Active Ribosome Profiling with RiboLace

Graphical Abstract

Highlights
- RiboLace isolates ribosomes in active translation by antibody-free and tag-free pull-down
- RiboLace works reliably with low amounts of input material in vitro and in vivo
- RiboLace provides positional data of active ribosomes with nucleotide resolution
- RiboLace estimates translation levels and predicts protein levels with accuracy

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In Brief
Clamer et al. present RiboLace, a method for isolating active ribosomes and associated proteins, intact mRNAs, or ribosome-protected fragments. RiboLace accurately quantifies translation levels, providing positional data of active ribosomes with nucleotide resolution. Requiring lower input than current ribosome profiling protocols, RiboLace can be used with challenging biological samples.
Active Ribosome Profiling with RiboLace

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SUMMARY

Ribosome profiling, or Ribo-seq, is based on large-scale sequencing of RNA fragments protected from nuclease digestion by ribosomes. Thanks to its unique ability to provide positional information about ribosomes flowing along transcripts, this method can be used to shed light on mechanistic aspects of translation. However, current Ribo-seq approaches lack the ability to distinguish between fragments protected by either ribosomes in active translation or inactive ribosomes. To overcome this possible limitation, we developed RiboLace, a method based on an original puromycin-containing molecule capable of isolating active ribosomes by means of an antibody-free and tag-free pull-down approach. RiboLace is fast, works reliably with low amounts of input material, and can be easily and rapidly applied both in vitro and in vivo, thereby generating a global snapshot of active ribosome footprints at single nucleotide resolution.

INTRODUCTION

The process of protein synthesis is a core regulator of numerous critical physiological pathways ranging from cell growth (Goyer et al., 1993) and development (Kondrashov et al., 2011; Xue et al., 2015) to immune response (Piccirillo et al., 2014). Local protein synthesis in neurons (Jung et al., 2014) also plays fundamental roles in memory formation (Fioriti et al., 2015; Kandel et al., 2014; Martin et al., 1997) and synaptic plasticity (McCamphill et al., 2015). Hence, dysregulation of translation is a major driver of important pathologies such as cancer (Bhat et al., 2015; Topisirovic and Sonenberg, 2015) and neurodegenerative diseases (Bernabò et al., 2017; Darnell et al., 2011).

During the last few years, methodological approaches such as ribosome profiling (Ribo-seq) (Ingolia et al., 2009) have contributed considerable insights into the translation process. Ribo-seq has been largely used to identify translated RNAs (both coding and, unexpectedly, non-coding), map upstream open reading frames (ORFs), and estimate translation levels in different biological conditions. Ribo-seq has been used to estimate translation efficiencies and “protein synthesis levels” (Ingolia et al., 2014; Li et al., 2014) in a variety of organisms, from prokaryotes (Li et al., 2014) to yeast (Ingolia et al., 2009), Caenorhabditis elegans (Stadler et al., 2012), zebrafish (Bazzini et al., 2014; Chew et al., 2013), plants (Juntawong et al., 2014), mouse (Ingolia et al., 2011), and human (Fritsch et al., 2012; Lee et al., 2012; Liu et al., 2013).

Despite its unquestionable discrimination power and wide applicability, Ribo-seq still faces a number of challenges and presents some limitations. For example, translationally inactive mRNAs can be sequestered into ribonucleoprotein particles (mRNPs) and stalled or paused in polysomes as a consequence of physiological surveillance mechanisms, stress stimuli, and regulatory mechanisms (Yordanova et al., 2018). The contribution of these phenomena in multicellular organisms is particularly important in specific tissues, and it has been shown to occur especially in neurons (Chapman and Walter, 1997; Darnell et al., 2011; Doma and Parker, 2006; Graber et al., 2013; Higashi et al., 2013). As such, while inactive ribosomes unbound to transcripts do not present a problem, Ribo-seq does not necessarily discriminate “authentic” protected footprints of translating polysomes from RNA fragments protected by inactive or stalled ribosomes, leading to possible misinterpretations of translation occupancy profiles. Therefore, to generate optimal insight into the translation process, Ribo-seq is still open to further optimization and refinements, incorporating aspects from the laboratory bench to data analysis (Aeschimann et al., 2015).

Here, we present RiboLace, a methodological approach to study active translation based on a newly developed reagent: a puromycin analog molecule. The aim of our study was to purify active ribosomes by immobilizing puromycylated ribosomes frozen on the transcript by the chain elongation inhibitor cycloheximide, which impedes the dissociation of ribosomal subunits (David et al., 2012). Our data show that RiboLace is useful for ribosome purification and the co-purification of associated proteins, intact mRNAs, and nuclease-protected footprints using
40 times less material than classical ribosome profiling. RiboLace provides a valuable technique, with clear applications in vitro and in vivo.

RESULTS

Design and Synthesis of an Analog of Puromycin

Puromycin is an aminonucleoside antibiotic able to bind the ribosome and the nascent peptide chain, causing ribosome disassembly and disruption of protein synthesis (Nissen et al., 2000; Welch et al., 1995; Wilson, 2014; Yarmolinsky and Haba, 1959). It has been extensively used to quantify global protein synthesis, taking advantage of radioactive (Gambetti et al., 1972) and biotinylated molecules (Aviner et al., 2014) or anti-puromycin antibodies (Schmidt et al., 2009). Leveraging its ability to maintain contact with the ribosome (Kukhanova et al., 1979; Odom et al., 1990; Pestka et al., 1972; Schmeing et al., 2002), puromycin has also been used to covalently bind an mRNA to the corresponding protein during its synthesis (Biyani et al., 2006; Roberts and Szostak, 1997). In addition, puromycin can be modified to create cell-permeable analogs suitable for direct and in situ imaging of newly synthesized proteins (Ge et al., 2016; Starck et al., 2004). These methods require the irreversible reaction of the α-amino group of puromycin with the carbon on its carbonyl group, acylating the 3′ hydroxyl group of the peptidyl-tRNA buried in the P-site of the ribosome.

Motivated by the evidence that molecules containing puromycin modified at its α-amino group can bind to the large subunit of the active ribosome (Schmeing et al., 2002), we covalently coupled puromycin to a biotin moiety through two 2,2′-ethylene-dioxy-bis-ethylamine units to obtain a compound that is still able to bind ribosomes by mimicking the 3′-tRNA in the acceptor site (A-site). We synthesized the molecule (Figures 1A, 1B, and S1A), characterized it by nuclear magnetic resonance (NMR) and liquid chromatography-electrospray ionization-mass spectrometry (LC-ESI-MS) (Figures S1B and S1C), and called it 3P. We verified the activity of the biotin moiety, taking advantage of its absorbance spectrum, and tested the binding on polystyrene or agarose beads.

Figure 1. An Analog of Puromycin Inhibits Translation and Can Be Used for Functionalization of Agarose and Polystyrene Beads

(A) Schematic representation of the 3P structure. Biotin is the residue binding the surface; two 2,2′-ethylene-dioxy-bis-ethylamine units form the “L” linker, and puromycin is the residue binding the ribosome. Steps 1–3 refer to the chemical synthesis procedure. CDI, N,N′-dicyclohexylcarbodiimide, jeffamine, 2,2′-ethylene-dioxy-bis-ethylamine; NHS, N-hydroxysuccinimide.

(B) Chemical formula of the puromycin analog 3P.

(C) Depletion assay of 3P with streptavidin-coated agarose (purple) and polystyrene beads (black). Absorbance of the supernatant at 275 nm is measured after addition of streptavidin-coated magnetic beads to 100 pmol 3P. Data represent the mean of triplicate experiments. The gray bar marks the quantity of beads used in all of the experiments for each sample.

(D) Comparison between the efficiency of puromycin (left) and 3P (right) to inhibit the protein production of firefly luciferase. ε-Labeled biotinylated lysine-tRNA is used to monitor the protein production by SDS-PAGE (top). Histograms represent the change in protein production after the addition of different concentrations of puromycin or 3P with respect to the control. Error bars represent SDs calculated from triplicate experiments; n = 3; t test; *p < 0.05.
agarose beads. We observed that the biotin group allows the binding of 3P to commercially available streptavidin beads (Figure 1C).

To demonstrate that the 3P molecule maintains an inhibitory effect on translation, we compared its effects to that exerted by puromycin, using a eukaryotic in vitro cell-free transcription-translation system and the firefly luciferase as a reporter gene. We monitored total protein production by SDS-PAGE (Figure 1D) and luminescence assay (Figure S1D) in the presence of puromycin and 3P at different concentrations. As expected, puromycin induced conspicuous decay of protein production at nanomolar concentrations. In the case of 3P, we observed a decreased level of translation, which reached >70% of inhibition at concentrations >1 μM (Figure 1D). We concluded that even if it has slightly lower efficacy than puromycin, 3P can robustly inhibit eukaryotic translation in vitro, interfering with ribosome function, and can therefore be used to produce functionalized beads.

3P-Functionalized Beads Capture mRNAs in Active Translation In Vitro

Figure 2. 3P-Functionalized Beads Can Capture mRNAs in Active Translation In Vitro

(A) Experimental design: 3P beads are used to pull down transcripts in a cell-free in vitro transcription-translation system. Briefly, from step 1 to step 5: (1) plasmids were added to the IVTT reaction mix, and the reaction was stopped by the addition of cycloheximide after 40 min; (2) beads were functionalized with 3P; (3) the IVTT reaction was incubated with 3P beads for 1 hr on a wheel at 2 rpm and 4°C; (4) beads were washed to remove unspecific binding; (4a) RNA was extracted and treated with DNase I to avoid possible DNA contaminations (4b); and (5) RNA samples were analyzed by RT-qPCR to detect the reporter gene.

(B) Immunoblotting of total EGFP protein at different incubation times, without (+) harringtonine (harr) or with (+) harringtonine (2 μg/mL for 3 min). Immunoblotting showing the comparison between total EGFP expression from the pPR-IBA2 plasmid (blue) and EGFP expressed from the pBluescript II KS+ plasmid (red).

(C) EGFP RNA enrichment on RiboLace in the presence or absence of harringtonine (harr) during time in different translational conditions: low (red pBluescript II KS+) and high translation efficiency (blue, pPR-IBA2 plasmid). At right, the total RNA content without (-) or with harringtonine (harr) in the two IVTT reactions; incubation time 25 min; t test; *p < 0.05.

(D) Experimental protocol for IVTT reaction with the firefly luciferase (luc) reporter with harringtonine (2 μg/mL) and high translation efficiency (blue, pPR-IBA2 plasmid). At right, the total RNA content without (-) or with harringtonine (harr) in the two IVTT reactions; incubation time 25 min; t test; *p < 0.05. All of the data presented represent the mean values and SDs of three to five independent experiments.

(E) Fold change values relative to the total amount of transcript captured by 3P beads were compared to the control beads (mP), henceforth referred to as the “enrichment.” t test; *p < 0.05. All of the data presented represent the mean values and SDs of three to five independent experiments.

First, we monitored the ability of 3P-functionalized magnetic beads (3P beads) to purify transcripts of reporter genes with different levels of protein expression in in vitro translation systems. To this end, we developed the following protocol (Figure 2A): (1) in vitro transcription translation (IVTT) reaction to induce the production of the reporter protein, (2) functionalization of beads with 3P, (3) stopping reaction with the translation inhibitor cycloheximide and incubation of 3P beads and control beads functionalized with a biotin-glycol conjugate (mP beads), (4) washing of beads, and (5) extraction of RNA associated with the beads for downstream analyses. In parallel, the production of the protein was followed by detection of whole protein extracts.

We applied this protocol to study the ability of our 3P functionalized beads to purify mRNAs associated with high translational states. We took advantage of two reporter plasmids characterized by different translational efficiencies. The EGFP reporter genes showed differential efficiency in protein production, depending on the expression vector used (pPR-IBA2) (Arosio et al., 2015).
specifically activate protein synthesis, we rescued cells from

1100

subunits of the ribosome, respectively. After immunoblotting

Calnexin is a chaperone protein in the endoplasmic

ribosums, therefore monitoring the translational state of cells.

Thus, to demonstrate that this result was not dependent on

The enrichment can be observed in the absence of transcrip-

tional changes (Figure 2C, right) and with respect to the unspec-

cific binding on control beads (Figure S2).

RiboLace Captures Active Ribosomes and Associated

Next, we wanted to establish whether RiboLace was capable of

isolating ribosomes and mRNAs under active translation from

more complex systems than in vitro mixtures. We used RiboLace

on whole cellular lysates under different translational states

and monitored its efficiency to capture mRNAs and proteins

associated with ribosomes and polysomes in cellular lysates

(Figure 3A). We took advantage of established cellular stimuli

in order to induce cells into translationally active or inactive states. To

place translation, we used cell starvation and oxidative,

proteotoxic stress, heat shock, sodium arsenite). In all of the cases,

translation markers were reduced (Figure S3D). We then tested

RiboLace on a mouse motor neuron-like cell line, NSC-34, under

to further validate this finding, we applied other stress stimuli

known to elicit repression of global protein synthesis (e.g., pro-

teotoxic stress, heat shock, sodium arsenite). In all of the cases,

translational markers were reduced (Figure S3D). We then tested

RiboLace on a mouse motor neuron-like cell line, NSC-34, under

normal growth conditions. We observed an ~4-fold enrichment of

RPL26 and an ~4-fold enrichment of RPS6 with respect to

control beads (Figure S3E), demonstrating that RiboLace can

isolate ribosomes from both human and mouse cell lines.

Consistent with these results, the pelota protein (mammalian

ortholog of the yeast Dom34), known to promote the dissociation

of stalled ribosomes (Guydosh and Green, 2014; Pisareva et al.,

2011), was not enriched in RiboLace applied to control, starved,

or arsenite-treated lysates of MCF7 and HEK293T cells (Figures

3F and S3F, respectively).

These findings prompted us to investigate whether RiboLace

provides an improved estimation of protein level with respect
to the use of total RNA or polysomal RNA (Figure 4A). To this end, we compared at transcriptome-wide scale the proteome of MCF7 cells with all three RNA quantifications: total RNA levels (classical transcriptome), polysomal RNA levels (classical translome), and RiboLace. The levels of ~2,700 proteins were determined by MS and label-free quantification (LFQ). We report in Figure 4B (left) that RNA levels obtained using RiboLace displayed the highest correlation with protein levels (0.48), significantly improving the correlation obtained with polysomal RNA (0.41) and total RNA (0.18). It is worth mentioning that some translational changes may show up as a shift in the position of an mRNA within a sucrose density gradient rather than as a change in the fraction of the mRNA found in polysomes, thereby penalizing the sensitivity of polysome-associated RNA sequencing. Our results prove that RiboLace provides a reliable estimation of the translation state of cells.
To further validate these results, we isolated RNA from total RNA, polysomal RNA, and RiboLace from MCF7 cells before and after serum starvation or EGF stimulation (Figures S4A–S4F). As in control cells (Figure 4B, left), we observed in starved MCF7 cells a significantly higher correlation between protein levels and RiboLace levels with respect to polysomal and total RNA levels (Figure 4B, right). We next searched for genes that contributed the most to this difference in correlation.

Figure 4. Comparison of RiboLace to RNA-Seq or POL-Seq Reveals that RiboLace Is an Accurate Proxy of the Cellular Proteome

(A) Experimental design for comparing the global RNA repertoire of RNAs associated with RiboLace by next-generation sequencing, total RNA sequencing (RNA-seq), and polysomal sequencing (POL-seq) to the cellular proteome.

(B) Correlation analysis between the MCF7 proteome, determined by mass spectrometry, and total RNA, polysomal RNA, and RiboLace RNA, respectively, determined by deep sequencing. Pearson correlation values are displayed (error bars refer to the 95% confidence interval). The significance of the differences between Pearson coefficients was measured using Williams' test (**p < 0.01, ***p < 0.001).

(C) Single gene comparisons of protein (Prot), RiboLace (RL), polysomal RNA (Poly), and total RNA (Tot) levels in MCF7 control and starved cells (empty and filled bars, respectively). Data are represented as means ± SEMs (*p < 0.05, **p < 0.01, and ***p < 0.001, based on proteomics or next-generation sequencing [NGS] differential analysis). A total of 15 representative genes, with the most significant differences between RiboLace and conventional approaches, were chosen for display, showing either increase (top row), decrease (middle row), or no change (bottom row) at the protein level.

To further validate these results, we isolated RNA from total RNA, polysomal RNA, and RiboLace from MCF7 cells before and after serum starvation or EGF stimulation (Figures S4A–S4F). As in control cells (Figure 4B, left), we observed in starved MCF7 cells a significantly higher correlation between protein levels and RiboLace levels with respect to polysomal and total RNA levels (Figure 4B, right). We next searched for genes that contributed the most to this difference in correlation. Comparing the control and starved conditions, we considered cases in which significant protein changes were reported in
proteomics. We identified a population of 80 genes changing at the protein and mRNA levels in proteomics and RiboLace, respectively. The mRNA shifts of these 80 genes were either not detected or reversed at both polysomal and total RNA levels (Figure 4C, upper row for upregulated proteins and middle row for downregulated proteins).

Next, we considered a scenario in which the researcher would exclude unchanging proteins from the analysis of translation. We found a population of 201 genes showing no change in protein levels and no change in RiboLace levels, but showing significant changes in polysomal and optionally total RNA levels (Figure 4C, bottom row). For each of these populations, five representative genes showing the most significant differences between RiboLace and conventional approaches were chosen for display (Figure 4C). Further validations for the consistency between RiboLace and conventional approaches were chosen for display.

Overall, these results establish the important proof-of-concept that RiboLace can capture ribosomes under active translation and estimate protein levels and their variations with accuracy and reliability.

**In Vivo Active Ribosome Profiling Using RiboLace: Active Ribo-Seq**

Given the unique opportunity to purify active ribosomes from low amounts of input material, we next wanted to combine our purification strategy with Ribo-seq to capture active ribosome dynamics along transcripts, thereby improving ribosome profiling experiments. To facilitate this, we modified our original protocol by including an endonuclease digestion step and applying it to cellular and tissue lysates (Figure 5A).

First, we demonstrated that RiboLace can capture isolated ribosomes after endonuclease digestion, as shown by the enrichment of eEF1α, calnexin, RPL26, and RPS6 on RiboLace in HEK293T and HeLa cells (Figure 5B). Second, we confirmed that RiboLace was able to enrich ribosome-protected fragments (Figures S5A and S5B). Third, we applied RiboLace on as few as 15 μL of whole mouse brain ribosome-protected fragments (RPFs). In parallel, we performed ribosome profiling from polysomes, pre-purified using sucrose gradients (750 μL of total brain lysate, Ribo-seq; Figure 5C). After sequencing, we analyzed both RiboLace and Ribo-seq ribosome-protected fragments using the dedicated pipeline riboWaltz (Lauria et al., 2018) to obtain sub-codon information and identification of the trinucleotide periodicity. The distribution of read lengths highlighted a main population of ribosome-protected fragments at ~29 nt (Figure S5C) (Archer et al., 2016; Lareau et al., 2014). As expected for ribosome footprints, we observed an enrichment of signal from mapped reads along coding sequence regions in both RiboLace and Ribo-seq data (Figure 5D), demonstrating that RiboLace is indeed able to capture bona fide ribosomes. Occupancy meta-profiles, derived from the aggregation of signals on single genes, presented the typical trinucleotide periodicity of the ribosome P-site along coding sequences, which is suggestive of signal derived from ribosomes moving along transcripts (Figures 5E and 5F). The comparison between meta-profiles obtained with RiboLace and Ribo-seq highlights for our method an accumulation of ribosomes at the start codon and at the fifth codon, a feature that has previously been reported to be associated with a productive elongation phase of translation (Han et al., 2014).

To further reinforce our findings, we systematically compared position-specific profiles from RiboLace and standard Ribo-seq using an additional dataset from the widely used HEK293 human cell line (Figure S6A). According to Arava et al. (2003), in fully growing, actively translating cells, an overall decrease of ribosome density is expected along the transcripts. We therefore analyzed our data by calculating the ratio between the average number of P-sites on the coding sequence and the average number of P-sites on the first five codons for each transcript. The lower this value, the lower the ribosome density ratio along the transcript. By comparing the distributions obtained with RiboLace and Ribo-seq, RiboLace showed a stronger decrease in ribosome density along coding sequences than Ribo-seq, both in mouse brain and in HEK293 cells (Figure 6A), which is in line with active translation (Arava et al., 2003). Conversely, a comparison of codon-specific ribosome densities revealed a high correlation between RiboLace and Ribo-seq (0.98 in mouse brains and 0.93 in HEK293 cells), suggesting the absence of differences at this general level.

Finally, we compared gene-specific translation estimates obtained using RiboLace and Ribo-seq. In this way, we identified populations of genes whose translation signal was specifically enriched in either RiboLace or Ribo-seq (Figure 6B). This result shows that, for both case studies we generated, the two techniques significantly differ in translation estimates for hundreds of genes. Functional enrichment analysis of these populations suggested that RiboLace-enriched transcripts were more pertinent to the biological system under study (embryonic kidney cells and brain extracts, respectively) in comparison to Ribo-seq-enriched transcripts (Figure 6C). In addition, RiboLace-specific transcripts were enriched for translation-related genes in both datasets (Figure 6C).

These results demonstrate that RiboLace is capable of providing positional data of active ribosomes with nucleotide resolution, starting from much less material than available protocols, thereby facilitating reliable descriptions of bona fide translational events in vitro and in vivo.

**DISCUSSION**

During their lifetime in the cytoplasm, mRNAs are regularly stored, degraded, and transported, with only a fraction being actively translated to produce proteins at a specific point in time (Morisaki et al., 2016; Wu et al., 2016). We estimated this fraction to be between 80% and 60% of the total population, depending on the cell type (Figure 5E). All of these stages of the mRNA life cycle are governed by cis- and trans-factors that tightly regulate the uploading of mRNAs on polysomes and the subsequent production of proteins. To generate an improved understanding of these sophisticated and dynamic processes, different methodological approaches have been developed to determine, at the genome-wide scale, changes in RNA steady-state levels (e.g., RNA sequencing [RNA-seq]), their engagement with the translational machinery (e.g., Ribo-seq, polysomal profiling) (Ingolia et al., 2011; Tebaldi et al., 2012), and changes in protein production (e.g., stable isotope labeling by amino
acids in cell culture [SILAC], puromycin-associated nascent chain proteomics [PUNCH-P] (Aviner et al., 2014; Liu et al., 2017; Ong and Mann, 2006). Although Ribo-seq remains a complex technology that requires relatively large volumes of experimental material, it has been shown to be extremely powerful for identifying ORFs and translation initiation sites from (1) cell lysates or ribosome pellets; (2) purified polysomal fractions (Junta-wong et al., 2014); or, more recently, (3) tagged ribosomes (Jan et al., 2014; Shi et al., 2017). Unfortunately, however, the use of cell lysates and ribosome pellets often introduces unwanted background signals.

Here, we specifically designed RiboLace for ribosome profiling experiments, to facilitate understanding of ribosome dynamics along transcripts and to allow accurate estimates of translation levels based on active ribosome footprints. We sought to enhance Ribo-seq approaches by developing a...
molecule (3P) that facilitates the selective capture of ribosomes under active translation. We focused our attention on puromycin, the well-known structural analog of the 3’ end of aminoacyl-tRNA, and tethered the α-amino group to a biotinylated linker. We observed that 3P can interfere with eukaryotic translation in vitro. We used 3P-functionalized beads (RiboLace) to capture and enrich transcripts undergoing translation in eukaryotic in vitro and in vivo systems. We observed that the elongation factor eEF1α, a key protein involved in delivering tRNAs to the ribosome, was the most enriched protein on RiboLace in all of our experiments. This suggests that in the presence of cycloheximide treatment, 3P binding to the A-site of the ribosome in the not-rotated state of the ribosome (Ferguson et al., 2015; Lareau et al., 2014) is favored. Given the elongation speed of ~6 aa/s (Ingolia et al., 2011) and the duration of cycloheximide treatment, all ribosomes in a different phase of the elongation cycle at the beginning of the treatment (i.e., after peptide bond formation but before translocation) have more than enough time to move into the post-translocation cycle, be blocked by cycloheximide, and be captured by RiboLace.

Comparing RiboLace sequencing with proteomics on a transcriptome-wide scale, we obtained evidence to suggest that RiboLace is a powerful and reliable alternative to quantify the translation state of cells compared to standard transcriptome and translatome profiling methods. We then demonstrated that RiboLace is capable of providing positional data with nucleotide resolution of translational events when used for ribosome profiling, requiring ~40 times less material than current methods.
Ribo-seq protocols. We observed that >95% of ribosome-protected fragments were mapped on the coding region, with the characteristic trinucleotide periodicity suggestive of active ribosomes flowing along the transcripts and almost no signal on either the 5'- and 3'-UTRs of mRNAs.

We showed in two different case studies that RiboLace and Ribo-seq significantly differ in translation estimates for hundreds of genes. Functional annotation analysis suggests that Ribo-seq-enriched transcripts are more pertinent to the biological system under study with respect to Ribo-seq-enriched transcripts. In addition, RiboLace-specific transcripts are enriched for translation-related genes.

Overall, our data suggest that RiboLace is an effective approach for ribosome profiling experiments, in terms of required-sample input and accuracy in ribosome-protected fragments detection. RiboLace protocols can be further adjusted to (1) isolate ribosomes from other organisms than human and mouse or to (2) isolate ribosomes from specific eukaryotic cellular compartments such as the endoplasmic reticulum or organelles such as mitochondria.

In summary, RiboLace can be used to capture ribosomes in active translation with challenging or troublesome biological samples with low-input material for reliable ribosome profiling. Our method empowers scientists to efficiently and reproducibly determine the actual translational state of a biological system.

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SUPPLEMENTAL INFORMATION

Supplemental Information includes six figures and three tables and can be found with this article online at https://doi.org/10.1016/j.celrep.2018.09.084.
puromycin-associated nascent chain proteomics (PUNCH-P). Nat. Protoc. 9, 751–760.


# STAR METHODS

## KEY RESOURCES TABLE

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## Deposited Data

| Sequencing data (mouse brain) | This study | GEO: GSE102354 |
| Sequencing data (HEK293)      | This study | GEO: GSE112353 |

(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Gabriella Viero (gabriella.viero@cnr.it).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

**Tissues, cell lines and growth conditions**

Human MCF7 (ATCC catalog no. ATCC® HTB-22), HeLa cells and HEK293 cells and murine NSC34 (CEDARLANE catalog no. CLU140) cell lines were seeded on adherent plates and maintained at 37°C, 5% CO2 in DMEM supplemented with 10% Fetal Bovine Serum (FBS), 2 mM L-glutamine, 100 units/mL penicillin and 100 µg/mL of streptomycin. Cells were used at 80% of confluence. For starvation treatments cells were kept 0.5% FBS, 2 mM L-glutamine, 100 units/mL penicillin and 100 µg/mL of streptomycin for at least 12. For EGF treatment, after starvation EFG was added at 1 µg/mL for 4 h.
Wild-type mice mice were obtained from breeding stocks at the University of Edinburgh. All procedures were performed under licensed authority from the UK Home Office (PPL 60/4569).

**METHOD DETAILS**

**Chemical synthesis of the 3P molecule**

A solution of DCC in DMF was added dropwise to Biotin and NHS in DMF, to obtain a white precipitate (Biotin-NHS). Biotin-NHS in MeCN was added to a jellifame (2,2’-Ethylenedioxy)bis(ethylamine) - MeCN solution to yield a white hygroscopic solid (BJ1), then dissolved in dry pyridine and reacted with CDI to obtain a product called BJ1’. Puromycin was dissolved in pyridine, reacted with CDI and added to a BOC protected jellifame in DCM. The product (PJ) was partitioned between DCM and water, evaporated and triturated in ethyl acetate. Trifluoroacetic acid was added dropwise to a stirred suspension of PJ in DCM. The solution was cooled and stirred overnight, diluted with chloroform and the solvent evaporated to give a yellow oil product (PJ’). BJ1’ and PJ’ were dissolved in pyridine and stirred overnight under N2. The product (3P) was purified by column chromatography and preparative HPLC.

**Depletion assay of 3P with streptavidin coated beads**

A volume of 50 µL of 2 M NaCl, 1 mM 3P, 1 mM EDTA, 10 mM Tris-HCl, pH 7.5 in DEPC water, was added to different amount of magnetic beads and the suspension was incubated for 5 min at RT. After separation on a magnetic rack, the absorbance of the supernatant was measured at 272 nm.

**In vitro - cell free Transcription/Translation**

*In vitro* translation reactions of the full-length Luciferase (inserted in SP6 DNA plasmid, Promega) and full-length EGFP (inserted in the IP-PR2 plasmid) were obtained with the TnT Quick Coupled Transcription/Translation System (Promega) according to manufacturer’s instructions.

**Preparation of RiboLace beads**

For each sample, a volume of 20 µL of Dynabeads® MyOne Streptavidin C1 (Life Technology) or 30 µL of Streptavidin Mag Sepharose (10% slurry, GE Healthcare Life Sciences) were washed for 5 min with a 0.05 M NaCl, 0.1 M NaOH, in DEPC-treated water. Then, beads were washed with 500 µL of nuclease free water and with Binding Buffer (2 M NaCl, 10 mM Tris-HCl, pH 7.5 in DEPC water). For functionalization with 3P, 30 µL of a 1 mM solution of 3P in Binding Buffer was added to the beads followed by an incubation of 1 h, mixing at 1400 rpm at 20°C. Beads were end-capped by incubating for 10 min at 1400 rpm at RT in the presence of biotin-methoxypolyethylene glycol conjugate (mP, Creative Pegworks) at 0.5 mM. RiboLace beads were washed with 500 µL of nuclease free water, placed on a magnet for 2 min, and washed with 500 µL of nuclease free water. The efficiency of binding was calculated by measuring the ratio between the absorbance of the supernatant at 270 nm, and the absorbance at 270 nm of a 1 mM starting solution. Finally, beads were washed twice with 500 µL of W-buffer (10 mM NaCl, 10 mM MgCl2, 20 µg/mL cycloheximide, 10 mM HEPES, pH 7 in DEPC water) and used for active ribosome pull-down. As a negative control, beads were functionalized with a 0.5 mM solution of a biotin-methoxypolyethylene glycol conjugate (1000 Daltons, Creative Pegworks).

**RiboLace and IVTT system**

IVTT mix reaction (Promega) was used with the abovementioned plasmids for 40 min. The reaction was stopped adding cycloheximide (10 µg/mL) for 3 minutes before addition of W-buffer. The solution was divided into 3 vials, each containing 150 µL. The first was used for the extraction of total RNA, the second used with RiboLace, and the third for control mP-beads. For RiboLace, the reaction mix was added to the functionalized beads and incubated for 1 h in orbital rotation at 2 rpm at 4°C. The tube was then kept on ice on a magnetic stand for 5 min to pellet the beads–bounded-ribosomes. The supernatant was separated and beads washed two times with 500 µL of W-buffer. Beads were dissolved in 100 µL of 100 mM NaCl, 10 mM MgCl2, Trizma HCL, pH 7.5 in DEPC water. RNA was extracted from the beads–bounded-ribosomes with Trizol (Thermo Fischer Scientific) and solubilized in 30 µL of RNase-free water and DNase I treated. cDNA was obtained synthesis and RT-qPCR were run using KAPA SYBT FAST qPCR kit (KAPA Biosystem) according to the following protocol: 3 min - 95°C activation; 2 s - 95°C; 20 s - 57°C; 25 cycles; 65°C to 95°C melting ramp. Primers are listed in Table S1. Data were processed with Bio-Rad CFX-Manager 1.6 software. The fold change ratio was determined as the ratio between the ΔΔCt of treated and not treated sample. The delta cross-threshold (ΔCt) was determined respect to the control, and the relative ΔΔCt calculated respect to the control (mP-beads ΔCt).

For determining the total amount of newly synthesized proteins, we employed ε-labeled biotinylated lysine-tRNA complex using Transcend Non-Radioactive Translation Detection Systems according to the manufacturer’s instructions (Promega) and the effective translation verified by SDS–PAGE.

**Luciferase assay**

Real-time measurements of luciferase activity were recorded at 37°C with 5% CO2 using the Infinite 200 PRO reader (Tecan), according to manufacturer’s instructions. In brief, the IVTT reactions were plated in a 96-well plate and the luciferase translation efficiency was monitored by means of luminescence signal using the Bright-Glo Luciferase Assay System (Promega).
Purification of active polysomes and ribosomes with RiboLace

MCF7, HeLa, HEK293 or NSC-34 cells were seeded at 1.5 × 10⁶ cells/dish and kept in culture until reaching 80% of confluence. Cells were then treated with 10 μg/mL of cycloheximide (CHX) for 5 min at 37°C before lysis. Cell lysates were obtained according to (Viero et al., 2015). Tissues were dissected immediately following sacrifice and pulverized under liquid nitrogen using a pestle and a mortar and the lysates was obtained according to (Bernabo et al., 2017). The lysate was aliquoted and stored at −80°C for not more than a month, avoiding more than one freeze/thaw cycle.

RNA absorbance was measured (260 nm) by Nanodrop ND-1000 UV-VIS Spectrophotometer and the lysate diluted to 0.5 - 1.7 a.u. A260/μL with W-buffer (10 mM NaCl, 10 mM MgCl₂, 20 μg/mL cycloheximide, 10 mM HEPES, pH 7 in DEPC water). For polysome profiling with RiboLace, the solution obtained was added directly to the functionalized beads and the suspension incubated on a wheel for 1 hour in orbital rotation, 3 rpm (StarLab Rotator), at 4°C. The tube was then kept on ice on a magnetic stand for 5 min to pellet the beads–bounded-ribosomes and washed twice with W-buffer. RNA was extracted by acidic-phenol chloroform separation. The quality of the RNA samples was assessed using Agilent Bioanalyzer 2100 and Nanodrop ND-1000 Spectrophotometer (Thermo Scientific). Whole transcriptome library preparation was performed starting from 1 μg of total RNA with RIN ≥ 8. Following enrichment of poly-A containing mRNA molecules using poly-dT oligo-attached magnetic beads, all recovered RNA was processed using the Illumina TruSeq RNA Sample Preparation Kit sequenced. Sequencing was carried out on Illumina HiSeq 2000 using the protocol HCS 1.5.15.1 in single reads.

For ribosome profiling with RiboLace, a total volume of lysate corresponding to 0.3 A.U.260nm was treated with 1.5 U of Artseq nuclease (Epicenter) or RNaseI for 45 min at 20°C. The reaction was stopped with 10U SUPERase In RNase inhibitor, 10 min on ice. Then, the sample was incubated with RiboLace beads for 1 hour in orbital rotation, 3 rpm (StarLab Rotator) at 4°C. The tubes were then kept on ice on a magnetic stand for 5 min to pellet the beads–bounded-ribosomes. The supernatant was discarded (unbound fraction) and beads washed twice with W-buffer. Finally, beads were solubilized in 200 μL of W buffer containing 20 U SUPERase In. The RNA was extracted using acid-phenol:chloroform (Ambion) after incubation with 1% SDS, 0.1 mg of proteinase K (Euroclone) at 37°C for 75 min. Library preparation was adapted from a previous protocol (Ingolia et al., 2012). The ribosome profiling library PCR forward primer and indexed reverse primers are in Table S2. Libraries (175 nt) were PAGE purified from a 8% not-denaturing TBE polyacrylamide and characterized using the Agilent 2100 Bioanalyzer (High-Sensitivity DNA assay).

Ribo-seq data were obtained using the protocol in (Ingolia et al., 2012).

RNA extraction for RNA-Seq and POL-Seq

For RNA-Seq the RNA was extracted from MCF7 cell lysates, obtained as previously described, with acid phenol:chloroform (Ambion catalog no. AM9720) extraction. For total polysomal RNA extraction, polysomes were obtained according to (Viero et al., 2015) using sucrose gradient fractionation. Briefly, sucrose fractions were collected and treated with 200 μg/mL proteinase K (Life technologies), 1% SDS in DEPC water and RNase Inhibitor (0.4 a.u./μL) for 1.5 h at 37°C. After phenol:chloroform extraction and isopropanol precipitation, polysomal RNA was quantified by Nanodrop ND-1000 UV-VIS Spectrophotometer and Agilent 2100 Bioanalyzer with RNA 6000 pico kit (Agilent).

Whole transcriptome library preparation was performed starting from 1 μg of total RNA with RIN ≥ 8. Following enrichment of poly-A containing mRNA molecules using poly-dT oligo-attached magnetic beads, all recovered RNA was processed using the Illumina TruSeq RNA Sample Preparation Kit (Illumina #FC-122-1001 #FC-122-1002) and the protocol v2 Rev. C. Completed libraries were enriched of poly-A containing mRNA molecules using poly-dT oligo-attached magnetic beads, all recovered RNA was processed using the Agilent 2100 Bioanalyzer and Illumina HiSeq 2000 using the protocol HCS 1.5.15.1 in single reads. Experiments were performed in biological duplicate.

Immunoblotting

Cell lysates were prepared in hypotonic lysis buffer (10 mM NaCl, 10 mM MgCl₂, 10 mM Tris–HCl, pH 7.5, 1% Triton X-100, 1% sodium deoxycholate, 5 U/mL DNaseI, 200 U/mL RNase inhibitor, 1 mM dithiothreitol and 10 μg/mL cycloheximide) or RIPA buffer. Proteins were separated in SDS–polyacrylamide gel electrophoresis and transferred onto PVDF membranes. The membranes were blocked in 5% BSA (Sigma) in TBS-Tween (0.1% Tween) for 1 hour, incubated in primary antibody o.n. The primary antibodies used are listed in the STAR Methods section. After incubation with secondary antibodies conjugated to horseradish peroxidase and extensive washing, the blots were developed using ECL Plus (GE Healthcare,) or SuperSignal West Femto Maximum Sensitivity Substrate (Thermo Scientific). Signals were acquired with ChemDoc-It (Bio-Rad) and analyzed with ImageJ software (v 1.45 s). All experiments were run in triplicate.

qPCR

RNA expression was analyzed by TaqMan assay, the gene name, aliases, chromosome location and TaqMan ID are reported in Table S3. All TaqMan probes were purchased from Life Technologies. Relative quantification of target genes was determined calculating the delta cross-threshold (ΔCT) respect to the 18S housekeeping gene and the relative ΔΔCT calculated respect to the total RNA sample, according to the Pfaffii method. RT was performed using random hexamers, single strand reverse transcriptase (RevertAid RT Reverse Transcription Kit, Life Technologies #K1622).
Protein sample preparation for MS
MCF7 cells were grown in 100 mm Petri dishes in complete medium (10% FBS) or under starvation (1% FBK) for 12 h. Two dishes were lysed with RIPA buffer (50mM Tris pH 7.4, 150mM NaCl, 0.25% Igepal, 1mM EDTA and 0.5% Sodium Deoxycholate) in the presence of 0.5% Phosphatase Inhibitor Cocktails, 5 μg/mL Peptastin A and 0.25% Protease Inhibitor Cocktail. After protein extraction according to the methanol-chloroform procedure, proteins were solubilized in 6 M urea/2 M thiourea, 10 mM HEPES pH 8.0 and incubated at RT for 30min in the presence of 5mM DTT and for additional 20min with 5.5mM iodoacetamide. Peptides were obtained by digestion with 1μg of LysC solution/50μg of protein for 3 hours at RT and then with 1μg of Trypsin/50μg protein overnight. Peptides were purified on C18 Pipette Tips (Pierce, Thermo Scientific). Mass spectra were acquired in the Orbitrap Fusion Tribrid analyzer at Thermo Fisher Scientific, Switzerland. Experiments were performed in triplicates.

QUANTIFICATION AND STATISTICAL ANALYSIS

Data analysis of RNA-seq and POL-seq experiments
Fastq files were checked for quality control with FastQC. Reads generated from each sample were aligned to the human genome with Tophat (version 2.0.14), using the Gencode 22 transcript annotation as transcriptome guide. All programs were used with default settings. Mapped reads were assembled into transcripts guided by reference annotation (Gencode 22) with Cufflinks (version 2.2.1). Expression levels were quantified by Cufflinks with normalized FPKM (fragments per kilobase of exon per million mapped fragments). Differential expression analysis was performed with Cuffdiff.

Data analysis of MS
The raw MS data were analyzed with MaxQuant software version 1.6.1.0 using default settings and Label Free Quantification (LFQ).

Data analysis of ribosome profiling experiments
For all ribosome profiling analyses we used riboWaltz (Lauria et al., 2018). Briefly, reads were processed by removing 5’ adapters, discarding reads shorter than 20 nucleotides and trimming the first nucleotide of the remaining ones (using Trimomatic v0.36). For the mouse brain and the HEK293 dataset, respectively, reads mapping on the collection of M. musculus and H. sapiens rRNAs (from the SILVA rRNA database, release 119) and tRNAs (from the Genomic tRNA database: gtrnadb.ucsc.edu/) were removed. Remaining reads were mapped on the mouse transcriptome (using the Gencode M6 transcript annotations) or human transcriptome (Gencode 25): antisense reads were removed and reads entirely mapping to the same nucleotides were considered identical and collapsed. All the alignments were performed with Bowtie2 (v2.2.6) employing default settings. Normalization among replicates was performed with the trimmed mean of M-values normalization method (TMM) implemented in edgeR. Protein coding transcripts used for further analyses were selected using a threshold on their signal (FPKM and CPM values > 80th percentile). Differential analyses between RiboLace and Ribo-Seq were performed with the edgeR Bioconductor package. Significantly enriched transcripts were selected with the following thresholds: absolute log2 fold enrichment > 1, P value < 0.05, normalized number of ribosome-protected fragment > 1 per million. Functional annotation enrichment analyses were performed with Enrichr (http://amp.pharm.mssm.edu/Enrichr/).

DATA AND SOFTWARE AVAILABILITY
Raw and analyzed sequencing data have been deposited under GEO: GSE102354 (mouse brain), GSE112353 (HEK293), GSE112295 (MCF7). The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository with the dataset identifier PXD009417. Classic ribosome profiling (Ribo-Seq) data from mouse brain were retrieved from GEO: GSE102318.