A Transcriptome-wide Atlas of RNP Composition Reveals Diverse Classes of mRNAs and lncRNAs

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SUMMARY

Eukaryotic genomes generate a heterogeneous ensemble of mRNAs and long noncoding RNAs (lncRNAs). LncRNAs and mRNAs are both transcribed by Pol II and acquire 5’ caps and poly(A) tails, but only mRNAs are translated into proteins. To address how these classes are distinguished, we identified the transcriptome-wide targets of 13 RNA processing, export, and turnover factors in budding yeast. Comparing the maturation pathways of mRNAs and lncRNAs revealed that transcript fate is largely determined during 3’ end formation. Most lncRNAs are targeted for nuclear RNA surveillance, but a subset with 3’ cleavage and polyadenylation features resembling the mRNA consensus can be exported to the cytoplasm. The Hrp1 and Nab2 proteins act at this decision point, with dual roles in mRNA cleavage/polyadenylation and lncRNA surveillance. Our data also reveal the dynamic and heterogeneous nature of mRNA maturation, and highlight a subset of “lncRNA-like” mRNAs regulated by the nuclear surveillance machinery.

INTRODUCTION

High-throughput transcriptome analyses in eukaryotes have revealed pervasive transcription at most, if not all, genomic loci. This generates many long noncoding RNAs (lncRNAs), which lack protein-coding capacity and are distinct from well-characterized structural RNAs (rRNAs, tRNAs, snRNAs, and snoRNAs) or small regulatory RNAs. LncRNAs arise from intergenic, antisense, or promoter-proximal regions and range in size from ~200 nt to >20 kb. Many features are shared between lncRNAs and mRNAs; both classes of RNA possess 5’-methylguanosine caps (Neil et al., 2009) and poly(A) tails (David et al., 2006) and have broadly similar lengths. Moreover, both are transcribed by RNA polymerase II (Pol II) from similar preinitiation complex assemblies (Rhee and Pugh, 2012) and can be regulated by common transcription factors. Despite these similarities, the fates and functions of lncRNAs and mRNAs are substantially different. Most mRNAs are rapidly exported to the cytoplasm, where they engage with the protein synthesis machinery. In contrast, diverse nuclear functions have been attributed to lncRNAs, including the assembly of nuclear domains, directing chromatin-modification, resetting of epigenetic marks, and the regulation of mRNA transcription.

Several studies have identified classes of lncRNA with distinct features. For example, stable unannotated transcripts (SUTs) are detectable in wild-type yeast, whereas cryptic unstable transcripts (CUTs) are apparent only in the absence of the nuclear surveillance factor Rrp6, and Xrn1-sensitive unstable transcripts (XUTs) are apparent only in the absence of the cytoplasmic exoribonuclease Xrn1 (van Dijk et al., 2011; Xu et al., 2009). This indicated that distinct classes of lncRNAs can be distinguished from each other, as well as from mRNAs, but the features that might differentiate these species were unclear.

All mRNAs interact with a defined series of protein factors during their transcription, packaging, processing, export, and turnover (see Figure 1A), forming ribonucleoprotein particles (RNPs). We hypothesized that lncRNAs and mRNAs must diverge at some point along this maturation pathway. We therefore systematically analyzed the in vivo, transcriptome-wide targets of key factors in this pathway in budding yeast (Figure 1A). We anticipated that this atlas of RNP compositions would provide a comprehensive picture of the dynamic events during canonical messenger RNP (mRNP) assembly, and give insights into the definition and behavior of different classes of mRNAs and lncRNAs.

Overall, our data reveal how distinct transcript classes are defined and how RNP composition relates to function, and have enabled us to begin to tackle the overwhelming complexity of the transcriptome.

RESULTS

Transcriptome-wide Analysis of RNP Composition

To establish how and when different classes of Pol II transcripts are distinguished in the cell, we determined the transcriptome-wide targets for 13 key mRNA biogenesis and turnover factors (listed in Figure 1A and Table 1, with references therein). We included nuclear surveillance factors (Mtr4 and Trf4), a component of the nuclear cap-binding complex (Cbc1), components of the TREX RNA packaging complex (Gbp2 and Tho2),...
Figure 1. Transcriptome-wide Analysis of RNP Composition

(A) mRNA maturation and decay factors selected for analysis.

(B) Relative recovery of spliced mRNAs versus unspliced pre-mRNAs bound to the tested proteins, expressed as the ratio of RNA fragments spanning exon-intron-exon junctions.

(C) Average binding distribution of the tested proteins across mRNAs. For each protein, average hit densities were calculated for 120 bins spanning their 1,000 most abundantly bound mRNAs (including 2×10 bins for 100 nt 5′ and 3′ flanking regions). 5′-proximal hits can arise from binding to promoter-proximal ncRNAs or to the 5′ end of full-length mRNAs.

(D–I) Total hits for each protein in RPL3 and HTB2 pre-mRNAs, snoRNAs, CUTs, SUTs, and XUTs, as a percentage of all hits in Pol II transcribed RNAs (mRNAs, CUTs, SUTs, snRNAs, and snoRNAs; Table S1).

(J–M) Hit distributions along individual transcripts, at the indicated scales (hits per million hits in Pol II transcribed RNAs). Note the different scale used for Hrp1 data in K and M due to the high level of binding. See also Figure S1 and Table S1.
pre-mRNA 3’ cleavage and polyadenylation factors (Hrp1, Nab2, and Pab1), a nuclear-cytoplasmic export factor (Mex67), an mRNA localization factor (Hek2), a cytoplasmic translation factor (Tif1), and cytoplasmic mRNA turnover and surveillance factors (Ski2 and Xrn1). Hrp1 and Nab2 function in mRNA cleavage and polyadenylation but additional roles are reported for Nab2 in mRNA packaging (Batisse et al., 2009), export (Iglesias et al., 2010), and nuclear surveillance (Schmid et al., 2012), and for Hrp1 in nuclear and cytoplasmic surveillance (González et al., 2000; Kuehner and Brow, 2008).

For target site identification, we used the crosslinking and analysis of cDNA (CRAC) technique (Granneman et al., 2011). Actively growing cells expressing HTP-tagged (His6-TEV-Protein A) proteins under the control of the endogenous promoter were UV irradiated to fix direct protein:RNA contacts. After stringent, multi-step affinity purification, mild RNase digestion, and radiolabelling, RNPs were isolated by SDS-PAGE (Figure S1A available online). Bound RNA fragments were amplified by RT-PCR and analyzed by high-throughput sequencing. Identical conditions were used for all proteins tested, and in most cases replicate data sets acquired (Table S1). We also repeated the analysis for the poly(A)-binding protein Nab2 in an rrp6Δ background, which is reported to stabilize its transient binding (Schmid et al., 2012). Comparison of the number of reads mapping to each annotated transcript in replicate data sets revealed good reproducibility, with most Spearman rank correlation coefficients $r \geq 0.75$ (Figure S1B). Furthermore, the most highly enriched 10% of mRNAs in Hrp1, Nab2, and Hek2 data sets showed significant overlap with published immunoprecipitation analyses ($\chi^2 \leq 0.001$) (Batisse et al., 2009; Hasegawa et al., 2008; Kim Guisbert et al., 2005).

A breakdown of hits by transcript class revealed a broad range of substrate specificities for the tested proteins (Figure S1C). The poly(A)-binding protein Pab1 predominantly bound to mRNAs, whereas tRNAs were prevalent targets of the nuclear surveillance factor Mtr4, consistent with reports of extensive nuclear pre-tRNA degradation (Gudipati et al., 2012b). In Hrp1 data sets, snoRNAs were abundant, supporting a role in snoRNA biogenesis originally indicated by a ChIP study (Kim et al., 2006). Many proteins crosslinked to rRNAs at regions distinct from those typically detected as background (Figure S1D). Mex67 is reported to bind 60S and 40S (Faza et al., 2012) preribosomal particles, in good agreement with our data, and the peak of Xrn1 binding in ITS1 supports its role in cytoplasmic degradation of this excised spacer region. The nuclear surveillance factors Mtr4 and Trf4 assist in the degradation of the 5’ ETS, and we identified binding within this region. The specific rRNA association of the cytoplasmic helicase Ski2 is likely to

### Table 1. Proteins Selected for Analysis

| Yeast Protein | Human Homolog | Function | References |
|--------------|---------------|----------|------------|
| **Nuclear Surveillance** |
| Mtr4 | hMtr4/SKIV2L2 | RNA helicase (Mtr4) and noncanonical poly(A) polymerase (Trf4) within the TRAMP complex; assist the nuclear exosome in RNA degradation | Reviewed in Porrua and Libri, 2013 |
| Trf4 | hTRF4-1/POLS | Cleavage and polyadenylation factor (Hrp1) | Kessler et al., 1997 |
| **Early mRNA Biogenesis** |
| Cbc1/Sto1 | CBP80 | Nuclear cap binding complex subunit; mRNA stabilization, processing, export and decay | Göremann et al., 2005; Wong et al., 2007 |
| Tho2 | Thoc2 | Components of the TREX complex; transcription elongation and mRNA export | Reviewed in Rondón et al., 2010 |
| Gbp2 | | | |
| **Cleavage and Polyadenylation** |
| Hrp1 | TDP-43 | Cleavage and polyadenylation factor | Kessler et al., 1997 |
| Nab2 | ZC3H14 | Nuclear poly(A)-binding protein; poly(A) tail length control, mRNA export and nuclear surveillance of pre-mRNAs | Iglesias et al., 2010; Schmid et al., 2012; Viphakone et al., 2008 |
| Pab1 | PABPC1 | Poly(A)-binding protein; mRNA export, translation and stability | Reviewed in Parker, 2012 |
| **Export and Translation** |
| Mex67 | NXF1/Tap | mRNA export receptor | Hieronymus and Silver, 2003 |
| Hek2/Khd1 | hnRNP K, hnRNP E and poly(C)-binding proteins | mRNA localization, translational inhibition and stability | Hasegawa et al., 2008; Irie et al., 2002; Mauchi et al., 2010; Paquin and Chartrand, 2008; Vogel et al., 2011; Wolf et al., 2010 |
| Tif1/eIF4A | eIF4A | Helicase within the cytoplasmic cap-binding complex; ribosome scanning of the 5’ UTR | |
| **Cytoplasmic Decay** |
| Ski2 | SKIV2L | Helicase that assists the cytoplasmic exosome in 3’–5’ mRNA turnover | Reviewed in Parker, 2012 |
| Xrn1 | XRN1 | 5’ to 3’ exonuclease in the major cytoplasmic mRNA decay pathway | Reviewed in Parker, 2012 |
reflect interactions with translating ribosomes during mRNA degradation by the exosome. Other proteins showed distributed binding on the pre-rRNA (Figure S1D), but the significance is currently unclear.

Assembly and Architecture of mRNPs

Analyses of hits in mRNAs provided a high-resolution picture of the dynamic assembly of mRNPs. For each protein, we calculated the ratio of reads mapping across exon-exon (EE) versus intron-exon (IE) junctions (Figure 1B) (Schneider et al., 2012). IE junctions are exclusively present in unspliced pre-mRNA, and EE junctions in spliced mature mRNAs, so the EE/IE score indicates when, relative to splicing, each protein associates with the mRNP. Low scores were found for the nuclear cap-binding protein Cbc1, TIF1, and Gbp2, and nuclear surveillance factors Mtr4 and Trf4, consistent with these factors acting during or shortly after transcription. In contrast, high scores for the translation initiation helicase Tif1 and cytoplasmic surveillance factors Xrn1 and Ski2 indicate that they function late in the mRNP lifecycle. The slightly lower score for Xrn1 is consistent with its role in surveillance of unspliced pre-mRNAs and lariat intermediates (Hilleren and Parker, 2003).

The EE/IE analysis was most informative for shuttling proteins. For example, Mex67 and Nab2 load onto mRNA in the nucleus and contribute to export (Iglesias et al., 2010) and are removed at the cytosolic face of the nuclear pore (Lund and Guthrie, 2005; Tran et al., 2007). The lower EE/IE scores for Nab2 suggest that Nab2 enters the mRNP before Mex67, which is perhaps only recruited when cleavage and polyadenylation is complete. Nab2 and Pab1 are both implicated in poly(A) tail length control, but it has been unclear which acts first. The EE/IE score for Pab1 is lower than that of Mex67 and similar to that of Nab2, supporting early roles for both Nab2 and Pab1 in the nucleus. The cleavage factor Hrp1 also shuttles and can contribute to cytoplasmic surveillance (González et al., 2000). The low EE/IE score, however, suggests that Hrp1 primarily functions early in mRNP biogenesis, consistent with prolonged cytoplasmic binding of Hrp1 denoting an aberrant mRNP. Conversely, the high score for Hek2, which contributes to translational regulation and mRNA localization (Irie et al., 2002; Paquin and Chartrand, 2008), suggests that it binds late in the nucleus and is predominantly associated with cytoplasmic mRNPs.

Although the architecture of several ribonucleoprotein complexes, such as the ribosome and spliceosome, has been studied in detail, little is known about the topology of mRNPs. We therefore examined their organization by plotting the distribution of reads mapping to snoRNAs, CUTs, and SUTs (Figures 1J–1L), Cbc1 and Tif1 reside in complexes that interact with the 5'-proximal RNA fragments rather than to the 5' ends of full-length mRNAs (see below). Nab2 binding was observed throughout the body of mRNAs, consistent with previous ChIP analyses (González-Aguilera et al., 2011). In addition to binding poly(A), Nab2 shows nonspecific RNA-binding activity (Viphakone et al., 2008) and may be an architectural component of mRNPs (Batisse et al., 2009).

In the cytoplasm, the 5' to 3' and 3' to 5' mRNA decay pathways are preceded by deadenylation of the poly(A) tail to ~10–12 nt. Pab1 is then displaced and the 3' end becomes accessible to the Ski2/3/8 complex and exosome for 3' degradation, or the Lsm1-7/Pat1 complex that activates decapping and 5' degradation by Xrn1. We observed prominent peaks at the 3' end of mRNAs for Ski2 and, less expectedly, for the 5' to 3' exonuclease Xrn1. This indicates that the oligo(A) tail is the site of a rate-limiting step in mRNA turnover, perhaps reflecting assembly of the surveillance machinery or regulated initiation of decay. Indeed, Xrn1 interacts with the Lsm1-7/Pat1 complex that crosslinks primarily to mRNA 3' ends (Mitchell et al., 2013). The 3' peak of Xrn1 hits might also reflect slowed degradation of the 3'UTR due to the presence of RNA-binding proteins not displaced by translating ribosomes. The absence of clear peaks of Xrn1 crosslinking elsewhere along the mRNA body is consistent with its high processivity, perhaps following the last translating ribosome (Hu et al., 2009). Supporting this model, we observed a moderate accumulation of Xrn1 upstream of mRNA stop codons (Figure S2A). Conversely, Ski2 binding was distributed across the body of mRNAs, suggesting that 3'-5' decay is slower or is more prone to pausing, perhaps due to collisions with translating ribosomes.

RNP Composition Defines Distinct Transcript Classes

Having obtained a picture of an “average” mRNP, we next investigated how RNP composition varies between mRNAs and between classes of Pol II transcripts. For each protein tested, we extracted all hits in Pol II transcribed RNAs (Table S1) and plotted the proportion mapping to snoRNAs, CUTs, SUTs, and two mRNAs. Combining the data for all 13 proteins produced “RNP profiles” for these six transcript types (Figures 1D–1I). Cbc1, Gbp2, and Tho2 were moderately abundant in all six RNP profiles and bind with similar distributions along mRNAs, CUTs, and SUTs (Figures 1M and S2B), suggesting they are universal RNP components and that early RNP assembly is similar for mRNAs and IncRNAs. These data corroborate reports that the CBC is present in snoRNP assembly intermediates (Schwer et al., 2011), and that the THO complex regulates snoRNA expression by binding at the 3' end (Figure S2C) (Larochelle...
et al., 2012). Hrp1 and Nab2 have multiple functions in RNA metabolism and were present in snoRNPs, IncRNPs, and the HTB2 mRNP, indicating that they too are ubiquitous constituents of RNPs.

Despite these similarities in early RNP assembly, there were striking differences in binding of different transcript classes to cytoplasmic and nuclear surveillance factors. RPL3 was strongly bound by Xrn1 and Ski2, but not Mtr4 or Trf4, indicating that it is predominantly degraded in the cytoplasm, whereas the reverse was seen for snoRNAs, SUTs, and particularly CUTs. Furthermore, RPL3 and HTB2 were bound more extensively by the export receptor Mex67 than were snoRNAs, CUTs, and SUTs. Collectively, these data suggest that CUTs and SUTs, like snoRNAs, are predominantly confined to the nucleus and that the distinction between IncRNPs and mRNPs occurs after early RNP packaging but prior to Mex67 recruitment. Notably, CUTs and SUTs were significantly bound by Pab1, Hrp1, and Nab2 suggesting that 3’ end formation on lncRNAs initially resembles that of mRNAs but culminates in nuclear retention rather than export.

In addition to differences in RNP composition between transcript classes, we identified heterogeneity within each class. For example, Trf4 bound to HTB2 more strongly than RPL3, consistent with reports that Trf4 regulates HTB2 expression (Reis and Campbell, 2007). This heterogeneity raised the question of whether the scarce IncRNA hits in Xrn1, Ski2, and Mex67 data sets arise from a general low level of binding to IncRNAs or robust interactions with a few atypical lncRNAs. To assess heterogeneity among mRNPs and IncRNPs, we performed a k-medians clustering analysis of mRNAs, CUTs, and SUTs based upon their individual RNP profiles (Figure 2A). These profiles were derived from the number of hits for each transcript in Cbc1, Mtr4, Nab2, Mex67, Xrn1, and Ski2 data sets (Figure 2A, columns 3–8), with hits normalized for each row (transcript). Within the Pab1 data set, some transcripts with relatively low numbers of total hits displayed a sharp peak of binding at the 3’ end. To distinguish
site-specific binding at these putative polyadenylation sites, from broadly distributed, potentially non-specific, interactions across transcript bodies, we generated a “peak sharpness” score (Figure 2A, column 9). For this, we identified the highest peak in each transcript and divided the value of this by the maximum obtained when reads in the surrounding 400 nt region were randomly placed.

The cluster analysis recapitulated the major findings from Figure 1, with CUT and SUT classes of IncRNAs predominantly falling into clusters (I–III) distinct from mRNAs (IV–X). Furthermore, whereas cluster I was enriched for CUTs, clusters II and III contained more SUTs (Table S2), revealing differences not readily apparent from the class-wide analysis in Figure 1. In comparison to the “CUT cluster” (I), the “SUT clusters” (II and III) had more specific Pab1 binding. Cluster III also has less dominant binding to the nuclear surveillance factor Mtr4. This suggests that although all IncRNAs are predominantly retained and degraded in the nucleus, CUTs are more rapidly degraded so might represent byproducts of functional transcription, whereas SUTs are more stable so perhaps function as transcripts. Furthermore, 34.4% of SUTs fell into mRNA clusters (IV–X), compared to just 6% of CUTs, indicating that CUTs are distinct from mRNAs but SUTs and mRNAs often overlap.

The PHO84-as IncRNA, which falls into an mRNA cluster, functions in trans (Camblong et al., 2009) and accumulates in the cytoplasm (Castelnuovo et al., 2013). Furthermore, “mRNA-like” SUTs (clusters IV–X) bound more strongly to Mex67 than SUTs in the IncRNA clusters (I–III) (Figure S2D). The “mRNA-like” IncRNAs therefore appear to behave and function differently from the “standard” IncRNAs in clusters I–III, perhaps representing functional transcripts exported to the cytoplasm. Indeed, inspection of published transcriptome profiling data revealed that cluster IV–X SUTs accumulate less than cluster I–III SUTs in nuclear surveillance mutants (Figure S2E) (Gudipati et al., 2012b), suggesting they are degraded in the cytoplasm by Xrn1, with which they interact (Figure 2A, column 7). Hek2 has a role in mRNA stabilization (Mauchi et al., 2010; Vogel et al., 2011) and localization, and SUTs in clusters IV–X bound more abundantly to Hek2 than those in clusters I–III (Figure S2D). Hek2 might therefore be one factor that helps discriminate mRNAs and “mRNA-like” IncRNAs from “standard” IncRNAs, based on sequence-specific binding. Indeed, SUTs in clusters VI–X have particularly long CNN repeats (Figure S2F) to which Hek2 binds (Figure S2G).

Clusters I–III contained 411 mRNAs (Table S2), and these are likely to behave like IncRNAs, with retention and/or degradation in the nucleus. Rp6 is active in CUT surveillance (Neil et al., 2009; Xu et al., 2009), and cluster I–III mRNAs are highly stabilized in strains lacking this exonuclease (Figure S2D). Furthermore, NRD1 (cluster I) transcripts undergo attenuation dependent on the Nrd1–Nab3 complex (Kuehner and Brow, 2008), which functions in the termination and nuclear surveillance of many CUTs. Additional cluster I–III mRNAs, such as CTH2 (Clais et al., 2008) and RPL9B (Gudipati et al., 2012a) are also terminated and processed/degraded via Nrd1-dependent pathways. Other cluster I–III mRNAs including UR8 and IMD2 are regulated by promoter-proximal CUTs (Kuehner and Brow, 2008; Thiebaut et al., 2008). We predict that the remaining cluster I–III mRNAs either behave like IncRNAs, with retention and degradation in the nucleus, or are regulated by overlapping IncRNAs. Indeed, comparison to ribosome profiling data (Brar et al., 2012) (Figure S2H) reveals that cluster I–III mRNAs are ~5-fold less abundant on ribosomes than cluster IV–X mRNAs, relative to their transcription rate (gauged by Gfp2 binding; Figure S2H).

We noted significant heterogeneity between the mRNA clusters (IV–X). Cluster IV was most similar to the IncRNA clusters (I–III), with binding to Mtr4 as well as cytoplasmic surveillance factors. Several cluster IV mRNAs overlap CUTs (UR2, SER3, ADE12, IMD3, and LEU4) (Davis and Ares, 2006; Thiebaut et al., 2008), and we suggest that the “mixed” RNP profile of cluster IV reflects genes where mRNAs and IncRNAs are transcribed concurrently (perhaps in distinct subpopulations of cells). GO term analyses (Table S3) revealed that some mRNA clusters were enriched for transcripts related to particular cellular processes, suggesting that mRNP composition is linked to the function of the encoded protein. For example, transcripts encoding ribosomal proteins were prevalent in cluster V, which showed high binding by Nab2 (Figure 2A) and strong stabilization in Rpr44 mutants (Figure S2D) (Gudipati et al., 2012b). This is consistent with reports that Nab2 and Rrp44 act in the nuclear surveillance of ribosomal protein gene pre-mRNAs (Bousquet-Antonelli et al., 2000; Gudipati et al., 2012b; Schmid et al., 2012). Transcripts encoding proteins with functions critical to the nucleus, such as nucleic acid metabolism, were enriched in clusters VI and VII, with high Obc1 and Nab2 binding, whereas those encoding proteins participating in predominantly cytoplasmic processes such as glucose metabolism were enriched in cluster X, with high levels of Ski2, Xrn1, and Mex67 binding. This suggests that mRNAs are preferentially regulated in the cellular compartment most appropriate to the function of their encoded protein, perhaps facilitating rapid feedback regulation.

### RNA Classes Are Defined by Distinct Modes of 3’ End Formation

The largely distinct behavior and RNP composition of mRNAs, CUTs, and SUTs lead us to question how they are distinguished in the cell. Both CUTs and SUTs bound early, cotranscriptionally recruited, mRNP packaging components (Cbc1, TREX, Hrp1, and Nab2) but were underrepresented (particularly CUTs) among Mex67 targets. These results indicated that the distinction between CUTs, SUTs, and mRNAs is made following transcription elongation but prior to the acquisition of export competence. This suggested that there might be crucial differences in 3’ end formation, which generally proceeds via one of two possible mechanisms: stable mRNA 3’ ends are generated via cotranscriptional cleavage and polyadenylation, whereas the 3’ ends of some CUTs arise directly from Nrd1-dependent transcription termination coupled to oligoadenylation and turnover. Formation of the 3’ ends of SUTs has not been studied in detail.

Plotting the average distribution of Pab1 hits across mRNAs, CUTs, and SUTs (Figure 3A, red) revealed 3’ peaks for mRNAs and SUTs but distributed binding across CUTs, consistent with the lower Pab1 peak scores in the CUT cluster (Figure 2, cluster I). Thus 3’ end processing only of mRNAs and SUTs results in the acquisition of a stable, Pab1-bound poly(A) tail.
Together with the clustering analysis in which SUTs and mRNAs overlapped and were distinct from CUTs, this leads to a working model in which SUTs undergo cleavage and polyadenylation like mRNAs. In contrast, CUTs are terminated by a distinct mechanism, most likely dependent on Nrd1-Nab3, coupled to rapid turnover.

Cleavage and polyadenylation of pre-mRNAs is dependent on the recognition of a precise configuration of sequence motifs by a large multicomponent complex. To investigate whether SUT 3' ends are defined by similar motifs, we needed to precisely localize sites of 3' end formation on SUTs and mRNAs. We therefore searched the Pab1 data set for chimeric reads in which the mapped sequence is followed by adenosine residues that are not genome encoded (nonencoded A-tails). These represent junctions between mRNA 3' ends and poly(A) tails, and extracting the genomic coordinate of the last encoded nucleotide provided us with a transcriptome-wide set of precise poly(A) (pA) sites. Plotting these sites across mRNAs and SUTs gave sharp 3' peaks (Figure 3A, blue), indicating that 3' ends were detected with high precision. For many transcripts, we found multiple pA sites, consistent with a recent transcript isoform sequencing study (Pelechano et al., 2013) and indicating that alternative pA sites are prevalent. To compare the sequence features defining mRNA and SUT 3' ends, we generated logos for the genomic sequence flanking the most frequently identified pA site for each mRNA (n = 5,453) or SUT (n = 370). (B–F) Analysis of the set of genomic sequences flanking the most frequent pA site for each mRNA (n = 5,453) or SUT (n = 370). (B) Sequence logo of these regions. (C) Sequence logo of the most highly enriched motif in these regions. (D and E) Occurrence of UAUAUA and AAUAAA motifs. (F) Nucleotide frequency distributions.

Figure 3. RNA Classes Are Defined by Alternative Modes of 3' End Formation

(A) Red: average distribution of Pab1 hits across mRNAs, SUTs, and CUTs (300 nt flanks included). Blue: average distribution of poly(A) (pA) sites, defined as the last genome-encoded nucleotide within Pab1-bound RNA fragments with a non-genome-encoded poly(A) tail.

(B–F) Analysis of the set of genomic sequences flanking the most frequent pA site for each mRNA (n = 5,453) or SUT (n = 370). (B) Sequence logo of these regions. (C) Sequence logo of the most highly enriched motif in these regions. (D and E) Occurrence of UAUAUA and AAUAAA motifs. (F) Nucleotide frequency distributions.
UAUAUA as highly enriched in the 3' regions of SUT and mRNA genes, most frequently located ~30–70 nt upstream of the pA site (Figures 3C and 3D). This motif corresponds to the efficiency element, originally identified ~50 nt upstream of mRNA cleavage sites. Messenger RNA 3' ends are also defined by the positioning element, AUAAAA, 10–30 nt upstream of the pA site, and U-rich regions flanking the pA site. We detected both of these features in genes encoding SUTs and mRNAs when we plotted the frequency of AUAAAA motifs (Figure 3E) or nucleotide base composition (Figure 3F) around pA sites. We conclude that mRNAs and SUTs possess stable Pab1-bound poly(A) tails, and their 3’ ends are defined by common sequence elements.

mRNA Cleavage and Polyadenylation Factors Participate in CUT Surveillance

The mode of 3’ end formation of mRNAs and SUTs is apparently distinct from CUTs, so it was surprising that CUTs bound to the mRNA cleavage and polyadenylation factors Hrp1 and Nab2. Hrp1 specifically binds the UAUAUA efficiency element via its tandem RRM domains (Pérez-Canadillas, 2006). This motif was enriched at Hrp1-binding sites in mRNAs but also within SUTs and CUTs (Figure 4A). Single-nucleotide deletions in cDNA reads indicate the precise nucleotide crosslinked to the bait protein, and for Hrp1 hit density and sequence deletions were elevated over UAUAUA motifs in all three classes of transcript indicating that this is a direct binding site (Figure 4B). However, whereas mRNAs and SUTs displayed a 3’ peak of Hrp1 binding, CUTs did not (Figure 4C). Furthermore, ~40% of Hrp1-binding sites at the 3’ end of mRNAs and SUTs possessed a UAUAUA element, but there was no such enrichment among the low number of Hrp1-binding sites at the 3’ end of CUTs (Figure 4D). This indicates that Hrp1 binds directly to the efficiency element to promote 3’ processing of mRNAs and SUTs, but binds in a more distributed manner throughout CUTs. Notably, even within SUTs and mRNAs, the majority of Hrp1 is bound to promoter-proximal regions (Figure 4C), and ~90% of binding sites lack a UAUAUA motif (Figure 4A). This suggested that Hrp1 has additional functions unrelated to cleavage and polyadenylation and independent of binding to UAUAUA.

The poly(A)-binding protein Nab2 also participates in mRNA 3' end formation and showed increased binding at the 3’ ends of mRNAs and SUTs. However, like Hrp1, Nab2 binding was more distributed over CUTs, with additional binding toward the 5’ ends of mRNAs and SUTs (Figure 4C). The lack of binding to the 3’ end of CUTs was not simply due to their rapid turnover, as a 3’ peak of Nab2 binding was still absent from full-length CUTs stabilized in an rnp6J strain (Figure 4C). We conclude that both Hrp1 and Nab2 function in mRNA and SUT 3’ end formation but act in a different manner on CUTs and promoter-proximal regions.

To compare the functions of Nab2 and Hrp1 when bound to mRNAs and CUTs, we constructed strains with endogenous NAB2 and HRP1 genes under the control of glucose-repressible PGAL promoters. The effects of Hrp1 and Nab2 depletion were assessed for transcripts identified as targets in the CRAC analyses. Northern analysis of CCW12, which falls into a typical mRNA cluster (X), revealed a slight increase following Nab2 depletion and dramatic decrease upon Hrp1 depletion (Figure 4E). This is consistent with the essential role of Hrp1 in mRNA synthesis and with the mild general increase in mRNA expression observed in a previous analysis of Nab2 depletion (Schmid et al., 2012). In contrast, the abundance of two tested CUTs, CUT479 and CUT200, was increased up to 15-fold following depletion of Nab2 or Hrp1 (Figure 4E). Many of the RNA fragments isolated with Hrp1 or Nab2, including those mapping to CUTs, possessed short nonencoded oligo(A) tails (Figures 5A–5C). These are hallmarks of nuclear decay intermediates, suggesting that CUTs bound by Hrp1 and Nab2 were undergoing active degradation.

Previous analyses showed that Nab2 binds the surveillance factors Rpr6 and Trf4 (Schmid et al., 2012) and participates in degradation of intron-containing pre-mRNAs, whereas Hrp1 was implicated in Nrd1-dependent termination coupled to pre-mRNA turnover at the NRD1 and HRP1 loci (Kuehner and Brow, 2008). We conclude that Hrp1 and Nab2 participate in the nuclear turnover of CUTs, in addition to their roles in the generation of stable mRNAs and perhaps SUTs. These dual roles support a model in which 3’ end processing is a key step in determining transcript fate.

Early Termination Generates Promoter-Proximal IncRNAs

Finally, we sought to determine the origin of the 5’ proximal binding of surveillance factors to mRNAs. RNAs carrying non-encoded A-tails were identified for many proteins (Figure 5A) and indicate that transcripts have been released from the poly-merase. Generally, stable mRNAs possess long Pab1-bound poly(A) tails that promote export and translation but are deadenylated to ~10–12 adenosines prior to cytoplasmic turnover by Xrn1 or the Ski complex and exosome. In contrast, short (4–5 nt) oligo(A) tails mapping throughout a gene arise from the adenylylation activity of the TRAMP complex and characterize nuclear surveillance intermediates. In agreement with these roles, (1) A-tails in Xrn1 and Ski2 data sets were ~1–12 nt long (Figures 5B and 5C), absent from transcript classes such as tRNAs (Figure 5D), and almost exclusively present at the 3’ end of mRNAs (Figure 5E); (2) Pab1 bound to long poly(A) tails (present on 74.2% of recovered fragments) but not oligo(A)~10 (Figures 5B and 5C); and (3) Mtr4 substrates universally possessed short (4–5 nt) oligo(A) tails (Figures 5B–5D), some of which mapped across mRNAs (Figure 5E). Therefore, oligo(A) tails are a universal feature of TRAMP activity and are exclusively associated with nuclear surveillance, whereas the longer A-tails in Xrn1 and Ski2 data sets reflect 3’ poly(A) tails on mRNAs and some other Pol II transcripts. The abundant short oligo(A) tails in Nab2 data sets (Figure 5B) support a noncanonical role in surveillance.

Most oligo(A)-tailed mRNA reads in Mtr4 data sets mapped to promoter-proximal regions (Figure 5E) and even for long genes did not extend beyond the first ~400 nt (Figure 5F). Promoter-proximal Mtr4-bound fragments are therefore unlikely to represent decay intermediates of full-length mRNAs but might instead arise from high levels of early transcription termination. In Nab2 data sets, A-tails were shorter for promoter-proximal reads than for reads mapping to mRNA 3’ ends and largely absent from reads mapping to central regions of mRNAs (Figure 5G). Nab2 therefore binds to poly(A) tails on full-length mRNAs, and...
Figure 4. Dual Roles of Hrp1 and Nab2 in mRNA Cleavage/Polyadenylation and in CUT Surveillance

(A) Enrichment scores for 6 nt motifs in Hrp1-bound RNA fragments. Inset: proportion of Hrp1-binding sites containing the UAUAUA motif in mRNAs, CUTs, and SUTs.

(B) Distribution of Hrp1 hits (black) and deletions (red) around UAUAUA motifs in mRNAs, CUTs, and SUTs.

(C) Heatmap showing the enrichment of Hrp1 and Nab2 binding sites around the UAUAUA motif in mRNAs, CUTs, and SUTs.

(D) Proportion of Hrp1 binding sites containing UAUAUA motif.

(E) Western blot analysis of CCW12 and 25S rRNA for different strains and treatments.

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to oligo(A)-tailed unstable transcripts from promoter-proximal regions, whereas interactions with mRNA central regions apparently reflect an adenosine-independent-binding activity. Promoter-proximal transcripts of a similar length were previously identified in yeast depleted of Rrp6 and Trf4, which stabilizes full-length CUTs (Figure 5H) (Neil et al., 2009). This supports the notion that promoter-proximal transcripts do not originate from longer precursors.

To test whether promoter-proximal fragments arise from early termination, we compared the distribution of Pol II (Churchman and Weissman, 2011), the termination factors Nrd1 and Nab3 (Figure 5H) (Wlotzka et al., 2011), and the 5′ exonuclease Rat1 (Granneman et al., 2011) across mRNAs. The promoter-proximal enrichment of Pol II has been interpreted as stalled elongation complexes that are competent to resume transcription. However, Nrd1, Nab3, and Rat1 are all enriched in this region, which coincides with the promoter-proximal oligoadenylated fragments. This suggests that stalled Pol II is susceptible to early termination, triggered either by Nrd1-Nab3 or by cleavage/decapping followed by a Rat1-dependent “torpedo” mechanism. The resultant 5′ fragments account for most Mtr4, Trf4, Nab2, and Hrp1 hits mapping to protein-coding genes, thus removal of early termination products is a major function of the nuclear surveillance machinery. Furthermore, these transcripts bind the same factors as CUTs (Mtr4, Trf4, Hrp1, Nab2), indicating that protein-coding loci give rise to both classical mRNP and an abundant class of promoter-proximal transcripts that assemble and behave like unstable IncRNAs.

**DISCUSSION**

Our data reveal that distinct transcript classes are defined during 3′ end formation, with RNP compositions tailored to the functions and fates of the transcripts. These classes loosely align with existing annotations, but we identified hundreds of exceptions. To address the extensive overlap between, and heterogeneity within, annotated transcript classes, we suggest an improved RNP-based classification, which reflects how transcripts are regulated and how they might function.

**Tailored RNP Composition**

There has been much debate about the function of pervasive transcription in eukaryotes. We find that CUTs and SUTs are predominantly retained and degraded in the nucleus, suggesting that their functions primarily arise from the act of transcription rather than the transcript itself. This refutes the notion that SUTs are generally “stable,” but agrees with recent analyses identifying SUTs among exosome substrates (Gudipati et al., 2012b; Schneider et al., 2012). However, SUTs were mildly less prone to nuclear turnover than were CUTs, and their RNP composition overlapped with that of mRNAs, suggesting that some SUTs might function as stable transcripts. Xrn1 and the cytoplasmic exosome do not appear to function widely in bulk IncRNA turnover, but we suggest they degrade mRNA-like SUTs and provide a fail-safe for leaky nuclear surveillance. A substantial IncRNA class, termed XUTs, was reported to be exported and degraded in the cytoplasm by Xrn1 (van Dijk et al., 2011), but we see little evidence for this (Figure 1). We also uncovered extensive heterogeneity in mRNP composition, with one mRNA class regulated by IncRNAs or subject to IncRNA-like turnover in the nucleus, and another regulated by Nab2 and Rrp44 (Gudipati et al., 2012b; Schmid et al., 2012). Other mRNAs were primarily subject to cytoplasmic regulation. We suggest that tailored RNP compositions enable transcripts to be regulated and localized in a way appropriate to their function (or that of the encoded protein).

**The Multicolored Transcriptome**

Conventional transcriptome profiling experiments (e.g., RNA-Seq) struggle to distinguish overlapping transcripts, particularly where one is less abundant. By combining high-resolution binding data for many RNP proteins, we obtained a “multicolored” view of the transcriptome and could readily distinguish overlapping transcripts with different RNP profiles. Most strikingly, this revealed that the nuclear surveillance machinery targets a major class of promoter-proximal IncRNAs apparently generated by early transcription termination and with an RNP composition resembling CUTs.

Our analyses suggest that this early termination is prevalent for “IncRNA-like” mRNAs, but occurs to some extent for most mRNAs, and we speculate that this reflects a checkpoint in Pol II transcription. Mtr4 hits peaked within ~150 nt of mRNA TSSs, coincident with locations of Pol II pausing (Churchman and Weissman, 2011). Here, transcription initiation factors exchange for elongation factors (Mayer et al., 2010) prior to the polymerase traversing the +2 nucleosome dyad ~90 nt further downstream, which can impede elongation. If remodeling of the transcription complex is unsuccessful or slow, we suggest that Pol II transcription is terminated. Termination might involve Nrd1, Nab3, and/or Rat1, which crosslink to promoter-proximal IncRNAs (Creamer et al., 2011; Wlotzka et al., 2011 and unpublished data) and are implicated in the early termination of some mRNAs and IncRNAs (Geisler et al., 2012). The oligoadenylated 3′ ends that we detect suggest that termination generates an entry site for the TRAMP and exosome complexes. This is consistent with either an Nrd1-dependent mechanism or endonuclease cleavage followed by Rat1-dependent termination, which was recently identified in humans (Wagschal et al., 2012). The distribution of Mtr4 hits suggests that this checkpoint is restricted to the first ~500 nt, consistent with the exclusion of termination factors from the midregions of genes by Y1P modification of the Pol II CTD (Mayer et al., 2012). The extent to which these early terminating transcripts function as ncRNAs remains to be determined.

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(C) Average distribution of binding sites for Hrp1 (top) and Nab2 (bottom) across mRNAs, SUTs, and CUTs in the wild-type background. For Nab2 binding to CUTs an rpd3S strain is also shown.

(D) Frequency of UAUAUA motifs in Hrp1-binding sites near the 3′ end of mRNAs, CUTs, and SUTs (Xu et al., 2009).

(E) Northern analysis of CUT479, CUT200, and CCW12 abundance in wild-type, *P*Δ*na*–Nab2 and *P*Δ*na*–Hrp1 strains after glucose-dependent repression. Northern signals were quantified for replicate experiments (right). Bottom: western analysis of HA–Hrp1, HA–Nab2, and Mtr4 abundance.
Figure 5. Nuclear Surveillance Factors Bind to Promoter-Proximal lncRNAs Arising from Early Termination within Protein-Coding Genes

(A) Frequency of non-genome-encoded A-tails in CRAC data sets.

(B) Length distribution of non-genome-encoded A-tails at the 3' end of mapped reads in selected data sets (relative to the abundance of 2 nt tails).

(C) Length distribution of A-tails on raw reads (% of all analyzed reads). In comparison to (B), this analysis can detect longer A-tails but does not distinguish between genome-encoded and nonencoded tails.

(D) Hits in tRNAs

(E) Hit density across mRNAs

(F) Mtr4 hits in mRNAs

(G) Nab2 - different A-tail lengths

(H) Mtr4 binding (CRAC)

(Nrd1 binding (CRAC)

(Nab3 binding (CRAC)

(Elongating Pol II)

(3' ends of “CUTs”)

Graphs scaled to maximum height

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Determining Transcript Fate

Our analyses also revealed characteristics by which transcript classes are distinguished in the cell, with 3' end formation emerging as a key step (Figure 6). For mRNAs and SUTs we detected the hallmarks of cleavage and polyadenylation, including an appropriate configuration of sequence elements, a Pab1-bound poly(A) tail, and Hrp1 bound to an efficiency element /C24 50 nt upstream of the pA site. In contrast, these were absent from CUTs, which therefore undergo a distinct termination pathway. This is most likely Nrd1-dependent termination, which is associated with exosome recruitment potentially explaining the inherently low stability of CUTs (Vasiljeva and Buratowski, 2006). The RNP composition of CUTs was related to that of snoRNAs (Figure 1), for which Nrd1-dependent termination is well established.

Analyses of two mRNA cleavage and polyadenylation factors, Hrp1 and Nab2, revealed additional roles in the surveillance of CUTs and promoter-proximal RNAs, supporting our conclusion that 3' end formation is a key step in determining transcript fate. We propose that Hrp1 and Nab2 have dual-functions. Sequence nonspecific binding may be a default activity that is associated with recruitment of the nuclear surveillance system. In contrast, Hrp1 bound to the UAUAUA motif and Nab2 bound to poly(A) at the 3' end of mRNAs, in the context of a cleavage/polyadenylation complex, assist in correct mRNP maturation. (D) Prevalence of tRNAs among all (blue) or A-tailed (red) RNA fragments recovered for Mtr4, Ski2, and Xrn1. (E) Average distribution of all (gray) and A-tailed (blue) Ski2 and Mtr4 hits across scaled mRNAs. (F) Distribution of Mtr4 hits across individual mRNAs aligned by their TSSs (n = 1,000). (G) Average distribution of all (gray) and A-tailed (blue) Nab2 hits across scaled mRNAs (left), and average distribution of Nab2 hits with different length A-tails across scaled mRNAs (right). (H) Average distribution around mRNA TSSs and 3' ends of (1) Mtr4-, Nrd1-, and Nab3-binding sites (CRAC) (Wlotzka et al., 2011), (2) elongating Pol II (Churchman and Weissman, 2011) and (3) 3' ends of unstable transcripts (“CUT fraction”) (Neil et al., 2009).

Functions for Nab2 in RNA surveillance are consistent with its interactions with Rrp6 and Trf4, and its role in intron-containing pre-mRNA turnover and Rrp6-dependent autoregulation of the NAB2 transcript (Roth et al., 2009; Schmid et al., 2012). A function for Hrp1 in surveillance is consistent with reports that Hrp1 participates in Nrd1-dependent termination (Kuehner and Brow, 2008) and cytoplasmic nonsense-mediated decay (NMD) (González et al., 2000). The extensive contacts made by Hrp1 and Nab2 across the body of transcripts is consistent with ChIP data (González-Aguilera et al., 2011; Kim et al., 2004), whereas the human Hrp1 homolog TDP-43 is a ubiquitous RNP component with IncRNAs among its targets (Tollervey et al., 2011). These data suggest that Nab2, Hrp1, and TDP-43 function widely as RNA packaging factors. Nab2 is reported to fold poly(A) tails into a particular configuration (Viphakone et al., 2008), and in Nab2 mutants mRNAs are both hyperadenylated and aberrantly compacted (Brockmann et al., 2012). This suggests that RNA packaging and processing are tightly coupled, and appropriate RNA folding by Nab2 and Hrp1 might regulate access to the mRNA 3' end and/or recruitment of surveillance and cleavage/polyadenylation factors. RNA fate may be determined not only by the protein factors bound but also by the folding of the RNA.

The ability of SUTs to undergo mRNA-like cleavage and polyadenylation may explain why SUTs are more stable than CUTs (Figure 6). However, unlike mRNAs, most SUTs are retained in the nucleus, with only a minority showing “mRNA-like” RNP compositions. We propose that additional layers of regulation following cleavage and polyadenylation
determine whether a transcript is retained or exported. One candidate is Hek2, which bound to mRNAs and some “mRNA-like” SUTs but not to nuclear-restricted IncRNAs (Figure S2C). Hek2 has roles in mRNA stabilization and localization, and a human homolog (xCP2) enhances the 3’ end processing and stability of hz-mRNA (Ji et al., 2011). We suggest that Hek2 contributes to a decision point associated with 3’ end formation and selectively marks mRNAs, and some SUTs, for export. We predict that other sequence-specific binding proteins assist in the export of the “mRNA-like” SUTs that are not bound by Hek2.

In conclusion, our systematic analysis of RNP composition sheds light on how diverse classes of transcripts are distinguished in the cell and provides insights into the functions of these transcripts and of key RNP proteins.

EXPERIMENTAL PROCEDURES

Crosslinking and Analysis of cDNAs
We used the CRAC method as previously described (Granneman et al., 2011). In vivo protein-RNA crosslinks were generated by irradiating yeast cultures with UV light (254 nm, 100 s). Illumina sequencing reads were aligned to the yeast genome (SGD v64) using Novaalign. To quantify hits for particular genomic features and identify binding motifs, we used the pyCRAC package developed by Sander Granneman and custom Python/AWK scripts (available upon request). Analyses of hits in spliced versus unspliced transcripts and of non-genome-encoded oligo(A) tails were performed as previously described (Schneider et al., 2012; Wlotzka et al., 2011), and k-medians clustering used Cluster 3.0. Further information is provided in the extended experimental procedures and Table S5.

ACCESSION NUMBERS

The GEO accession number for all sequence data reported in this paper is GSE46742.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, two figures, and five tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.07.047.

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