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Ribosome Profiling Reveals Pervasive Translation Outside of Annotated Protein-Coding Genes

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SUMMARY

Ribosome profiling suggests that ribosomes occupy many regions of the transcriptome thought to be noncoding, including 5′ UTRs and long noncoding RNAs (lncRNAs). Apparent ribosome footprints outside of protein-coding regions raise the possibility of artifacts unrelated to translation, particularly when they occupy multiple, overlapping open reading frames (ORFs). Here, we show hallmarks of translation in these footprints: copurification with the large ribosomal subunit, response to drugs targeting elongation, trinucleotide periodicity, and initiation at early AUGs. We develop a metric for distinguishing between 80S footprints and nonribosomal sources using footprint size distributions, which validates the vast majority of footprints outside of coding regions. We present evidence for polypeptide production beyond annotated genes, including the induction of immune responses following human cytomegalovirus (HCMV) infection. Translation is pervasive on cytosolic transcripts outside of conserved reading frames, and direct detection of this expanded universe of translated products enables efforts at understanding how cells manage and exploit its consequences.

INTRODUCTION

Identifying the genomic regions that are transcribed and translated is a fundamental step in annotating a genome and understanding its expression. A variety of microarray- and sequencing-based approaches can reveal the mRNA content of the cell (Bertone et al., 2004; Carninci et al., 2005; Wang et al., 2009), but it has proven more challenging to experimentally define translated sequences within the genome or the transcriptome. Historically, protein-coding sequences were discovered by search for long (>100 codon) open reading frames, which are unlikely to occur in the absence of selection against stop codons. Widespread use of this approach has also been based on the assumption that short peptides are unlikely to fold into stable structures and thus perform robust biological functions. Recently, more-sophisticated conservation-based metrics, such as PhyloCSF, were developed for the computational identification of sequences that appear to encode proteins over a broad size range (Lin et al., 2008, 2011). However, these approaches focus on identifying regions of the genome experiencing selective pressure to maintain a reading frame encoding a functional protein. The questions of which parts of the genome are translated and whether or not the protein product has an adaptive function in the cell are related but distinct; the former can be answered by experimentally finding the locations of ribosomes on mRNAs.

Global profiling of transcription and mRNA abundance has revealed a class of transcripts with no clear protein-coding potential (Bertone et al., 2004; Carninci et al., 2005; Guttman et al., 2009). Many of these RNAs were long RNA polymerase II products, transcribed from genomic regions far from known protein-coding genes and thus were named long noncoding RNAs (lncRNAs).

The discovery of these surprising RNAs in the transcriptome as well as the existence of short upstream open reading frames (uORFs) in 5′ leader regions (often referred to as 5′ UTRs; Calvo et al., 2009; Wethmar et al., 2014) highlight the need for comparable direct, experimental maps of translation. Whereas, based on both lack of conservation and the distribution of ribosome-protected fragments, there is strong evidence that most lncRNAs do not encode proteins with conserved adaptive cellular roles (Cabili et al., 2011; Chew et al., 2013; Guttman et al., 2013), these computational approaches could miss functional coding sequences, particularly those that are short and/or species specific (Reinhardt et al., 2013). Furthermore, translation and protein synthesis have impacts beyond the production of stable proteins with discrete molecular functions—polypeptide products from all cellular translation must be degraded and noncanonical translation products yield unanticipated...
Figure 1. Fragment Length Analysis Distinguishes True Ribosome Footprints on Coding and Noncoding Sequences

(A–E) Distribution of fragment lengths mapping to nuclear coding sequences (CDSs) compared to (A) the telomerase RNA Terc, (B) mitochondrial coding sequences, (C) snoRNA host gene Snhg5, (D) ENCODE lncRNAs, and (E) 5' UTRs of protein-coding genes in ribosome-profiling data from emetine-treated mESCs. (F) Metric comparing the similarity of two length distributions.

(legend continued on next page)
antigens that may play roles in viral detection or in autoimmunity (Starck et al., 2012). Finally, the process of translation can affect the stability of the template message by triggering cotranslational decay pathways including nonsense-mediated decay (NMD) (Rebbaapragada and Lykke-Andersen, 2009). Knowing what transcripts are translated has important implications for the fate of the RNA, the ribosome, and the cell. The ribosome-profiling technique provides a unique opportunity to experimentally address this question.

Ribosome profiling is an approach for mapping the exact position of translating ribosomes across the transcriptome by deep sequencing of the mRNA footprints that are occupied by the ribosomes and thereby physically protected from nuclease digestion (Ingolia et al., 2009; Steitz, 1969; Wolin and Walter, 1988). Analysis of these ribosome-protected mRNA fragments yields a quantitative and detailed map of ribosome occupancy that reveals translation in the cell with single-nucleotide resolution. Most ribosome footprints fall within known coding sequences, where they showed three-nucleotide periodicity reflecting the triplet nature of the genetic code. However, ribosome-profiling data suggested that some predicted noncoding regions of the transcriptome were translated (Ingolia et al., 2011). In some cases, these footprints were organized on single reading frames that closely resembled known coding sequences except for their shorter length (Brar et al., 2012; Stern-Ginossar et al., 2012). In other cases, footprints were not restricted to a single predominant reading frame based on metrics such as the ribosome release score (RRS) or the disengagement score (DS) (Chew et al., 2013; Guttman et al., 2013). This second group of predicted translated sequences, present on some IncRNAs as well as the 5’ leaders of many mRNAs, can be distinguished both from conserved protein-coding genes, where one single reading frame does predominate, and from the 3’ UTRs of most mRNAs, which are devoid of ribosome footprints (Chew et al., 2013). The high ribosome occupancy on some of these regions, comparable to that on protein-coding genes, suggests a similar stoichiometry of polypeptide production.

The broad implications of pervasive translation and the discrepancy between ribosome profiling and conservation analysis pose an immediate question: do the footprint sequences detected in these profiling experiments indicate the presence of assembled (80S) ribosomes? Here, we address this question and present several ways to distinguish true 80S footprints in ribosome-profiling data. We first classify protected RNA fragments based on their size distribution, a purely computational analysis that can be applied to existing data and to new profiling data collected without experimental modification. Our analysis discriminates cleanly between true footprints and known sources of contamination. We validate the results from our fragment length classifier with two new lines of experimental evidence: drugs that target the elongating 80S ribosome specifically and affinity purification of the large ribosomal subunit, both of which support the translation of IncRNAs and 5’ UTRs. We also show that footprints on these noncoding sequences demonstrate hallmarks of eukaryotic translation. Finally, we verify the accumulation of protein products from noncanonical translation and demonstrate the potential functional impact of novel human cytomegalovirus (HCMV) proteins as a source of viral antigens. Our results show that the universe of translated regions extends beyond long conserved regions encoding large, well-conserved proteins.

RESULTS

The Characteristic Length of Ribosome Footprints Distinguishes Them from Background RNA Fragments

The ribosome physically encloses its mRNA template and protects a characteristic length of this RNA from nuclease digestion (Steitz, 1969; Wolin and Walter, 1988). In ribosome-profiling data, the overall size distribution of fragments derived from protein-coding sequences, which should predominately reflect true ribosome footprints, differs from the lengths of the abundant rRNA contamination found in profiling samples (Ingolia et al., 2009, 2011). We reasoned that fragment size could likewise distinguish true ribosome footprints from other, nonribosomal contaminants, such as RNA regions that are protected by protein complexes or stable RNA secondary structure. The exact length distribution of protected fragments can vary slightly between samples, likely due to differences in digestion conditions (Ingolia et al., 2012). Furthermore, distinct ribosome conformations can lead to significantly different mRNA footprints lengths (Lareau et al., 2014), and the predominant conformation may vary between samples. In order to avoid these confounding effects, we compared the size distributions of fragments derived from noncoding sequences to those on protein-coding genes within a single sample, treated with translation elongation inhibitors that should capture most ribosomes in a specific state (Lareau et al., 2014; Wolin and Walter, 1988). We gathered new ribosome-profiling data from mouse embryonic stem cells (mESCs) treated with the translation elongation inhibitor emetine in order to obtain footprints with stronger reading frame bias (Ingolia et al., 2011, 2012). Fragment size distributions in this sample clearly distinguished true ribosome footprints, which predominate on coding sequences, from background RNA contained in nonribosomal ribonucleoprotein (RNP) complexes such as telomerase (Figure 1A). They also separated (G) Fragment length analysis plot of total reads per transcript and FLOSS relative to the nuclear coding sequence average. An FLOSS cutoff is based on an extreme outlier threshold for annotated coding sequences. IncRNAs resemble annotated, nuclear protein-coding genes, whereas functional RNAs and mitochondrial coding sequences are distinct.

(H) As (G), comparing 5’ UTRs and coding sequences of nuclear-encoded mRNAs.

(I) Read count profile on Malat1 with an inset showing ribosomes on a non-AUG uORF and the first reading frame at the 5’ end of the transcript. An inset shows the fragment length distribution for the first reading frame, which matches the overall coding sequence average, and the whole transcript, which does not.

(J) Fragment length analysis showing the shift from the entire Malat1 transcript, which contains substantial background, to the first Malat1 reading frame, which contains true ribosome footprints.

(K) Read count profile across the primary Gass transcript with the snoRNAs and the fully spliced transcript shown.

(L) As (J) for the primary GASS transcript, containing snoRNA precursors, and the fully spliced product.
footprints of the 80S ribosome from fragments of mitochondrial coding sequences that likely reflect footprints of the distinct mitochondrial ribosome (Figure 1B) and noncoding short RNAs that associate with the cytosolic ribosome or its precursors, such as small nuclear RNAs (snRNAs) (Figure 1C). By contrast, RNA fragments derived from IncRNAs and from 5′ UTRs showed a size distribution much like that seen on coding sequences (Figures 1D and 1E). This similarity provides evidence that the protected fragments on these two classes of noncoding sequences consist principally of 80S ribosome footprints and, thus, that translation occurs outside of annotated protein-coding regions.

**Classifying the Translation Status of Individual Transcripts and Subregions**

We next adapted our fragment length distribution analysis to distinguish between individual transcripts that show substantial background fragments from those having true 80S footprints. When hundreds or thousands of ribosome-footprint-sequencing reads are available for a single transcript, their length distribution should converge to match the characteristic ribosome footprint size. We define a fragment length organization similarity score (FLOSS) that measures the magnitude of disagreement between these two distributions, with lower scores reflecting higher similarity (Figure 1F). Thousands of well-expressed protein-coding transcripts almost uniformly scored well, and the similarity improved with increasing read counts, as expected (Figure 1G).

As with many sequencing-based analyses, this metric is less informative on transcripts with few reads—an inevitable consequence of sampling error in estimating the fragment length distribution—but we are most interested in the transcripts with many reads and, thus, clear FLOSS results.

In order to contrast nonribosomal background with true ribosome footprints, we needed canonical set of nontranslated RNAs to compare with annotated protein-coding sequences. We selected transcripts with well-established molecular functions as RNAs and features likely to suppress their translation, such as an absence of 5′ methyguanosine caps or assembly into stable ribonucleoprotein structures inaccessible to the translational machinery. Many of these transcripts, defined in previous studies as “classical” noncoding RNAs (Guttman et al., 2013), in fact yielded very few protected fragments. We did find several (including telomerase RNA, vault RNA, and RNase P) that we could test, however, and found that each could be distinguished clearly from annotated coding sequences. Likewise, every individual mitochondrially encoded message stood out from nuclear genes. We concluded that this metric discriminates reliably between true 80S ribosome footprints and background RNA fragments on specific transcripts as well as on broad classes of RNAs.

FLOSS analysis revealed that ribosome-profiling-derived reads from IncRNAs and 5′ UTRs overwhelmingly reflect true ribosome footprints. Protected fragments on nearly every individual IncRNA showed a FLOSS value very similar to that seen on coding sequences, in contrast to background from classical noncoding RNAs (Figure 1G). Individual 5′ UTRs also grouped very well with coding sequences (Figure 1H).

We formalized this classification by defining a threshold FLOSS value, excluding transcripts that differed greatly from annotated protein-coding genes. We set this threshold based on the read counts and FLOSS values for known protein-coding genes using Tukey’s method, a widely accepted nonparametric criterion for extreme outliers (Tukey, 1977). This cutoff eliminated all classical noncoding RNAs with substantial (>100 reads) expression while retaining almost all annotated mRNAs (99.6%). The perfect specificity and extraordinary sensitivity likely overestimate the true performance of this metric, especially on transcripts that contribute a mixture of true translation and background. Nonetheless, the vast majority of 5′ UTRs (96%) and IncRNAs (90%) were classified with protein-coding genes (Figures S1A and S1B). Not all 5′ UTRs or IncRNAs produced protected RNA fragments in profiling experiments, but when fragments did appear, they generally resembled the ribosome footprints of coding sequences, suggesting true translation in these regions.

We previously reported apparent ribosome occupancy on the abundant and prototypical IncRNA Malat1, which is predominantly nuclear and thus is largely separated from the translational apparatus (Wilusz et al., 2008). This surprising result led us to investigate protected Malat1 RNA fragments more closely (see Figure 1I). We saw a pattern that was highly suggestive of ribosome occupancy near the 5′ end of the transcript, covering the first AUG-initiated reading frame with substantially lower ribosome density after the corresponding in-frame stop codon. We also saw several other sites in Malat1 that produced abundant protected RNA fragments. Whereas the overall distribution of Malat1 fragment lengths did not resemble the profile of true ribosome footprints, the first short reading frame did appear to contain 80S ribosomes (Figures 1I and 1J). Similarly, whereas the full Malat1 transcript stood out from protein-coding genes by fragment length analysis, the upstream reading frame resembled those of ordinary protein-coding genes. Thus, Malat1 RNA fragments appear both to contribute nonribosomal background, like telomerase or RNase P, and also to represent footprints from ribosomes translating its first ORF. As Malat1 is predominantly nuclear, whereas the translation occurs in the cytosol, it would be interesting to find the ribosome density and the relative background contribution in the cytoplasmic fraction. MALAT1 is also unusual in that the mature form is not polyadenylated, but the triple-helix structure that protects its nonadenylated 3′ end also supports efficient translation (Wilusz et al., 2012); the role of these ribosomes, if any, in the function of Malat1 remains to be determined.

The noncoding RNA Gas5 also yielded a complex mixture of translation and background RNA that could be separated by fragment length analysis. Gas5 is a snoRNA host gene whose introns contain several snoRNAs; there are no long or conserved reading frames in the mature message. Nonetheless, the spliced RNA associates with ribosomes in order to trigger its degradation by NMD (Smith and Steitz, 1998). Fragment length analysis of the primary Gas5 transcript indicates that it is a source of background RNA in profiling experiments, corresponding principally to the intronic snoRNAs (Figure 1K). Fragments that mapped to the fully processed Gas5 transcript, with no remaining snoRNA sequences, resembled 80S footprints on coding sequences (Figures 1K and 1L). They were also concentrated in reading frames near the 5′ end of the transcript, where translation is expected to occur.
Taken together, these analyses show that fragment length analysis can discriminate between true 80S footprints and background RNA reads in ribosome-profiling data. Furthermore, this simple metric can be applied to existing profiling data sets as well as incorporated into computational workflows with no change to experimental protocols. It provides strong evidence for the presence of ribosomes based on comparisons with RNAs whose biology is well understood. As this analysis is correlative, however, we performed direct experimental tests to confirm that footprints on noncoding sequences reflected true translation.

Drugs that Inhibit Translation Specifically Affect Elongating Ribosome Footprints on Coding and Noncoding Sequences

Diverse translation inhibitors target distinct sites on the ribosome with high affinity and selectivity (McCoy et al., 2011; Schneider-Poetsch et al., 2010). We previously observed that mammalian cells treated with one such drug, cycloheximide, yielded ~1 nt shorter ribosome footprints over the body of open reading frames than those treated with another, emetine (Figure 2A; Ingolia et al., 2011). Both emetine and cycloheximide target the ribosome specifically, and so the differences observed in mammalian cells between these two drugs should appear only in true footprints of elongating ribosomes.

We set out to use the selectivity of these drugs for the ribosome as an additional test to distinguish true footprints. In aggregate, fragments on IncRNAs and on 5' UTRs showed a similar but more modest length shift to that seen on protein-coding genes—the cumulative length distribution on both noncoding regions is larger in emetine than in cycloheximide (Figures 2B and 2C). Drug treatment may affect footprints on noncoding RNAs less than those on coding sequences because the translated reading frames on these RNAs are short, and thus terminating ribosomes, whose footprints appear to differ slightly from elongating ribosomes (Ingolia et al., 2011), comprise a larger fraction of the total ribosomes. Alternately, a fraction of these footprints may reflect posttermination ribosome footprints, which can accumulate in yeast defective for ribosome-recycling factors and which should not respond to drugs targeting elongation (Guydosh and Green, 2014). Nonribosomal background fragments do not shift in length between these two elongation inhibitors (Figure 2D).

We gathered new ribosome-profiling data from cycloheximide-stabilized yeast polysomes in each sample in order to monitor any differences in the extent of nuclease digestion between samples (Figure 2E). The true ribosome footprints on annotated coding sequences were again shorter from cycloheximide-treated than from emetine-treated mESCs.
Figure 3. Ribosome Affinity Purification Separates 80S Footprints from Background RNA
(A) Schematic showing that affinity purification of tagged 60S ribosome subunits recovers 80S footprints but depletes background from nonribosomal RNPs, potential scanning 40S footprints, and footprints of untagged yeast 80S ribosomes.
cells, though the difference was less pronounced (Figure 2F). The length of footprints on IncRNAs also shifted in response to treatment with elongation inhibitors (Figure 2G), and these length shifts were significant on protein-coding genes ($p < 1 \times 10^{-5}$), 5’ UTRs ($p < 1 \times 10^{-6}$), and on IncRNAs ($p < 0.01$; Figure S2). In contrast, the footprints from the yeast ribosomes included as an internal control showed, if anything, a very modest shift in the opposite direction (Figure 2H) that did not rise to the level of significance ($p > 0.05$; Figure S2), arguing that the reproducible difference between cycloheximide- and emetine-treated polysomes did not result from differences in nuclease digestion or library generation that affect all RNA fragments in a sample.

Ribosome Footprints on Classical Coding Sequences, 5’ UTRs, and IncRNAs Copurify with the Large Ribosomal Subunit

We next sought to verify that footprints seen outside of annotated coding regions copurified specifically with the ribosome. Ribosome affinity purification would provide strong evidence that footprints on IncRNAs and on 5’ UTRs were bound to the ribosome (Figure 3A). We typically recover ribosomes by sedimentation in an ultracentrifuge, but this purification provides little specificity for ribosomes over other large RNPs. The most prominent classical noncoding RNAs that contribute to background in ribosome-profiling experiments are components of nonribosomal RNPs, such as RNase P, telomerase, and the vault RNP (Figure 1G). We infer that these RNP assemblies both protect RNA fragments from digestion and then sediment with ribosomes, and it seemed possible that some apparent ribosome footprints on IncRNAs actually reflected the incorporation of the IncRNA into a similar RNP complex.

Specific affinity purification of the ribosome would deplete background from these RNPs. The large (60S) subunit joins at the last step in translation initiation and does not associate with mRNA prior to this time, and so any footprint associated with the 60S subunit derives from a ribosome that has completed initiation and begun translation (Aitken and Lorsch, 2012). Ribosome-profiling data are unlikely to include footprints of small (40S) subunits scanning 5’ UTRs prior to initiation, because these complexes are unstable in the absence of chemical crosslinking and are expected to protect a different mRNA footprint size from assembled 80S ribosomes (Valásek et al., 2007). Nonetheless, we wished to verify that footprints on 5’ UTRs reflected postinitiation-assembled (80S) ribosomes.

In order to purify 80S (and 60S) ribosomes specifically, we developed an affinity-tagged version of large subunit ribosomal protein L1 (formerly RPL10A). Several ribosome epitope tags have been developed for lineage-specific polysome isolation, including the translating ribosome affinity purification tag, in which L1 is fused to enhanced GFP (Heiman et al., 2009). We believed that in vivo biotinylation of L1 would offer advantages over epitope tags, allowing us to exploit the high affinity and rapid association of biotin and streptavidin to purify tagged ribosomes. We placed a biotin acceptor peptide at the end of a long, flexible linker at the C terminus of L1 and coexpressed this tagged protein along with birA, the cognate E. coli biotin ligase, in human embryonic kidney 293 (HEK293) cells. Tagged L1 was biotinylated, dependent on the presence of birA, and L1-biotin was incorporated into ribosomes.

In order to test our enrichment of tagged ribosomes, we mixed lysate from human cells expressing L1-biotin (in addition to their endogenous L1) with a control yeast lysate lacking biotinylated ribosomes and compared the fate of the human ribosome footprints to footprints from yeast genes. We performed nuclease footprinting of this mixture, collected all ribosomes by filtration through Sephacryl S400 columns, and purified the tagged human ribosomes by streptavidin affinity. Footprints from human protein-coding genes were strongly enriched in the streptavidin-bound sample relative to footprints from yeast transcripts (Figure 3B). The only exception was the yeast gene ACC1, which encodes the endogenous yeast biotin carrier protein. We assume that this is biotinylated cotranslationally in vivo and so footprints recovered by affinity purification through the nascent chain. Consistent with this model, only footprints from the 3’ end of ACC1, corresponding to ribosomes that have synthesized the biotin acceptor site of Acc1p, are enriched. Importantly, the observed specificity for human mRNAs also excluded postlysis association of human ribosomes to yeast mRNAs, arguing strongly that footprints seen in ribosome-profiling experiments reflect translation that initiated in vivo prior to cell lysis. Fragment length distribution analysis provided further evidence against human ribosomes subject to affinity enrichment on yeast mRNAs, as protected fragments on human and yeast ribosomes are distinct in the mixed lysate and there was no evidence for a shift toward human fragment lengths on yeast messages following affinity purification. Human snoRNA reads also copurified with biotinylated L1, though somewhat less efficiently than ribosome footprints, as we expect due to their binding to preribosomal complexes in order to guide pre-rRNA modification (Figures S3A–S3C).

We then investigated the fate of other human-derived background reads following affinity purification of ribosomes. As noted above, profiling data after conventional ribosome sedimentation in HEK cells contained fragments mapping to several classical noncoding RNAs that also appeared in the mESC profiling, such as RNase P. Fragment length analysis using the
FLOSS reliably discriminated this background from footprints on coding sequences (Figure 3D). These same transcript fragments were also depleted in affinity-purified profiling samples, at least as strongly as were yeast-coding sequences (Figures 3E and 3F). Fragments from mitochondrial coding sequences were also strongly depleted, as the mitochondrial ribosome, which is entirely distinct from the cytosolic ribosome, lacked a biotin tag.

Having established affinity purification as a physical separation of background RNA fragments from true ribosome footprints, we next turned to investigate the status of apparent ribosome footprints in noncoding regions. We first verified that, as in mESCs, the protected fragments size distribution on HEK cell 5′ UTRs closely resembled ribosome footprints from the coding sequences (Figure S3D). These 5′ UTR protected fragments also copurified with the large ribosomal subunit in nearly all cases (Figure 3C). We thus conclude that these fragments are true 80S ribosome footprints and do not reflect scanning 40S subunits. Likewise, we found that protected fragments on most HEK lncRNAs are physically bound to the ribosome and likely reflect true translation of these noncoding RNAs (Figures 3G–3I). Furthermore, the small number of lncRNAs yielding substantial non-ribosome-associated fragments were independently identified as sources of background by the FLOSS analysis.

**Translation on lncRNAs Occurs in AUG-Initiated Reading Frames near the 5′ End of the Transcript**

lncRNAs lack a conserved, protein-coding reading frame by definition, and accordingly, ribosome footprints on these transcripts are not organized into a single, discrete reading frame without downstream translation in the manner seen on mRNAs (Chew et al., 2013; Guttman et al., 2013; Ingolia et al., 2011). Translation on lncRNAs and on mRNAs could differ fundamentally, however, and we wished to determine whether ribosome occupancy on lncRNAs show key features of eukaryotic translation. Whereas translation outside of annotated protein-coding regions often initiates at a variety of near-cognate codons in overlapping reading frames, obscuring some features of translation that manifest clearly on transcripts encoding a conserved protein, initiation should nonetheless be strongly biased toward AUG codons near the 5′ end of RNAs, and elongating ribosomes should show enrichment in the reading frame that follows until it ends at a stop codon. In order to evaluate the pattern of translation on lncRNA, we analyzed the initiation-site-profiling data we gathered from mESCs (Ingolia et al., 2011). We previously reported that brief treatment with the drug harringtonine causes ribosomes to accumulate at start codons while allowing run-off depletion of ribosomes over the rest of the coding sequence. This can be used to robustly identify translation initiation sites (Ingolia et al., 2011; Stern-Ginossar et al., 2012). Here, we use a simplified criterion to detect peaks of ribosome occupancy over AUG codons following harringtonine treatment (Figure 4A). This approach is robust against the possibility of concurrent translation of other, overlapping reading frames. Whereas we considered only AUG codons as candidate start sites, we found that, on the majority of lncRNAs, the start site we selected was the highest occupancy ribosome position of the entire RNA (Figure 4B), suggesting that this assumption was reasonable.

Initiation sites on lncRNAs detected in harringtonine-profiling data showed hallmarks of eukaryotic translation. In the canonical initiation pathway, factors bound to the 5′ cap recruit a preinitiation complex that scans the RNA directionally to identify a start codon. Consistent with this mechanism of translation, the start sites detected in harringtonine profiling generally fell near the beginning of the lncRNA, within a few hundred nucleotides of the 5′ end (Figure 4C) and at one of the first AUG codons on the transcript (Figure 4D). This bias toward early AUG codons is well explained by the classical model of eukaryotic initiation. By contrast, it is not likely that background RNA fragments not indicative of translation would show a strong preference for AUG codons near the 5′ end of transcripts.

Based on these observations, we next looked for evidence of elongating ribosome footprints in the reading frames associated with these initiation sites. Earlier studies argued against the predominance of a single open reading frame on lncRNAs. Both studies employed similar metrics—the RRS or the DS—to demonstrate that the abrupt drop in ribosome occupancy at the end of coding sequences was not seen for short reading frames in 5′ UTRs and on lncRNAs (Chew et al., 2013; Guttman et al., 2013). The absence of clear termination in any single reading frame argues that multiple, overlapping reading frames are translated on these RNAs. Nonetheless, we expected that the start sites we detected should result in elevated ribosome occupancy in the downstream open reading frame relative to the overall transcript. Indeed, we found the observed number of ribosome footprints within predicted reading frames on lncRNAs exceeded the number expected based on the overall ribosome density the length of the reading frame, often 10-fold or more, and never strongly depleted relative to the transcript overall (Figure 4E). This comparison is related to the inside/outside score, the ratio of footprints inside versus outside a candidate reading frame, used by Chew et al. (2013). Furthermore, we found that footprints within the open reading frame immediately following the predicted strongest initiation site on a lncRNA showed codon periodicity relative to that start site, similar to the periodicity seen in annotated protein-coding genes, whereas footprints outside of these reading frames do not (Figure 4F). This pattern of footprint occupancy is consistent with substantial in-frame translation from the predicted start site occurring alongside translation of many other reading frames on the transcript, including those initiating at near-cognate, non-AUG sites. This translation, particularly the downstream component that lacks a reading frame signal relative to the strongest AUG start site and thus reflects overlapping translation in alternate reading frames, would reduce RRS and DS metrics on these lncRNAs relative to annotated mRNAs.

**Fragment Length Analysis Supports Translation on Novel Reading Frames in Meiotic Yeast**

In previous studies, we defined translated reading frames in meiotic budding yeast using ribosome-profiling data (Brar et al., 2012). We wished to determine whether FLOSS analysis could be applied in this distantly related organism to support our annotations. Cycloheximide-stabilized ribosome footprints lying within yeast-coding sequences show a tight size distribution, as we observed previously, which could be readily...
distinguished from background RNA fragments derived from nontranslated yeast RNAs, including tRNAs and isolated snRNAs, and from the validated yeast meiotic noncoding RNAs \textit{IRT1}, \textit{RME2}, and \textit{RME3} (Figure 5A). As in mammals, we also found fragments of mitochondrial mRNAs, likely representing footprints of the mitoribosome, which were larger than cytosolic ribosome footprints. By contrast, the protected fragments on the large majority of new, independent ORFs and on upstram ORFs in the 5′ UTRs of annotated protein-coding genes matched the size of true ribosome footprints closely (Figures 5B–5D). FLOSS analysis discriminated well between individual annotated coding sequences and noncoding transcripts (Figure 5E).
and classified nearly all novel ORFs with known protein-coding genes (Figure 5F). Thus, considered singly or as a group, our reading frame annotations, defined solely by ribosome-profiling data, represent the presence of 80S ribosomes and not background RNA fragments.

We also sought to test whether productive translation could be detected from the ribosomes occupying these novel short reading frames. We integrated a GFP reading frame at the 3' end of meiotically regulated short reading frames in yeast (Figures 5G and 5H). Fusion protein from one short (72 codon) non-AUG uORF was detected among meiotic cells (Figures 5G and 5H).
reading frame accumulated in midmeiotic cells, as predicted from translation data, and localized to mitochondria (Figures 5I and 5J). GFP fused to another, 78-codon reading frame showed robust expression in vegetative cells that decreased in meiosis, consistent with expression-profiling data (Figure 5K). The fusion protein colocalized with the nucleus in vegetative cells (Figure 5L). The translational fusion of these short peptides with the large and well-folded GFP may artificially stabilize the protein products and enhance their accumulation. Nonetheless, these data confirm that the novel ORFs defined by ribosome profiling result in the synthesis of proteins and, further, that these short proteins can confer specific localization on a GFP fusion, suggesting that they can display some molecular activity in the cell.

**Fragment Length Analysis Supports Translation on Novel Reading Frames in Human Cytomegalovirus**

We recently published an annotation of HCMV open reading frames based on ribosome profiling of infected human foreskin fibroblasts (Stern-Ginossar et al., 2012). This annotation included many entirely novel reading frames as well as alternate versions of known proteins. The translation of many of our novel HCMV reading frames was confirmed previously by epitope tagging and by direct detection of native protein products through mass spectrometry (Stern-Ginossar et al., 2012). Our fragment length analysis revealed little difference between human protein-coding genes, well-known viral coding sequences, and newly identified ORFs (Figures 6A–6D). We next tested the FLOSS on individual HCMV ORFs and found that nearly all fell among the annotated human protein-coding genes (Figures 6E and 6F).

We may fail to detect proteins from other novel reading frames, despite the fact that they are actually synthesized in the cell, if they are highly unstable and thus low abundance. However, all translated polypeptides can serve as antigens, even if they are rapidly degraded and never accumulate within the cell. In fact, breakdown products from cotranslational degradation may be preferentially targeted for display as antigens. The adaptive immune system thus records signatures of past protein expression, and we wanted to mine this record by testing the antigenicity of the novel reading frames we identified in HCMV. We reasoned that, if humans with a history of CMV infection displayed T cell responses against novel peptides, as they do against canonical CMV proteins (Sylwester et al., 2005), it would indicate that these peptides were produced in the course of the normal viral life cycle in a human host. Furthermore, the T cell response would directly demonstrate the functional impact of short reading frame translation in viral infection.

We focused on the beta 2.7 transcript in HCMV. Despite its designation as a long noncoding RNA, ribosome-profiling data identified eight new, moderately sized ORFs, two of which (ORF6C and ORF7C) were identified in lysates from infected cells by mass spectrometry (Stern-Ginossar et al., 2012; Figure 6G). Human T cells from anonymous HCMV-positive donors revealed robust cellular immune responses to ORF7C and ORF6C, as well as to other short reading frames on beta 2.7 and other ORFs that we had identified by ribosome profiling (Figures 6H and 6I). These responses were absent from HCMV-negative individuals (Figure S2), supporting the natural exposure of HCMV-infected individuals specifically to these newly annotated translation products. Neither ORF6C nor ORF7C resembled annotated reading frames by the RRS metric, consistent with the polycistronic and overlapping translation on the beta 2.7 transcript (Figure 6G), but the encoded proteins are synthesized in culture models and in infected humans.

**DISCUSSION**

In this study, we establish the validity of ribosome profiling as a global and experimental strategy for identifying translated regions of a genome. Profiling data are an excellent complement to computational analyses, which detect conserved protein-coding regions of the genome, and to proteomic approaches for identifying stable proteins. These three techniques answer different but related questions. Conserved functional proteins are a subset of the total polypeptide content of the cell, which in turn is a subset of all products that are produced, however transiently, by translation. Ribosome profiling thus provides the most expansive view of the proteome and has thereby helped us appreciate a wider universe of translated sequences.

We present multiple lines of evidence that true ribosome footprints are pervasive on cytosolic RNAs, independent of the presence of conserved reading frames. These footprints change in response to translation inhibitors, copurify with the large ribosomal subunit, and fall preferentially in reading frames near the 5' ends of transcripts. The size distribution of ribosome-protected mRNA fragments also distinguishes them from the background present in profiling data. This observation allowed us to develop a fragment length analysis, the FLOSS, that very accurately predicts the results of ribosome affinity purification, which separate true footprints from background RNA by physical rather than computational means. In fact, because some noncoding RNAs do associate with the ribosome for reasons that are unrelated to their actual translation, the FLOSS appears to exclude background more effectively than ribosome pull-down. The large majority of regions identified in profiling experiments reflect true translation; background originates from a handful of known, abundant noncoding RNAs. The FLOSS can be easily incorporated into ribosome-profiling workflows, and we here provide tools for applying it based on the widely used Bioconductor project (Gentleman et al., 2004). The specific length distribution and FLOSS cutoff for each individual data set can be determined empirically based on annotated protein-coding genes serving as examples of true translation. Adoption of the FLOSS should further increase confidence that profiling measurements on individual transcripts reflect their translation and aid in removing the small number of RNAs that yield nonribosomal background.

Pervasive ribosome occupancy outside of annotated coding regions has been seen in diverse organisms, and we here present further evidence for the existence of protein products resulting from translation by these ribosomes. The biological implications of this translation remain to be explored, however. In part, it may reflect an imprecision that leads to translation with no functional relevance. We do not know of molecular features that would enable the translational apparatus to distinguish an mRNAs from a capped, polyadenylated, cytosolic IncRNA, and so it may not be surprising to find ribosomes on many IncRNAs.
Figure 6. Novel Human Cytomegalovirus Reading Frames Based on True Ribosome Footprints Lead to Antigens in Humans

(A–D) Distribution of fragment lengths mapping to human nuclear CDSs compared to all annotated CMV-coding sequences after (A) 5 hr or (B) 72 hr of infection and of specifically the (C) previously annotated and (D) novel CMV-coding sequences after 5 hr of infection.

(E and F) Fragment length analysis of human coding sequences compared to (E) previously annotated CMV reading frames and (F) novel CMV annotations.

(G) Ribosome footprint organization on beta 2.7 transcript (Stern-Ginossar et al., 2012).

(H) ELISPOT assay of human donor T cell responses to novel CMV reading frames along with controls.

(I) Quantitation of ELISPOT data.

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Imperfect rejection of near-AUG codons during translation initiation, combined with the presence of actual AUGs, could explain ribosome occupancy in many 5' UTRs. However, translation of these noncoding sequences has many potential consequences and noncoding sequences likely experience selection against translation with harmful effects. For example, AUG codons are depleted in many 5' UTRs, as they interfere with translation of the downstream protein-coding sequence, though this interference is exploited as a regulatory mechanism controlling the expression of genes such as Atf4 (Sonenberg and Hinnebusch, 2009). Other side effects of noncoding translation may likewise be avoided in some RNAs and co-opted in others.

The translation of an RNA can impact the transcript itself, and IncRNAs with specific molecular functions are likely subject to selective pressure to manage this translation and avoid interference with their other activities. The translating ribosome acts as a potent helicase that can displace RNA structure and remove RNA-binding proteins, potentially disrupting functional ribonucleoprotein complexes. We have shown that initiation and translation are biased toward the 5' ends of IncRNAs, as expected in eukaryotes, and so noncoding cytosolic transcripts may experience selection for benign 5' reading frames that capture ribosomes and protect functional elements occurring in the 3' end of the RNA (Ulitsky and Bartel, 2013). Short reading frames with atypical amino acid composition may resemble those found in aberrant mRNAs and trigger RNA decay through NMD or no-go decay, which were originally characterized as mRNA quality control pathways (Pérez-Ortín et al., 2013). Translated sequences may also exert cis-acting effects through the peptides they encode, for example, by cotranslational recruitment of the nascent chain attached to the ribosome and the transcript, to specific structures in the cell (Yanagitani et al., 2009).

Translation results in the synthesis of a polypeptide, regardless of whether an RNA sequence encodes a functional protein constrained by selection, and we have now detected proteins synthesized from novel translated sequences predicted by ribosome profiling in yeast and given evidence for their presence in humans during CMV infection. These unconstrained peptide sequences may not adopt a specific fold and may occupy cotranslational folding or degradation machinery, and those peptides escaping surveillance may aggregate and contribute to the burden of unfolded proteins. Some subset of this large pool of newly identified short peptides may play cellular roles that we have yet to discover, akin to the important roles recently shown for the 11- and 32-amino-acid peptides synthesized from the polished rice and sarcocolamin loci in Drosophila and the 58-amino-acid peptide encoded by the zebrafish toddler gene (Kondo et al., 2010; Magny et al., 2013; Pauli et al., 2014).

All RNA sequences subject to translation will experience selection against encoding proteins with detrimental impact on the cell or on the organism. These benign proteins may occasionally provide an adaptive molecular function; for example, a surprisingly large fraction (~20%) of random nucleotide sequences encode functional secretion signals (Kaiser et al., 1987). Further evolution may refine their expression, folding, and activity, ultimately giving rise to the birth of a new gene (Carvunis et al., 2012; Reinhardt et al., 2013).

Regardless of their original cellular role, degraded proteins are the substrates for antigens presented to the cellular immune system, and proteins synthesized by noncanonical translation may be shunted preferentially for degradation and presentation as antigens, expanding the range of epitopes displayed by virus infected or transformed (Yewdell, 2011). The apparent elevation of noncanonical translation in stress could aid the body in detecting these pathological cells, and differences in translation between normal and transformed cells could yield cancer-specific antigens for immunomodulatory therapy (Mellman et al., 2011). The same processes producing cryptic viral and tumor antigens could also expose cryptic self-antigens that could initiate or sustain an autoimmune response.

In summary, translation of noncoding RNA has the potential to impact the cell directly and to constrain the evolution of genomic sequences. A better understanding of these molecular and evolutionary implications relies, first, on a reliable means for unbiased detection of translation. Ribosome profiling provides a starting point for exploring the role of the translational apparatus in truly noncoding RNAs as well as revealing novel short, functional proteins and offering a window into the murky gradations in between.

**EXPERIMENTAL PROCEDURES**

**Ribosome Footprinting**

Embryonic day 14 mESCs were pretreated with cycloheximide (100 µg/ml) or emetine (50 µg/ml) for 1 min as indicated, followed by detergent lysis and ribosome footprinting by RNase I digestion (Ingolia et al., 2012). Deep sequencing libraries were generated from 26–34 nt footprint fragments (Ingolia et al., 2012) and sequenced on an Illumina HiSeq.

**Ribosome Affinity Purification**

The ribosome affinity tag construct comprised human ribosomal protein L1 fused to the biotin acceptor peptide (Beckett et al., 1999; de Boer et al., 2003), coexpressed with a biotin ligase using a 2A peptide (de Felipe et al., 2006), as a stable transgene in HEK293 cells using the Flp-In system (Invitrogen). Yeast lysates were prepared as described (Ingolia, 2010). Following nuclease digestion, lysates were loaded onto a SepharylS-G400 gel filtration spin column (Boca Scientific) and the flowthrough was collected. One aliquot of flowthrough was bound to streptavidin-coated magnetic beads (Invitrogen), and RNA was recovered by Trizol extraction directly from beads; another aliquot was used directly for library generation following Trizol extraction. Extracted RNA was converted into deep sequencing libraries.

**Footprint Sequence Alignment**

Footprint sequences were trimmed to remove 3' adaper sequence and aligned using TopHat v2.0.7 (Kim et al., 2013) with Bowtie v0.12.9.0 and samtools v0.1.18.0. The composite reference genomes comprised either the mm10 mouse genome with Ensembl GRCm38.72 transcripts or the human hg19 genome with Gencode v17 transcripts (Harwood et al., 2012), supplemented with the yeast genome with de novo transcript annotations (Brag et al., 2012). Alignments were filtered to remove those containing more than one mismatch.

**Footprint Sequence Data Analysis**

Footprints were assigned to specific A site nucleotide positions ~15 bases from their 5' ends, depending on the exact fragment length, as described previously. Reads assigned between 15 nucleotides before the start codon and 45 nucleotides after the start codon were excluded, as were all reads falling after the position 15 nucleotides upstream of the stop codon. All footprint data analysis was implemented in R/Bioconductor and is provided in a format allowing the direct reproduction of the analyses presented here.
We used our previously published (Stern-Ginossar et al., 2012), simplified approach to detect sites of AUG-mediated initiation in harringtonine-treated mESCs. We identified all AUG codons and selected harringtonine peaks by finding codons where A site occupancy on the +1 codon (i.e., AUG in the P site) as greater than occupancy on the +2 codon and greater than the sum of occupancy on the −1 and the 0 codon in both replicates. Among these AUG harringtonine peaks, we then selected the highest footprint occupancy on the +1 codon.

We computed the footprint A site occupancy at all codons on the transcript (not restricted to AUG codons with a harringtonine peak) and found the rank of the candidate initiation site relative to all other codons.

We also indexed all AUG codons on the transcript, starting from the 5′ end, and found the candidate initiation site among all AUG codons on the transcript.

Fragment Length Organization Similarity Score
The FLOSS was computed from a histogram of read lengths for footprints on a transcript or reading frame. A reference histogram was produced using raw counts on all annotated protein-coding transcript, excluding those whose gene overlapped a gene annotated as noncoding. The FLOSS was defined as

\[ 0.5 \times \sum_{l=35}^{34} f(l) - f_{ref}(l), \]

where \( f(l) \) is the fraction of reads at length \( l \) in the transcript histogram and \( f_{ref}(l) \) is the corresponding fraction in the reference histogram. The FLOSS cutoff score, as a function of the total number of reads, was counted from a rolling window of individual annotated genes and the computing of the upper extreme outlier cutoff for each window.

Yeast Western Blotting and Microscopy
Novel ORFs were tagged with C-terminal GFP fusions by the Pringle method (Longtine et al., 1998). Samples were collected by trichloroacetic acid precipitation and subjected to western blotting (mouse anti-GFP antibody, M.R.W. and S.E.J.). We were also collected for microscopy, which was performed on a Zeiss Axioskop equipped for time-lapse imaging.

We also indexed all AUG codons on the transcript, starting from the 5′ end, and found the candidate initiation site among all AUG codons on the transcript.

T Cell Response Assays
T cell responses were measured using ELISPOT plates (EBioscience) were prepared, coated, and blocked, and clear cells were isolated by Lymphoprep (Axis-Shield) and depleted of either T cells were plated at 3.0 × 10⁵ cells per well in RPMI-10.

ACCESSION NUMBERS
The NCBI GEO accession number for the ribosome profiling data reported in this paper is GSE60095.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, four figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2014.07.045.

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Figure S1. Classification of individual IncRNAs and 5' UTRs using the FLOSS. Related to Figure 1. (A, B) Fragment length analysis of mESC nuclear coding sequences showing a score cutoff based on the extreme outlier threshold of 3 x interquartile range over the 3rd quartile along a rolling 200 gene window. Individual (A) IncRNAs and (B) 5' UTRs are classified according to the FLOSS cutoff.
Figure S2. Bootstrap analysis of drug-dependent fragment length shifts. Related to Figure 2. The actual shift in cumulative length distribution between cycloheximide and emetine is shown, along with the distribution of shifts in bootstrap trials (N = 10,000). Bootstrapping was performed by assembling two sets of transcripts through selecting (with replacement) individual transcripts from a combined set of cycloheximide- and emetine-treated transcript data without regard for their drug treatment condition. Length distribution was computed for each of these bootstrapped transcript sets, and the length shift between the two randomly selected transcript sets was used as the result for that individual bootstrap trial.
Figure S3. FLOSS analysis detects snoRNA-derived background that co-purifies with the ribosome. Related to Figure 3. (A) FLOSS analysis distinguishes snoRNA-derived background from true ribosome footprints. (B, C) SnoRNAs are substantially retained during ribosome affinity purification. (D) FLOSS analysis confirms that nearly all 5’ UTRs resemble coding sequences in total HEK cell ribosome profiling.
Figure S4. Antigen responses to novel CMV ORFs in CMV-positive but not CMV-negative donors. Related to Figure 6. Quilt plot summary of ELISPOT assays of four CMV positive donors and three CMV negative control.
Table S1. Fragment length analysis on mouse lncRNAs. Related to Figure 1.

Fragment length analysis data (read counts, FLOSS values, and classification relative to typical protein-coding genes) are shown for each mouse lncRNA transcript, along with genome annotation data.
Table S2. Affinity purification and fragment length analysis on human lncRNAs. 
Related to Figure 3. Affinity purification data (total input read counts and affinity-purified bound read counts) and fragment length analysis data (read counts and FLOSS values in total input profiling data, and classification relative to typical protein-coding genes) are shown for human lncRNAs with aligned footprint reads, along with genome annotation data for those transcripts. Fragment length analysis read counts include all aligned reads regardless of their length, whereas read counts for expression comparison include only reads in the 26 nt – 31 nt size range corresponding to ribosome footprints in this sample.
Table S3. Initiation site detection and reading frame bias on human lncRNAs.

Related to Figure 4. Fragment length analysis data from each mouse lncRNA transcript are provided, as in Table S1, accompanied by data regarding the detection of harringtonine-marked AUG initiation sites and the bias of ribosome footprints for the downstream reading frames.
EXTENDED EXPERIMENTAL PROCEDURES

Mouse ES Cell Footprinting

Cell Growth and Lysis
E14 mouse ES cells were seeded at 4 x 10^4 cells per cm^2 in gelatin-coated tissue culture dishes in ESGRO Complete medium (Millipore), which was changed after 24 hours. Cells were grown at 37º C in 5% CO_2. At 48 hours, cells were dissociated with ESGRO Accutase (Millipore), counted, and re-plated at 4 x 10^3 cells per cm^2 in gelatin-coated tissue culture dishes in ESGRO Complete medium. Each profiling sample was prepared from one 10 cm dish of ES cells plated 48 hours prior to harvesting. Immediately prior to harvesting, cycloheximide (100 µg / ml final from a 100 mg / ml stock in DMSO) or emetine (50 µg / ml final from a 50 mg / ml stock) was added by withdrawing ~1 ml of media, mixing it with the drug stock, and returning this to the dish. Drug treatment was conducted for 1 minute in an incubator. Dishes were removed from the incubator, media was aspirated, and the dish was placed on ice while adherent cells were washed once in 10 ml ice-cold 1 x PBS containing drug. The wash was aspirated thoroughly and 400 µl lysis buffer (ribosome buffer (see below) + 0.5% Triton X-100, 25 U / ml Turbo DNase (Life Technologies)) was dripped onto the dish. The dish was tilted slightly and cells in lysis buffer were scraped and then pipetted off, yielding ~550 µl lysate per dish. Lysate was transferred to a clean non-stick RNase-free tube and pipetted extensively, then incubated 10 min. on ice. Lysate was then triturated several times through a 26-gauge needle and debris was pelleted by centrifugation for 10 minutes at 20,000 x g, 4º C. The supernatant was recovered. Ribosome buffer is 20 mM Tris pH 7.4, 150 mM NaCl, 5 mM MgCl_2, 1 mM DTT, and 100 µg / ml cycloheximide.

Ribosome Footprinting
RNA concentrations in mouse ES cell lysates were estimated using the Quant-iT RNA Assay kit (Molecular Probes) with fluorescence measurements on a SpectraMax M5 (Molecular Devices). The RNA in 2.0 µl of a 5-fold dilution was measured at 60.6 ng (Chx) or 56.8 ng (Emet), indicating an undiluted RNA concentration of ~0.15 µg / µl in the lysate. A lysate volume containing 56.8 µg RNA was diluted to 400 µl with lysis buffer and 12.0 µl of a diluted yeast lysate, prepared as described (Ingolia, 2010), was added. Ribosome footprinting was performed by adding 2.0 µl E. coli RNase I (10 U /µl; Epicentre) and incubating 45 min. on a room temperature nutator. Digestion was stopped by the addition of 10.0 µl SuperAseIn (20 U /µl; Life Technologies) and samples were placed on ice.

Total Ribosome Recovery
Mouse ES cell footprinting reactions were loaded into 13x51 mm thick-walled ultracentrifuge tubes (Beckman Coulter) and an 0.90 ml sucrose cushion (1 M sucrose in ribosome buffer) was underlaid. Ribosomes were precipitated by ultracentrifugation for 4 hours at 70,000 rpm, 4º C in a TLA100.4 rotor. The supernatant was discarded and a glassy ribosome pellet was recovered. The ribosome pellet was dissolved in 700 µl QIAzol (Qiagen) and total RNA including small (< 200 nt) RNAs was purified from the dissolved pellet using the miRNEasy Micro Kit (Qiagen) according to the manufacturer’s
instructions. RNA was eluted in 100 µl total volume of nuclease-free water, yielding ~600 ng / µl RNA.

Footprint Library Generation
This RNA was precipitated by adding 1.0 µl GlycoBlue (Life Technologies), 12 µl 3M sodium acetate, and 150 µl isopropanol, followed by overnight precipitation on dry ice. Precipitated RNA was recovered by centrifugation for 30 minutes at 20,000 x g, 4º C, the supernatant was removed, and the RNA pellet was air dried for ~15 minutes. The RNA was then resuspended in 5 µl 10 mM Tris pH 8 and 5 µl denaturing loading dye (98% formamide, 10 mM EDTA, 300 µg / ml bromphenol blue) and separated by electrophoresis on a 15% polyacrylamide denaturing TBE-Urea gel (Life Technologies) at 200 V for 65 minutes. The gel was stained for 3 minutes in 1x Sybr Gold (Life Technologies) and visualized on a blue light transilluminator (Clare Chemical). A gel region between 26 nt and 33 nt was excised based on the migration of marker RNA oligos. The gel slice was transferred to a non-stick nuclease-free microfuge tube and 400 µl of RNA gel extraction buffer (300 mM sodium acetate pH 5.5, 1 mM EDTA, 0.25% SDS) was added. The gel slice was frozen on dry ice for 30 minutes and then transferred to a room temperature rocker overnight to extract RNA. Gel extraction buffer was recovered from the elution, 1.5 µl GlycoBlue was added, followed by 500 µl isopropanol, and RNA was precipitated and recovered as described above.

Size selected RNA was resuspended in 10 µl 10 mM Tris pH 8 and transferred to a new non-stick tube with 33 µl nuclease-free water. The same was denatured 90 s at 80º C and then equilibrated at 37º C. Dephosphorylation was carried out by adding 5.0 µl 10x T4 polynucleotide kinase buffer (New England Biolabs), 1.0 µl SuperAseIn, and 1.0 µl T4 polynucleotide kinase (New England Biolabs) and incubating 1 hour at 37º C. The enzyme was inactivated by heating for 10 min. at 70º C, and then RNA was precipitated by adding 39 µl water, 1.0 µl GlycoBlue, and 10 µl 3M sodium acetate pH 5.5 to each reaction, followed by 150 µl isopropanol. Precipitation and RNA recovery was carried out as described above.

Dephosphorylated RNA was resuspended in 8.5 µl 10 mM Tris pH 8 and 1.5 µl preadenylated linker at 0.5 µg / µl (AIR adenylated linker A, Bioo Scientific) was added. This sample was denatured 90 s at 80º C and returned to room temperature. 2.0 µl 10x T4 RNA ligase buffer (New England Biolabs), 6.0 µl 50% polyethylene glycol, 1.0 µl SuperAseIn, and 1.0 µl T4 Rnl2(tr) (New England Biolabs) was added and mixed well. Ligation was carried out for 3 hours at room temperature. RNA was precipitated by the addition of 156 µl water, 20 µl 3M sodium acetate pH 5.5, 2.0 µl GlycoBlue, followed by 300 µl isopropanol, followed by chilling and centrifugation as described above. Ligation products were resuspended in 5.0 µl 10 mM Tris pH 8 and 5.0 µl 2x denaturing loading dye was added. Samples were separated by electrophoresis on a 15% polyacrylamide denaturing TBE-Urea gel and ligation products were excised. RNA was recovered from excised gel slices as described above. Gel-purified ligation products were resuspended in 10.0 µl 10 mM Tris pH 8 and 2.0 µl reverse transcription primer at 1.25 µM was added. Nucleic acid samples were denatured 90 s at 80º C and then placed on ice. Reverse transcription reactions were set up with 4.0
µl 5x first-strand buffer, 1.0 µl dNTPs 10 mM each, 1.0 µl 0.1 M DTT, and 1.0 µl SuperAseIn, followed by 1.0 µl M-MuLV reverse transcriptase (New England Biolabs), and mixed well. Reverse transcription was carried out for 30 min. at 48ºC and then RNA was hydrolyzed by adding 2.2 µl 1N NaOH and heating 20 min. at 98ºC. Reverse transcription products were purified using an RNA Clean and Concentrator-5 column (Zymo Research) with elution into 7.0 µl 10 mM Tris pH 8. The first-strand cDNA was mixed with 7.0 µl 2x denaturing loading dye and separated by electrophoresis on a 15% polyacrylamide denaturing TBE-Urea gel. The reverse transcription product band was excised and DNA was extracted as described above, except using a DNA extraction buffer (300 mM NaCl, 10 mM Tris pH 8, 1 mM EDTA) in place of RNA extraction buffer.

First-strand cDNA was resuspended in 5.0 µl 10 mM Tris pH 8 and transferred to a new tube along with 1.0 µl 1 mM ATP, 1.0 µl 50 mM MnCl₂, 2.0 µl 10x CircLigase buffer, 10.0 µl water, and 1.0 µl CircLigase 2. Circularization was carried out for 60 min. at 60ºC and then the enzyme was inactivated 10 min. at 80ºC. Circles containing rRNA fragments were depleted by mixing 5.0 µl circularization reaction with 1.0 µl depletion oligo pool (10 µM each), 1.0 µl 20x SSC, and 3.0 µl water. Samples were heated to 100ºC for 90 s and then annealed at 0.1ºC / s to 37ºC, followed by a 15 minute incubation at 37ºC. MyOne C1 DynaBeads (Life Technologies), 25 µl per sample, were prepared by washing three times in 1x bind/wash buffer (1M NaCl, 0.5 mM EDTA, 2.5 mM Tris pH 7.5, 0.1 % Triton X-100) and resuspended in 10 µl 2x bind/wash buffer per sample. Annealing reactions were combined with 10 µl washed beads and incubated 15 min. at 37ºC with agitation. Beads were collected by placing the tubes in a magnetic rack and ~17.5 µl supernatant was recovered. DNA was recovered from the supernatant by adding 1 µl GlycoBlue, 6 µl 5M NaCl, 75 µl water, and 150 µl isopropanol, followed by precipitation as described above.

Depleted circles were resuspended in 10.0 µl 10 mM Tris pH 8 and used to prepare PCR amplification reactions with 12.0 µl 5x HF buffer, 1.2 µl dNTPs, 3.0 µl forward primer at 10 µM, 3.0 µl reverse indexed primer at 10 µM, 30 µl water, and 0.6 µl Phusion polymerase (New England Biolabs). Aliquots of 16.7 µl were amplified by thermal cycling (denaturation for 30s at 98ºC followed by cycles of 10 s at 98ºC, 10 s at 65ºC, and 5 s at 72ºC). Samples were amplified for 10 cycles of PCR and then separated by electrophoresis on an 8% non-denaturing polyacrylamide gel. The sequencing library band (~175 bp) was excised and the dsDNA library was extracted as described above. The library was validated and quantified using the High Sensitivity DNA kit on the BioAnalyzer 2000 (Agilent).

**Library Generation Primers**
Reverse transcription primer (NI-NI-9):
[Phos]AGATCGGAAGAGCACGTGTAGGGAAAGAGTGTAGATCTCGGTCGGTCTCAG\[Sp-C18\]CACTCA[Sp-C18]TTCAGACGTGTCTCTTCCGATCTATGGATGATGCCCTCAG
Forward primer (NI-NI-2): AATGATACGCACCTTATGCACAGATGTCTACACT
Reverse index primer for Chx sample (NI-NI-6):
CAAGCAGAAGACGGCATACGAGATATGCTGGTGACTGGAGTTTCAGACGTGTGCTCTTCCG
Reverse index primer for Emet sample (NI-NI-7):
CAAGCAGAAGACGGCATACGAGATACGTCGGTGACTGGAGTTTCAGACGTGTGCTCTTCCG

Subtraction oligos:
NI-NI-21 [BioTEG]tcccggggtacgcctgtgtgag
NI-NI-23 [BioTEG]gggcccaagtctctctcgatec
NI-NI-24 [BioTEG]gctctccagtccgcgcgag
NI-NI-63 [BioTEG]ACTCGCCGAATCCCCGGGCCGA
NI-NI-64 [BioTEG]GCAGACCGGCTCCGGGACGGCT
NI-NI-65 [BioTEG]TTCACTGACCCGGTGAGGCGG
NI-NI-356 [BioTEG]CTCGGTTGGCCYCGGATAGCCGG
NI-NI-357 [BioTEG]CTCGCTTCTGGCGCCAAGCGCCCG
NI-MH-1 [BioTEG]GAA GCCGAGCGCACGGGGTCGG
NI-MH-2 [BioTEG]GTCGGGGTTTCGTACGTAGCAGAGC
NI-MH-3 [BioTEG]CGATCTATTGAAAGTCAGCCCTCG
NI-MH-4 [BioTEG]GACTCTAGATACCTGGGGCGGC


Ribosome Affinity Purification

Ribosome Affinity Tagging

The ribosome affinity tag construct comprised human ribosomal protein L1 fused to the biotin acceptor peptide (Beckett et al., 1999; de Boer et al., 2003) with a 27 amino acid flexible linker, along with a human codon optimized biotin ligase co-expressed and separated by a 2A ribosome skipping peptide (de Felipe et al., 2006). This RPL1-GS-TEVsite-Avi-T2A-hBirA fusion was transgenically expressed from the pNTI194 expression vector. This plasmid is derived from the pcDNA5/FRT expression vector (Invitrogen) with CMV promoter and BGH poly-(A) site. The human RPL1 sequence was amplified from human cell cDNA and fused translationally to a linker and affinity tag sequence encoding the polypeptide

GGSSGSGSGSGGGSSGSGGSENLYFOGLNDIFEAQKIEWHE, where the underlined amino acids comprise a TEV recognition site not used in this study, the bold peptide sequence comprises a biotinylation motif (Beckett et al., 1999; de Boer et al., 2003) for the E. coli birA biotin ligase, and the bold italic lysine is the biotinylated residue. The biotin acceptor peptide is followed by the T2A translational skipping peptide (de Felipe et al., 2006; Szymczak-Workman et al., 2012) and then by hBirA, a human codon optimized version of the E. coli birA331,825 biotin ligase with Ala34Glu and
Arg33Leu mutations that abolish DNA binding without affecting enzymatic activity (Buoncristiani et al., 1986).

Stable pNT194 expression was established by co-transfection into Flp-In HEK cells (Invitrogen) with pOG44, which expresses the Flp recombinase and selection for hygromycin resistance (150 µg/ml) according to the manufacturer’s instructions. HEK cells expressing the affinity-tagged ribosome were propagated in the presence of hygromycin (150 µg/ml; InvivoGen) according to standard procedures. For ribosome profiling experiments, cells were plated in 10 cm dishes in the absence of hygromycin and expanded to ~50% confluency. Growth media was supplemented with 1 mM biotin (Sigma) 16 hours prior to cell harvesting. Cell harvesting and lysate preparation were performed as described for mouse ES cells. Affinity-tagged HEK lysates were mixed with yeast lysate containing an equivalent amount of total RNA and this mixture was subjected to nuclease footprinting as described above. Following nuclease digestion, lysates were loaded onto a Sephacryl S-400 gel filtration spin column (Boca Scientific) and the flow-through was collected. One aliquot of flow-through was bound to streptavidin-coated magnetic beads (Invitrogen) and RNA was recovered by Trizol extraction directly from beads after washing. Another aliquot was used directly. Extracted RNA from affinity purification (bound) and ultracentrifugation (input) were subsequently converted into deep sequencing libraries according to the same procedures used for total ribosome footprint profiling.

<table>
<thead>
<tr>
<th>Cells</th>
<th>Treatment</th>
<th>Mapped Reads</th>
</tr>
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<tbody>
<tr>
<td>mES Cells</td>
<td>Cycloheximide</td>
<td>254,516,745</td>
</tr>
<tr>
<td></td>
<td>Emetine</td>
<td>170,514,739</td>
</tr>
<tr>
<td>HEK + Yeast</td>
<td>Input</td>
<td>26,071,571</td>
</tr>
<tr>
<td></td>
<td>Bound</td>
<td>14,723,220</td>
</tr>
<tr>
<td>mES Cells</td>
<td>Harringtonine 120s</td>
<td>33,355,973</td>
</tr>
<tr>
<td>(Ingolia et al. 2011)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Harringtonine 150s</td>
<td>47,320,855</td>
</tr>
</tbody>
</table>

**Fragment Length Organization Similarity Score**

The fragment length organization similarity score (FLOSS) was computed by calculating a histogram of read lengths for all footprints that aligned to a specific transcript or reading frame, collapsing those below 26 nt or above 34 nt into the 26nt – 34nt range used in physical fragment size selection. A reference histogram was produced by summing individual raw counts (without normalization) for each annotated nuclear protein-coding transcript, excluding those whose gene overlapped a gene annotated as non-coding. The FLOSS was defined as...
\[0.5 \times \sum_{l=26}^{34} \| f(l) - f_{\text{ref}}(l) \|\]

where \( f(l) \) is the fraction of reads at length \( l \) in the transcript histogram and \( f_{\text{ref}}(l) \) is the corresponding fraction in the reference histogram. The FLOSS cutoff, calculated as a function of the total number of reads in the transcript histogram, was established by considering a rolling window of individual annotated genes and computing the upper extreme outlier cutoff for each window using Tukey’s method (Q3 + 3*IQR, where Q3 is the 3\textsuperscript{rd} quartile and IQR is the interquartile range).

**T Cell Response Assays**

Peptides constructed as sequential 15 amino acid peptides with 10 amino acid overlap, spanning ORFL2C, ORFL3C, ORFL4C, ORFL5C, ORFL7C, ORFL8C, ORFL293C and ORFL283W were obtained from JPT Peptide Technologies. Peptides were reconstituted to give a storage concentration of 40 mg/ml. Individual peptides were further diluted in RPMI 1640 to create a stock of 1 mg/ml. Peptide pools were made and were constructed to give 2 µg/ml of each individual peptide.

Blood was collected in heparin sodium (100 IU/ml), diluted 1:2 with RPMI-1640 containing no serum supplemented with 100,000 IU/ml penicillin, 100 mg/ml streptomycin, and 2 mmol/ml L-glutamine (RPMI-wash). Peripheral blood mononuclear cells (PBMC) were isolated by Lymphoprep (Axis- Shield, Norway) centrifuged at 800 g for 15 minutes.

ELISPOT plates were prepared, coated and blocked according to manufacturer’s instruction (EBioscience). PBMC directly ex vivo, previously frozen, or depleted of either CD4+ or CD8+ T cells by magnetic activated cell sorting (MACS), were plated 3.0×105 cells in 100 µl RPMI-10 per well (of a 96 well Multiscreen IP sterile plate (Millipore, UK)). Plates were incubated for 48 hours at 37°C 5% CO2, and developed according to manufacturer's instruction.

PBMC were depleted of either CD4+ or CD8+ T cells by MACS using either anti-CD4+ or anti-CD8+ direct beads (Miltenyi, U.K.), according to manufacturer's instructions and separated on LS columns (Miltenyi, U.K.). Efficiency of depletion was determined by staining cells with either anti-CD4 or anti-CD8 antibodies and analysed by flow cytometry. Depletions performed in this manner resulted in 0.1–0.8% CD4+ cells and 0.3–0.8% CD8+ cells, respectively.
Supplemental References


