracil (4tU, Sigma). This was performed simultaneously on the entire plate using a pipette robot (Tecan EVO200), after which cells were immediately fixed in pre-frozen and pre-KL-spiked methanol (600 μl pre-spiked frozen methanol to 500 μl SC cells). Samples were then kept at −80°C up to several weeks.

**RNA purification.** Cells fixed in frozen methanol were washed twice in ddw and RNA purification was performed as previously described (53) with minor modifications. Briefly, RNA was released from the cells by digestion with Proteinase K (Epicenter) in the presence of 1% SDS at 70°C. Cell debris and proteins were precipitated by centrifugation in the presence of potassium acetate. RNA was then purified from the supernatant using nucleic acid binding plates (96-well, 800 μl UNIFILTER Microplate, GE Healthcare) in the presence of 0.1 mM DTT, eluted in 1 mM DTT, and stored at −80°C.

**Metabolic labeling, and adaptation of SLAM-seq to yeast.** Metabolic labeling of recently transcribed RNA molecules was done as previously described (54,55). Briefly, 4tU was dissolved in NaOH and added to cells at a final concentration of 5 mM 4tU for the indicated times (6–10 min). To avoid pH change as a result of NaOH addition, MES buffer was added to the media prior to growth. RNA purification was performed as described above. Total RNA was subjected to thiol(SH)-linked alklylation by iodoacetamide (Sigma, 10 mM) at 50°C for 15 min, the reaction was stopped with 20 mM DTT. RNA was purified using nucleic acid binding plates (96-well, 800 μl UNIFILTER Microplate, GE Healthcare) and was stored with RNase-inhibitor at −80°C.

**polyA RNA-seq library preparation.** Library preparation was done as previously described (56). Total RNA was incubated with oligo-dT reverse transcription primers with a 7 bp barcode and 8 bp Unique Molecular Identifier (UMI) at 72°C for 3 min and transferred immediately to ice. RT reaction was performed with the SMARTScribe enzyme (Clontech) at 42°C for 1 h followed by enzyme inactivation at 70°C for 15 min. Barcoded cDNA samples were then pooled and purified using SPRI beads x1.2 (Agencourt AMPure XP, Beckman Coulter). DNA-RNA hybrids were tagmented using Tn5 transposase (loaded with oligos Tn5MEDS-A, Supplementary Table S2), and 0.2% SDS was added to strip off the Tn5 from the DNA (57), followed by a SPRI x2 cleanup. Barcoded Illumina adaptor
sequences (Supplementary Table S2) were added to the tagged DNA by PCR (KAPA HiFi HotStart ReadyMix, Kapa Biosystems, 12 cycles). And the DNA was cleaned with an x0.8 SPRI procedure. Libraries were sequenced using Illumina NextSeq-500 sequencer.

Data processing.

Demultiplexing. Pooled libraries were demultiplexed using Illumina’s bcl2fastq (version 2.20.0). Internal (sample) barcodes were demultiplexed with an awk command, not allowing any barcode errors in both cases.

Genome alignment. Prior to read alignment, we prepared several versions of the SC and KL genomes. First, we converted both genomes to accommodate alignments of partially (T→C)-converted reads. This was achieved by converting all the observed Ts in the genome to Cs per strand. In addition, we also generated a redacted version of the KL genome in which any 18-mer that is reference strand. In addition, we also generated a redacted genome without ACG conversion. This results in an ACG-only genome with one contig per read. And more generally, the likelihood of a collection of reads, $R(p_r)$, is:

$$L(p_r) = Pr(R \mid p_r) = \prod_{r \in R} [p_r R_{Y(r),X(r)} + (1 - p_r) O_{Y(r),X(r)}]$$

Thus, assuming the error and conversion rates are global (Supplementary Figure S1E), the likelihood is a function of a single parameter—$p_r \in [0, 1]$, and its maximum can be efficiently calculated given the other model parameters (see supplementary material).

Steady-state model. We assume the following dynamic system:

$$\dot{M} = \pi C - \delta M$$

$$\dot{C} = \gamma C$$
where $M$ is the total amount of mRNA, C is the number of cells (or, alternatively this can be biomass), $\pi$ is a gene-specific transcription rate per cell, $\delta$ is the gene-specific degradation rate, and $\gamma$ is the growth rate. Redefining $R = \frac{\pi}{\gamma}$, i.e. mRNA/cell, the system simplifies to:

$$R = \pi - (\delta + \gamma) R$$

We also assume that 4tU labeling does not disturb cells from their exponential growth pseudo-steady-state (Supplementary Figure S1G, H), in which case the steady state value of the system is given by:

$$R_{ss} = \frac{\pi}{\delta + \gamma}$$

Additionally, the differential equation yields the following dynamics for molecules transcribed during the labeling period ($N$):

$$N(t) = \frac{\pi}{\delta + \gamma} \left(1 - e^{-(\delta + \gamma)t}\right)$$

Thus, the proportion of labeled transcripts, $p_r$, evolves with labeling time ($t$), as follows:

$$p_r(t) = \frac{N(t)}{R_{ss}} = 1 - e^{-(\delta + \gamma)t}$$

**Half-life estimation.** Briefly, we fit/measure several global parameters (detailed description is given in the supplementary material): incorporation probability ($\xi$), error probability ($\epsilon$), conversion lag time ($t_0$), and growth rate ($\gamma$). Once these are determined, the only gene-specific parameter is the degradation rate ($\delta$), which can be calculated by fitting the BMM model to the data to obtain $p_r$, and by inverting the last equation we obtain the degradation rate:

$$\delta = -\frac{1}{t} \ln(1 - p_r) - \gamma$$

Which can be further transformed to a half-life ($ln(2)/\delta$).

**Auxin-induced degradation**

In all experiments, yeast cells were grown in YPD at 30°C overnight, supplemented with a final concentration of 10 mM MES pH 6.0 buffer. 60 minutes prior to the beginning of the time course, samples with OD ranging from 0.3 to 0.6 were split to deep well plates and grown at 25°C with constant pipette mixing during the time course. At indicated times auxin (3-indolo acetic acid, Sigma) was added at a final concentration of 1–2.5 mM (Supplementary Table S4). Auxin stock was dissolved in DMSO to 2.5 M, and diluted 1:1 in 1 M NaOH before being added to samples to prevent sedimentation in the aqueous medium. When mock treatment was appropriate, the same amount of DMSO and NaOH was added to control samples.

**cDTA-seq calibrations**

**Spike-in titration.** As a standard for mRNA levels in cDTA-seq experiments, we spike-in fixed amounts of exogenous cells to each sample. To evaluate this strategy, we performed a titration of spike-in cells (KL) in the 1%-5% range into an SC sample. SC and KL cells were grown to log-phase (od 0.5) and fixed separately in cold methanol (7.5 ml cells in 9 ml methanol). Fixed cells were mixed at varying ratios (1–5% KL) and were kept at −80°C. RNA purification and polyA RNA-seq libraries were prepared as described above.

**DNA-seq validation.** When comparing mRNA levels between samples, one can use different measures, such as mRNA/cell, mRNA/volume or mRNA/biomass. In our view, biomass or volume are the most relevant normalizing measures for the question of mRNA homeostasis, but cell counting remains common (22). In many experimental settings, the optical density (OD) of the culture is a good proxy for cell counts, but more generally OD quantifies the total sample biomass (which is correlated to cell counts (61)). In our experiments, there was a concern that different genetic backgrounds and cell states can affect the OD/cell ratio (62,63). We verified that our OD measures correspond to cell counts by comparing the DNA content extracted from 93 samples from various genetic backgrounds to their OD. To estimate the ratio between the number of SC and KL cells we extracted nucleic acids from various samples that underwent cDTA-seq and were therefore spiked-in with KL cells. These samples included various AID-tagged proteins (Dcp2, Xrn1, Rat1, Fcp1, Sth1, Med14, Pop2, Spt6) grown overnight with varying degrees of auxin, resulting in a wide range of growth rates (1.5–4 h doubling time) and morphological phenotypes (e.g. Xrn1-depleted cells are larger). When we compared the ratio between SC DNA and KL DNA in each sample with the OD of that sample, we observed a high correlation ($R^2 = 0.88$, Supplementary Figure S1B).

To prepare DNA, RNase (Sigma, 11119915001) was added to nucleic acid extracted as mentioned in the cDTA-seq protocol (0.1 µl RNase to 100 ng RNA in a reaction volume of 20 µl) and incubated at 37°C for 30 min. The remaining nucleic acid - genomic DNA - was fragmented using Tn5 transposasase (loaded with oligos Tn5MEDS-A and Tn5MEDS-B, Supplementary Table S2), and 0.2% SDS was added to strip off the Tn5 from the DNA (57). Barcoded Illumina adaptor sequences (Supplementary Table S2) were added to the tagged DNA by PCR (KAPA HiFi Hot-Start ReadyMix, Kapa Biosystems, 20 cycles), and the libraries were cleaned with a ×0.9 SPRI procedure. Libraries were sequenced on an Illumina NextSeq-500. Reads aligned to the SC genome were manually inspected to have uniform genomic distribution (with exceptions in the rDNA locus, transposable elements, etc.). Reads were also aligned to a reduced SC genome (removing any 20-kmer found in KL) and to a similarly reduced KL genome to obtain estimates of the amount of DNA from each organism in the sample. The ratio between these numbers was compared to the OD of each sample (both measures were normalized to their median for visualization in Supplementary Figure S1B). Outliers were not of a specific strain, batch, or condition, point-
ing to measurement noise (probably in the OD), rather than inherent biases in the technique.

Western blots
Yeast lysates were prepared as previously described (64) and proteins were analyzed using standard western blotting procedures with anti-FLAG M2 (Sigma F1804), and anti-Myc (Sigma M4439 clone 9E10).

Growth assays and OD measurements
Optical density was collected using a Tecan Infinite F200 for 96-sample plates. Each well/sample was measured at five different positions and the median value was used as the OD measure. A background level was measured in each plate and was subtracted from the measured OD.

Growth assays were conducted using a Tecan Freedom Evo 2000 liquid handling station. The 200 μl 96 sample plate was incubated at 30°C, for >24 h with an automatic scheduled OD measurement in the Tecan Infinite F200 executed every hour (65).

A log-linear fit was applied to each consecutive set of 10 data points, and the minimal doubling time was determined by the fit with the highest slope among the fits that passed the 5% significance threshold (test for linear fit, Bonferroni-corrected).

Microscopy
Samples along the depletion time course were fixed in 1% formaldehyde for 15 min and quenched in 125 mM Gly. Bright-field images were automatically collected in multiple fields per sample along an 8 μm z-stack (1 μm step size) using a Scan∧R system (Olympus). Cells were segmented using freely available yeast segmentation software (66).

Single-molecule FISH
Sample preparation. In the first experiment (Figure 3), 120 ml Spt6AID and 120 ml Xrn1AID cells were grown to mid-log, and split. Half of the culture was supplemented with auxin (final 2.5 mM) and half with mock treatment (50% DMSO, 0.5M NaOH at equal volume). After one hour of incubation at room temperature samples were fixed in 5% formaldehyde and prepared with a 3-component gaussian mixture model. (erosion/dilation/convolution) to account for uneven illumination. Segmented nuclei were filtered by size and manually inspected. Single-molecule counts were obtained using custom-made MATLAB software (69).

Microscopy. Images were acquired with a 100 × 1.4 oil UPLS APO objective, using an Olympus IX83 based live-imaging system equipped with CSU-W1 spinning disc (SC-MOS digital Scientific Grade Camera 4.2 MPixel, Oxford Instruments, Abingdon, UK). For each sample, at least 4 different positions were chosen. In each position, three-channel Z-stacks images were taken with a step size of 200 nm for a total of >8 μm: bright-field image, 488 nm laser with 100 mW; DAPI image, 405 nm laser with 120 mW, and exposure time of 250 ms; mRNA image, 561 nm laser with 100 mW and exposure time of 1000 ms. Each z-plane image was of size 2048 × 2048 pixels.

Image analysis. Nuclei and single molecules were segmented from the DAPI channel using a MATLAB script that uses basic image processing steps (erosion/dilation/convolution) to account for uneven illumination. Segmented nuclei were filtered by size and manually inspected. Single-molecule counts were obtained using custom-made MATLAB software (69).

dpolyA FACS analysis
Cells from the second FISH experiment were stained with a polyT-TAMRA probe. At least 7500 valid events (cells) were collected per sample on an Amnis CellStream high throughput flow cytometer after excitation with a 561 nm laser and acquisition with a 561–583 nm filter.

DNA staining
Wildtype, Xrn1AID and Sth1AID cells were split and exposed to auxin along a time course of 4.5 h. At the end of the time course, cells were collected into pre-frozen EtOH (final 70%). Cells were washed in 50mM Tris–HCl pH 8 (Sigma), incubated with RNaseA (Sigma R4875, final 1 mg/ml in 50 mM Tris–HCl pH 8), Proteinase K (Sigma P2308, final 2.5 mg/ml in ddw), and with SYBR green (Molecular Probes S7567, diluted 1:1000 in 10 mM Tris–HCl pH 8, 1 mM EDTA pH 8). Following an additional wash, cells were briefly sonicated and analyzed by FACS with a 525-centered filter (BD LSRII system, BD Biosciences). Cell-cycle phases were determined by fitting the DNA content distribution with a 3-component gaussian mixture model.

Strains and plasmids
Saccharomyces cerevisiae yeast strains used in this study are provided in Supplementary Table S1. Validations of AID degradation are also detailed in this table (see also Supplementary Figure S8 for a summary of western blots).

Yeast strains were generated using the LiAc transformation method (70). Auxin inducible degradation domain was PCR-amplified from plasmid pNat-AID*-9MYC or pHyg-AID*-6FLAG using matching primers (Supplementary Table S2) and introduced into TIR1 expressing cells immediately before the target gene stop codon (plasmids and TIR1 cells were a gift from Ulrich lab (71)).

Fresh Xrn1 knockouts were prepared using plasmids pNat-AID*-9MYC or pHyg-AID*-6FLAG, and oligos (xrn1_ko_F, xrn1_deg-R, see Supplementary Table S2)
RESULTS

High-throughput, quantitative dynamic transcriptome analysis by sequencing (cDTA-seq)

To study the dynamic equilibrium of mRNA in cells a dynamic measurement is needed. We aimed to assess absolute mRNA levels and transcription rates in multiple strains in detailed time-course experiments, resulting in hundreds of samples. Therefore, we developed a protocol (Figure 1A) inspired by Sun et al. (22), in which we spike in a constant amount of cells from a close yeast species for mRNA quantification, combined with a brief pulse with 4-thiouracil (4tU) to quantify recently-transcribed molecules. Following similar works (54,55), we alkylate the 4tU nucleotides with iodoacetamide, which results in their subsequent reverse-transcription to G instead of A (T→C on the sense strand). The conversion allows for the detection of 4tU at the same time as quantifying mRNA levels with RNA-seq. Thus, we avoid additional biochemical separation and unknown losses in each sample, which allows for accurate estimation of the fraction of recently-transcribed molecules (Figure 1B). We incorporated these steps into an RNA extraction and 3’ mRNA sequencing protocol we had developed that was specifically aimed at high throughput measurements (56). The final protocol (Figure 1A-B) allows a single person to quantify the transcriptome, including recently-transcribed mRNA, from 192 samples in a single day (Materials and Methods).

To study changes to mRNA concentration, defined as molecules/biomass (see Materials and Methods), we needed two points of reference per sample - a measure of mRNA levels and of biomass. We verified our ability to quantify relative sample mRNA levels by spike-in titration (Supplementary Figure S1A, \( R^2 = 0.99, P < 10^{-13} \)). For biomass estimation, we measured optical density (OD), and we verified that there are no gross deviations between biomass and cell counts in the different genetic backgrounds we worked with (see methods, Supplementary Figure S1B, \( R^2 = 0.88, P < 10^{-47} \)).

Next, we performed a 4tU labeling time course experiment and observed a significant increase only in T→C conversions (Figure 1C, Supplementary Figure S1C). To verify that we can detect changes in transcription rates, we also performed this analysis after transcription inhibition with thiolutin for 15’ and pulse-labeled the samples with 4tU to observe a >50% reduction in labeled molecules (Figure 1C). As previously observed (60), individual reads exhibit multiple conversion events (Figure 1C) bolstering confidence that they arise from newly transcribed molecules. We used this observation to estimate the percent of molecules that were transcribed during the labeling period (\( p_r \)) by fitting a probabilistic model to the data (Figure 1B, methods). We verified that the model fitted the observed data well (Figure 1D, Supplementary Figure S1D), and that the data is best explained by an increase in the proportion of recently-transcribed molecules (\( p_r \)), rather than changes to incorporation efficiency or other artifacts (Supplementary Figure S1E).

Assuming a steady-state and a first-order model for mRNA (Figure 1B, Supplementary Figure S1F–H), a half-life was calculated per transcript (Figure 1B, E, F, Supplementary Figure S1I, J). We compared our estimates with four published works that measured or estimated transcript half-lives using various methods (Figure 1G) (1,18,19,72). Our technique is clearly correlated to other studies, but there are global discrepancies between half-life estimates that were previously noted (Supplementary Figure S1K) (22,73,74), and are potentially explainable by technical discrepancies (e.g. maturation and polyadenylation time can cause an offset between techniques, see also Supplementary Figure S1L).

We conclude that cDTA-seq can be used to determine the relative mRNA level and estimate the half-life of individual transcripts from a single measurement, with the caveat that the absolute half-life numbers from any technique should be treated with care. Therefore, we include a wildtype sample in each experimental batch, which is used as a control for physiological anomalies and technical discrepancies between experiments.

Turnover rates are slowed in the absence of Xrn1, but Global mRNA levels are maintained

Previous studies found Xrn1 knockout to cause a global increase in mRNA levels per cell (18), or conversely, an unchanged level of mRNA (19,49). To address the question of absolute mRNA levels, we applied cDTA-seq to quantify the changes in mRNA levels in the absence of Xrn1 and the underlying mRNA dynamics. Given that Xrn1 is a major mRNA degradation factor, a knockout of Xrn1 is expected to cause an increase in total mRNA levels. However, using our protocol we do not observe any difference in global mRNA levels (normalized to biomass) when comparing freshly deleted Δxrn1 strains to wildtype cells (Figure 2A). Further, despite significant changes to ~400 transcripts (Supplementary Figure S2A), the overall distribution of mRNA was unchanged (Figure 2B). We then turned to examine the changes to transcription and degradation rates in the absence of Xrn1. Contradicting reports in the literature argue that transcription slightly increases (18) upon Xrn1 knockout, or is markedly reduced (19). In our data (two biological replicates in two fresh knockouts) we find a significant reduction in the fraction of recently-transcribed molecules from 22.8% to 14% (Figure 2A, t-test \( P < 0.004 \)). Using our data, we were able to estimate the degradation and transcription rates for ~4800 transcripts (>70% of genes). Unlike mRNA levels, we do observe a clear reduction in degradation rates (median decrease of 40%, with 746 transcripts becoming completely stable, Figure 2C). A corresponding reduction in transcription rates (median decrease of 70%, Supplementary Figure S2B) can be inferred from the mRNA levels and degradation rate estimates. When we examined the relationship between changes in production and degradation rates per transcript we found strong agreement, consistent with buffering (\( r = 0.87, P < 10^{-300} \), Figure 2D, Supplementary Figure S2C).

To better understand the global changes in degradation and transcription rates, we looked for functional signa-

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Oligos used in this study are listed in Supplementary Table S2.
Gene X changes in mRNA levels or degradation and transcription. We tried using various gene effects (Supplementary Figure S2G). Having excluded functional explanations, we tried using various gene sets in the knockout studies (Supplementary Figure S2F). Similarly, we found that in our data virtually only ribosomal protein genes and ribosome biogenesis genes were exceptional, pointing to indirect growth effects (Supplementary Figure S2G). Having excluded functional explanations, we tried using various gene/转录 features and sequence information to explain observed changes in mRNA levels or degradation and transcription rates. However, our models only explained a small fraction of the observed variance (Supplementary Figure S2H-J, supplementary note).

We conclude that cells maintain their global mRNA levels in the absence of Xrn1 by reducing global transcription rates, but the specific details underpinning the homeostasis are obscure in the knockout strain.

Conditional Xrn1 depletion reveals a global but transient increase in mRNA levels

We reasoned that if Xrn1 is a major exonuclease of mRNA (6,18), its removal must exert an effect that is somehow buffered by cells. To test this hypothesis, we set out to repeat the cDTA-seq experiment in an Xrn1 conditional knockdown strain. We generated strains in which Xrn1 is tagged with an Auxin Inducible Degron (AID) (71,77) and
validated that the Xrn1 protein is depleted rapidly from cells (half-life of ~6’, Figure 3A, Supplementary Figure S5G). To monitor the effects of Xrn1 depletion, we grew cells to an exponential growth phase and performed a 4-h time-course experiment following Xrn1 depletion (Figure 3B).

Multiple replicates reveal that mRNA levels increase significantly following Xrn1 depletion (20%-40% across multiple experiments and repeats; Figure 3B). However, after about 70 min from the time of auxin addition, the accumulation trajectory inverts, and we observe a decrease in global mRNA levels, resulting in a return to basal mRNA levels (90–105%) within 4 h. Importantly, the mRNA profile also converges to the Xrn1 knockout mRNA profile (Supplementary Figure S3A), validating the effectiveness of Xrn1 depletion.

We call the initial time period (0–55 min) the accumulation phase, the subsequent time (55–95 min) the adaptation phase, and during the final reversion phase cells settle back to their initial mRNA levels (95 min and onwards).

An examination of the mRNA profile along the time course reveals that it is not the result of an increase in specific highly-expressed transcripts, but that virtually all transcript levels are transiently increased (Figure 3C). We validated the increase in mRNA levels following 60 min of mock or auxin treatment by single-molecule RNA-FISH with probes targeting four different transcripts and recapitulated the results from our spike-in-normalized mRNA-seq counts (Figure 3D, E, Supplementary Figure S3B). Notably, in a different time course FISH experiment, we observed that the reduction in the smFISH signal is slower than the observed reduction in the cDTA-seq signal, and we have evidence to suggest that this is a result of the previously reported accumulation of non-polyadenylated transcripts in Xrn1-depleted cells, which are not captured in the cDTA-seq protocol (Supplementary Figure S3C, D) (8).

These results highlight the ubiquitous nature of degradation by Xrn1 as evident by the immediate increase in the vast majority of transcripts and reveal the dynamics leading to homeostatic mRNA levels.

mRNA accumulation correlates to transcription rates

While the response to Xrn1 depletion seems ubiquitous there are significant and reproducible differences between the response rate of different transcripts (Figure 3F). Given the direct nature of the perturbation, we wanted to test whether the observed changes to mRNA levels are consistent with a first order model for mRNA, whereby the immediate change in mRNA following a decrease in degradation is mainly a function of individual transcript production rate (see supplementary note). To test this prediction, we fit each transcript with a linear model for the change in mRNA during the accumulation phase of the response (Figure 3F). As predicted, the change in many transcripts is consistent with a linear increase (Supplementary Figure S3E, 10% FDR, $R^2 > 0.29, 2383/5164$ transcripts), and where we cannot reject the null hypothesis of constant levels, it is mostly due to sampling noise (low expression levels, see for example Msn2 and Suc2 in Figure 3F). We tested the correlation of the fitted slopes to the pre-perturbation transcription and degradation rates and found that the strongest correlation is to the production rate, as expected by a first-order model (Pearson $r = 0.55, P < 10^{-187}$, Figure 3G, Supplementary Figure S3F). To account for the potentially confounding effect of expression levels, which are correlated to both production and the slope (Figure 3G), we also calculated the partial correlation, which remained significant (partial Pearson $r = 0.28, P < 10^{-45}$, Supplementary Figure S3F). Furthermore, this dependency is significantly accentuated in transcripts with a longer half-life (Supplementary Figure S3G), consistent with masking effects of residual degradation following Xrn1 depletion.

Therefore, during the accumulation phase, mRNA increase is consistent with a scenario where degradation is abruptly reduced, while transcription remains largely unchanged.
Metabolic labeling through Xrn1 depletion uncovers a delayed global transcriptional adaptation

During the adaptation phase mRNA levels stop increasing and eventually revert to WT levels. How does this occur within a few hours? Since mRNA level is at a balance of transcription, degradation, and dilution by growth, there are multiple possible explanations.

Cells can adapt to reduced Xrn1-dependent degradation rates by activating alternative mRNA degradation mechanisms, increasing the dilution rate (which requires faster cell division and growth), decreasing transcription rates, or they can respond in some combination of these mechanisms.

We first tested the hypothesis that cells adapt by increasing their division rate or volume. We monitored various physiological aspects of cellular growth following Xrn1 depletion and found no significant inflection points in optical density, cell size, cell counts, colony-forming units, or cell-cycle fractions within two hours of auxin addition (Supplementary Figure S4A–E). Notably, some of these measures
do eventually change, as previously reported for the knock-out strain (e.g. increase in cell size, Supplementary Figure S4B). We also measured the growth rate in the absence of Xrn1 (knockouts and AID strains) and found a slower growth rate by 20–30% (Supplementary Figure S4F, G). We, therefore, exclude the possibility of volume or growth increase as possible explanations for the observed adaptation in mRNA levels within 2 h.

Another possibility is a compensatory increase in Xrn1-independent degradation. Xrn1 is the main 5′-3′ RNase in the cytoplasm, so the 3′-5′ degradation branch may be compensating for its absence. To test this hypothesis, we AID-tagged two components of the SKI complex (Sk12, Sk18) that were shown to be required for 3′-5′ degradation by the exosome (78) in addition to the Xrn1 AID tag. While these double-AID strains exhibited significantly slower growth when exposed to auxin (Supplementary Figure S4F, G), their immediate mRNA response to depletion was virtually identical to the Xrn1-AID strain (Supplementary Figure S4H), suggesting that the 3′-5′ degradation branch, as mediated by the SKI complex, does not take an active part in the observed reduction in mRNA levels.

To test the remaining possibility of transcriptional reduction, we examined the 4tU-labeled data obtained from cDTA-seq following Xrn1 depletion (Figure 4A). During the adaptation phase (55%–95% following Xrn1 depletion) we observed a concerted and significant reduction in transcription (Figure 4B–D). Indeed, even genes that were induced immediately following Xrn1 depletion show a significant decrease at this point (Figure 4C, D), suggesting global repression of transcription, which we term the transcription adaptation response.

Transient mRNA accumulation is recapitulated but dampened when upstream factors along the 5′-3′ degradation pathway are depleted

Xrn1’s function is largely attributed to 5′-3′ degradation after deadenylation and decapping (Figure 5A). We reasoned that we could pinpoint the molecular constituent sensed by cells by perturbing other factors in the mRNA degradation network and studying the changes to the transcription adaptation response. Therefore, we AID-tagged multiple components of this intricate network—Not1 (Cdc39), Dis3 (Rrp44), Rrp6, Rat1, Pop2, Pab1, Dcp2, Pan3, Nrd1, Sen1, Nab3 and Ccr4 (Figure 5A). We performed a depletion time-course experiment in these strains and applied cDTA-seq to hundreds of samples.

The changes in mRNA in this large dataset revealed the compartment, pathway, and protein-complex interactions between the depleted factors. For example, the effect of depletions of Rrp6, and Dis3—subunits of the nuclear and core exosome respectively (79)—are significantly correlated as expected, but are also correlated with depletion of other nuclear proteins (Figure 5B,D). We wanted to further dissect the response to Xrn1 depletion and focused on Xrn1-correlated factors - Rat1, Rrp6, Dis3, Not1, Dcp2 (Supplementary Figure S5A). We examined the overall mRNA profile following the depletion of these five additional targets (Figure 5E, Supplementary Figure S5B), and it became clear that responses upon Dcp2 and Not1 depletion were significantly more similar to Xrn1 depletion than all other ones (Supplementary Figure S5A).

Not1 and Dcp2 act upstream to Xrn1 in the 5′-3′ mRNA degradation pathway - Dcp2 is the catalytic component of the main decapping complex, and Not1 is the (essential) scaffold of the Ccr4–Not deadenylation complex. In all three depletion time courses (Xrn1, Dcp2, Not1) we observed an accumulation of mRNA followed by a reduction in mRNA levels, consistent with a general feedback mechanism that is triggered when this pathway is perturbed (Figure 5E, F). We compared the transcript profiles along the depletion time course (Figure 5F) and observed an overall high correlation between the perturbations (10−300 < P < 10−63, Supplementary Figure S5A). More specifically it seems that mRNA accumulation is most pronounced in the case of Xrn1 depletion, slightly dampened when Dcp2 is depleted, and further muted when Not1 is depleted (Figure 5E, F).

Further examination of the data revealed a subset of ~13% of transcripts that were more sensitive to Not1 depletion (Figure 5C, F). Functional analysis of these transcripts shows a strong enrichment for transcripts of proteolysis-related genes (q < 10−8, Supplementary Figure S5E). A link between Not1 and proteasome transcript regulation was reported in the literature, and recently, a co-translational complex assembly mechanism was suggested to be mediated by Not1 (80–83). The rapid and prominent increase in these transcripts, largely without a concomitant increase in transcription (Supplementary Figure S5F) suggests that these transcripts are especially susceptible to Ccr4–Not-dependent degradation directly or via the 3′-5′ degradation pathway. These results expand the previously reported link between Ccr4–Not and post-transcriptional regulation of the proteasome. Importantly, even in the Ccr4–Not-sensitive transcript cluster, mRNA accumulation following depletion of Xrn1 and Dcp2 is consistent with the global response pattern, suggesting that when Not1 is depleted a part of the observable increase in this cluster is due to accumulation emanating from the interference to the 5′-3′ degradation branch (Figure 5C,F, Supplementary Figure S5C, D).

These results demonstrate that mRNA generally accumulates in the same pattern when factors along the 5′-3′ degradation branch are perturbed, but the degree of accumulation depends on the specific element depleted (Not1 < Dcp2 < Xrn1).

Upstream perturbations in the 5′-3′ degradation pathway result in earlier onset of the transcription adaptation response

To understand the apparent association between the global mRNA accumulation profile and the 5′-3′ degradation pathway order, we excluded the Ccr4–Not sensitive genes from the analysis and turned to examine transcription along the depletion time-courses of these factors.

In all three cases, we observed a reduction in transcription, but strikingly, the timing order of the observed reduction recapitulated the observed order in the case of total mRNA, namely—Not1 caused the most immediate decline in recently-transcribed mRNA, followed by Dcp2, and then by Xrn1 (t1/2 of 33′, 52′ and 70′ respectively;
pairs and found a strong correlation (Figure 5H, then compared these measures between the different strain translation phase (changes in recently-transcribed mRNA during the accumulation) of the adaptation response when Not1 is depleted compared to Dcp2 depletion and in Dcp2 relative to Xrn1. To test this decreased faster in Dcp2 compared to Xrn1 should accumulate less mRNA in Dcp2 compared to Xrn1. To test this result suggested that the difference in accumulated mRNA between the strains is due to the earlier onset of the adaptation response when Not1 is depleted compared to Dcp2 depletion and in Dcp2 relative to Xrn1 depletion.

While the reduction in transcription affects the entire genome, there was still significant transcript-to-transcript variation in the exact dynamics (Figure 5G). We reasoned that if the earlier reduction in transcription explains the reduced degree of mRNA accumulation in Not1 relative to Dcp2 and relative to Xrn1, this relationship should also hold per transcript, namely, genes whose transcription is decreased faster in Dcp2 compared to Xrn1 should accumulate less mRNA in Dcp2 compared to Xrn1. To test this hypothesis, we calculated the difference between cumulative changes in recently-transcribed mRNA during the accumulation phase (Δnew), and the difference in maximal total mRNA during the accumulation and adaptation phase (Δmax mRNA, Figure 5H). We then compared these measures between the different strain pairs and found a strong correlation (Figure 5H, r = 0.4, P < 10^-70), i.e. genes with reduced transcription in Dcp2 depletion compared to Xrn1 depletion accumulated less mRNA, as expected. This was also the case (albeit to a lesser degree) when we examined the differences between Dcp2 and Not1 deletions (r = 0.31, P < 10^-98, Supplementary Figure S5H).

Taken together, these results point to a global mRNA accumulation pattern in response to a perturbation along the 5’-3’ degradation pathway. The degree of mRNA accumulation can be explained by the timing of the transcription response, and furthermore, the onset time of the response is earlier when upstream factors in the 5’-3’ pathway are perturbed.

Finally, we looked for sets of genes whose transcription decreased faster or slower than average and found a clear signature of G1- and S-phase genes (Figure 5I) (84). For further results and analyses regarding the cell cycle, see supplementary notes and Supplementary Figures S6 and S7).

**DISCUSSION**

We set out to study the mechanism of mRNA homeostasis that was observed in multiple conditions and organisms. The literature surrounding the question of feedback mechanisms between mRNA degradation and transcription is largely based on steady-state measurements. As a baseline, when we examined Xrn1 knockout we recapitulated a previous observation of a genome-wide reduction in degradation and transcription rates resulting in unchanged global mRNA (Figure 2). We reasoned that to study a feedback mechanism, steady-state measurements can be insufficient and potentially lead to incorrect interpretations. Therefore, we developed and applied a high-throughput sequencing-based implementation of the widely used cDTA technique (Figure 1). This allowed us to monitor total and
Figure 5. The transcription adaptation response is induced earlier when the 5'-3' pathway is perturbed upstream. (A) AID-tagged proteins in this figure. Nuclear factors to the left - Rat1 is the nuclear 5'-3' exoribonuclease, Nrd1, Sen1 and Nab3 survey aberrant transcripts and recruit the nuclear exosome (Dis3, Rrp6). Mature mRNA leaves the nucleus and will be deadenylated in the cytoplasm by the Ccr4–Not complex (Not1, Pop2, Ccr4). After deadenylation transcripts will continue to degrade 3'-5' by the cytosolic exosome (Dis3), or 5'-3' by Xrn1 after decapping by the DCP complex (Dcp2). Pab1 is the polyA binding protein, and Pan3 is part of an alternative deadenylation complex (37). (B) Factors noted in (A) were subjected to a 4-h depletion time course and cDTA-seq (as in Figure 4A). The matrix summarizes the Pearson correlation between the log-fold changes to transcripts relative to \( t = 0 \) after depletion of Xrn1 and Not1. Highlighted square corresponds to the scatterplot shown in (C). (C) Correlation between Xrn1 and Not1 depletion. Fold change relative to \( t = 0 \) after 90 min in Xrn1AID (x-axis) and Not1AID (y-axis). Changes are generally correlated. Set of Not1-sensitive transcripts are marked in blue, other points are colored by density. Data correspond to marked columns in (F) and marked square in (B). (D) Same as in (B), but comparing all time-course experiments. Highlighted square corresponds to the square depicted in (B). Note that the color scale is different. (E) Average changes to transcripts' mRNA relative to \( t = 0 \) (y-axis) along the time course (x-axis) for factors exhibiting significant correlation to Xrn1 (Supplementary Figure S5A, see Supplementary Figure S5B for the 240' timepoint). (F) mRNA changes upon interference to the 5'-3' cytosolic pathway are correlated. Changes to transcripts (rows) along the time-course (x-axis, 15, 30, 45, 60, 90, 120, 240 min) in three time-courses: upon depletion of Xrn1, Dcp2, and Not1. The mRNA log fold change relative to \( t = 0 \) is color-coded. Transcripts are split into Not1-sensitive (\( N = 672 \)), and the rest of the transcripts (\( N = 4539 \)). Rows in each set are sorted by the extreme point in a smoothed trajectory of the Xrn1 response. Highlighted columns correspond to the x- and the y-axis in (C). (G) The transcription adaptation response occurs earlier when the 5'-3' pathway is perturbed upstream. Each line represents the average change (over all transcripts) to recently-transcribed mRNA relative to \( t = 0 \) (y-axis) along the depletion time course (x-axis) in each one of the four strains. The transcript-change distribution in each strain at the 60' time point is detailed on the right, where the median of each distribution is denoted by a horizontal line. (H) Recently-transcribed mRNA differences explain total mRNA differences between strains. To compare the differences in response profiles between strains (in this case, comparing Xrn1 and Dcp2), we calculate the difference between the maximum observed change in mRNA per transcript (\( \Delta \text{max} \), y-axis in scatter) and between cumulative nascent trajectories (shaded gray area in example, \( \Delta \text{new} \), x-axis in scatter). We plot these statistics per transcript (dots in scatter, color denotes density) and found a significant correlation (Pearson \( r = 0.4 \), \( P < 10^{-17} \)). See Supplementary Figure S5H for the same comparison between Dcp2 and Not1 (Pearson \( r = 0.31 \), \( P < 10^{-98} \)). (I) A unique cell-cycle signature when the 5'-3' degradation pathway is perturbed. We compared the distributions of transcription changes (relative to \( t = 0 \)) in cell-cycle genes (rows) to the distributions of non-cycling genes (84). Significant deviations are denoted as colored triangles (purple/down—lower than non-cycling genes, orange/up—higher than non-cycling genes, size proportional to Kolmogorov–Smirnov \( q \)-value). Each triangle denotes the difference in a specific depletion time point (columns, x-axis time since auxin addition, same as in (B)). The bottom panels denote the average log fold change to mRNA and nascent mRNA in the same samples (same as in E and G). For further details, data, and analysis see supplementary material.
recently-transcribed mRNA in detailed dynamic settings for hundreds of samples, resulting in the biggest resource of metabolically-labeled mRNA in yeast to date. Applying cDTA-seq to cells undergoing rapid Xrn1 depletion (Figure 3A), we observed pronounced mRNA accumulation in virtually all transcripts, followed by a return to normal wildtype levels. The labeled mRNA profile provided by cDTA-seq revealed a striking reduction in transcription roughly 60 minutes following Xrn1 depletion (Figure 4). Finally, when we applied cDTA-seq to cells in which different RNA processing factors were depleted, we found that the response to Xrn1 depletion is not unique; detailed dynamic measurements revealed that the depletion of decapping and deadenylation factors results in a similar initial increase in mRNA levels. However, while the initial response was similar, the reduction in transcription occurred earlier when upstream factors along the 5'-3' degradation pathway were perturbed (Figure 5).

We focused on Xrn1 for two main reasons. First, it was implicated in mRNA homeostasis in two important but incongruent works by the Choder (19) and Cramer (18) groups. Secondly, Xrn1 degrades a large proportion of mRNA molecules in eukaryotic cells (6), so we expected a considerable response. Indeed, Xrn1 knockout exerts the most extreme alterations to mRNA profiles in published systemic knockout studies (Supplementary Figure S2A). Despite our technique being more akin to the cDTA protocol used by the Cramer group (Figure 1G), our data support the results from the Choder group, namely that mRNA levels are unchanged when cells lack Xrn1 and that degradation and transcription rates are significantly reduced (Figure 2).

To understand this homeostatic response, we applied cDTA-seq to cells undergoing rapid Xrn1 depletion (Figure 3A). We observed pronounced and transient mRNA accumulation immediately after the perturbation. To our knowledge, this is the first time that such a detailed view of a transient increase in mRNA levels is observed. Strikingly, in similar depletions of other factors (but not all, Figure 5E) cells revert to almost exactly the same mRNA levels they began with, suggesting some form of perfect adaptation taking place (85, 86). Further dynamic data in different settings will be useful to model this possibility. Importantly, this response does not seem to be a normalization artifact as it was measured in multiple different experimental batches and conditions (Figures 3, 5, and Supplementary Figure S7), and it was observed by single-molecule FISH in four different transcripts (Figure 3). The observed reversion to normal levels was harder to verify by smFISH, as cells lacking Xrn1 accumulate deadenylated mRNA molecules (8, 90) that also interact with the FISH probes. This makes the direct counting of poly(A+) molecules by FISH and microscopy challenging. As an alternative that circumvents direct molecule counting, we used polyA probes and FACS to measure the overall quantity of polyA RNA in cells and observed a temporal response consistent with the cDTA-seq signal (Supplementary Figure S3D).

Having excluded normalization, increased growth, or compensatory degradation as possible explanations for the reduction in mRNA levels, we could explain the return to wildtype levels by a global reduction in transcription roughly 60 min following Xrn1 depletion together with dilution by continued growth (Figure 4). The genome-wide nature of the reduction strongly supports a general mechanism for transcriptional regulation, rather than the existence of multiple gene-specific feedback loops. This result is further supported by a recent report that uses aneuploid cells to distinguish between these two possibilities (87).

Importantly, the observed delay in transcription reduction argues against the direct involvement of Xrn1 in this transcriptional reprogramming, as Xrn1 is degraded rapidly from cells (Figure 3A), but the transcriptional response occurs an hour later (Figure 4). Conversely, when we depleted Rat1 (the 5'-3' exonuclease operating in the nucleus that is involved in transcription termination (88)) we observed an immediate reduction in transcription (Supplementary Figure S6B). Alternatively, Xrn1 was hypothesized to indirectly affect global transcription through post-transcriptional regulation of Nrg1 (18). Therefore, we repeated the Xrn1 depletion time course experiment in Δnrg1 and Δnrg2 strains but we did not observe any difference in the total mRNA dynamics or in transcription (not shown), arguing against Nrg1/2 involvement in the feedback.

To further dissect the response, we expanded our experiments to include the depletion of dozens of RNA-related factors (shown in Figure 5A, and others, not shown). This expansive view revealed various response dynamics (Figure 5D, E) and will hopefully prove useful to understand different phenomena related to mRNA processing. Focusing on the adaptation response we observed in the wake of Xrn1 depletion, it was clear that Dcp2 and Not1 depletion elicited similar cellular reactions (Figure 5F). The observation that Xrn1 depletion caused the most significant accumulation of mRNA amongst all the factors we depleted, and that this accumulation is similar to the response upon Dcp2/Not1 depletion, argues that the 5'-3' pathway is the major pathway regulating mRNA levels in exponentially growing yeast in rich medium. Therefore, in our analysis, we put aside potential compensations by alternative degradation pathways (e.g. Not1-sensitive genes, Figure 5C, F), and focused on the 5'-3' degradation pathway.

Strikingly, when factors along the 5'-3' mRNA degradation pathway were depleted, the mRNA response began essentially the same, but seemed to be attenuated in the order of the factor along the pathway (Not1 < Dcp2 < Xrn1). Consistent with the reduced accumulation of mRNA along the 5'-3' pathway, we observed a delayed reduction in transcription, that also followed the order of the 5'-3' degradation pathway (Figure 5G), parsimoniously explaining the reduced accumulation of mRNA (Not1 < Dcp2 < Xrn1). We interpret these results to suggest that cells do not monitor accumulated deadenylated and decapped mRNA, or downstream byproducts such as p-bodies (89), for if this was the case, transcription inhibition would ensue earlier in the Xrn1 depletion time course (Figure 6). Extending the logic of this argument, and neglecting alternative degradation pathways, the agreement between the 5'-3' mRNA degradation pathway order and the onset time of the transcription adaptation response suggests that cells monitor a precursor that accumulates upstream of this pathway.

This model, when extrapolated, suggests local feedback along the lifecycle of the mRNA, rather than a closed-
A functional diversion of the Ccr4–Not complex itself due to mRNA accumulation and binding might cause the transcriptional response to the accumulation of mRNA may cause the transcriptional re-

A better understanding of the feedback mechanism employed by cells will likely have broader implications and might be relevant for critical processes in health and disease such as proliferation, apoptosis, and viral immune response (101). The data we presented here argue that detailed dynamic measurements are important for a deeper understanding of global mRNA regulation.

**DATA AVAILABILITY**

Sequencing data are available under GEO accession number GSE196937. Scripts and summary data for generating all the figures and supplementary figures in this manuscript are available at 10.5281/zenodo.6528287. Raw microscopy images are available upon request.
SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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