Translation of CircRNAs

Highlights

- Specific circRNAs are associated with translating ribosomes
- Ribosome footprinting reads match termination codon signature for circMbl
- circMbl3-derived protein is detected by mass spectrometry
- circRNAs are translated in vitro and in vivo in a cap-independent manner

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In Brief

Pamudurti et al. show that a subset of circRNAs is translated. These circRNAs generally share the start codon with the hosting RNA, encode proteins with specific protein domains, and are translated in a cap-independent manner.
Translation of CircRNAs


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SUMMARY

Circular RNAs (circRNAs) are abundant and evolutionarily conserved RNAs of largely unknown function. Here, we show that a subset of circRNAs is translated in vivo. By performing ribosome footprinting from fly heads, we demonstrate that a group of circRNAs is associated with translating ribosomes. Many of these ribo-circRNAs use the start codon of the hosting mRNA, are bound by membrane-associated ribosomes, and have evolutionarily conserved termination codons. In addition, we found that a circRNA generated from the muscleblind locus encodes a protein, which we detected in fly head extracts by mass spectrometry. Next, by performing in vivo and in vitro translation assays, we show that UTRs of ribo-circRNAs (cUTRs) allow cap-independent translation. Moreover, we found that starvation and FOXO likely regulate the translation of a circMbl isoform. Altogether, our study provides strong evidence for translation of circRNAs, revealing the existence of an unexplored layer of gene activity.

INTRODUCTION

Circular RNAs (circRNAs) in animals are a large class of particularly stable RNAs produced by circularization of specific exons (Jeck and Sharpless, 2014; Wang et al., 2014). In vivo, circRNAs are generated by the spliceosome via backsplicing: the 3’ end of an exon is covalently linked to the 5’ end of an upstream exon (Ashwal-Fluss et al., 2014; Starke et al., 2015). Most circRNAs originate from protein-coding genes and contain complete exons. Particularly in the fly brain (Westholm et al., 2014) and mammalian neuronal and muscle tissues (Rybak-Wolf et al., 2015), circRNAs are highly abundant and evolutionarily conserved.

The function of only a few circRNAs has been elucidated. At the molecular level, CDR1as acts as a microRNA (miRNA) sponge (Hansen et al., 2013; Memczak et al., 2013). Other circRNAs can regulate the function of RNA-binding proteins (Ashwal-Fluss et al., 2014) or the transcription rate of its host gene (Li et al., 2015). However, the functions of thousands of described circRNAs (Glazar et al., 2014) remain unknown.

Although it has been shown that in principle circRNAs can be translated in vitro and in vivo (Abe et al., 2015; Chen and Samow, 1995; Li and Lytton, 1999; Wang and Wang, 2015), the possibility that circRNAs are endogenously translated has so far been only indirectly tested (Guo et al., 2014; Jeck and Sharpless, 2014). In eukaryotes, the canonical translation process begins with the binding of the open pre-initiation complex, which contains the small ribosomal subunit. In addition ribosomes can be recruited to an internal start codon by a mechanism that depends on an internal ribosome entry site (IRES). Translation of several viral proteins is dependent on IRES, and cellular IRESs have been identified in many organisms, including humans, although their mechanism of action remains controversial (Jackson, 2013; Weingarten-Gabbay et al., 2016).

RESULTS

A Subset of Drosophila CircRNAs Is Associated with Translating Ribosomes

To determine the specific coding potential of Drosophila circRNAs, we identified those with open reading frames (ORFs) across the backspliced junction. We found that most circRNAs have coding potential (see list in Table S1). However, we obtained similar results with a set of control exons (data not shown).

After annotating the Drosophila circRNA ORFs (cORFs), we searched for evidence of their translation. We utilized previously published ribosome footprinting (RFP) datasets (Aspden et al., 2014; Dunn et al., 2013; Kronja et al., 2014; Mettinen and...
Minigenes of Ribo-CircRNAs Produce Proteins

We then generated intron-exon-intron minigenes for expression of specific circRNAs in *Drosophila* S2 cells. We focused on the ribo-circRNAs circMbl, circCdi, and circPde8 and used as negative controls circRNAs for which we did not observe RFP reads in this cell type: circHaspin and circCamK1. We engineered the minigenes to express a V5-tagged protein in case of a circularization event (Figure 2A). Cells transfected with circMblV5, circCdiV5, or circPde8V5 minigenes produced V5-tagged proteins, whereas those transfected with circHaspinV5 or circCamK1V5 minigenes failed to express a V5-tagged protein (Figure 2B). The two V5-tagged protein products in the cells transfected with the circPde8V5 minigene are due to the translation of two proteins generated from the circRNA with or without an internal intron. We detected a non-specific band above the circCdi protein (Figure 2B).

Moreover, the MBL-positive and V5-positive western signals originate from the same protein (Figure S1B). We detected stronger anti-MBL immunoreactivity in cells that expressed the circMbl minigene. Therefore, we visualized the RNA by northern blot (Figure S1C). Indeed, we found that circMbl and circMblV5 minigenes generate linear concatemers that can also potentially produce the observed peptides (Figures 2C, S1D, and S1E). The circMbl minigene produces more protein than the circMblV5 minigene (see “Input” in Figure S1B) despite much lower levels of concatemers, suggesting that the circRNA is the main source of the detected protein. The circPde8V5 minigene construct also produces substantial amounts of linear concatemers (Figures S1F and S1G). Interestingly, other minigenes (i.e., circCamK1V5 and circCdiV5; Figures S1G–S1I) produce lower levels or no concatemers.

To determine whether the produced proteins are generated from the circRNA, we determined whether minigene-derived circRNA molecules are associated with translating ribosomes. We co-transfected *Drosophila* S2 cells with the circMblV5, circCdiV5, and circCamK1V5 minigenes, a plasmid driving expression of GFP, and a plasmid driving expression of a circRNA in which a split Cherry protein is under the control of the circMbl cUTR (UTR of circMbl: all the sequences in circMbl except the cUTR). We co-transfected *Drosophila* S2 cells with the circMblV5, circCdiV5, or circPde8V5 minigenes without the V5-tag sequence, despite similar amounts of circMbl RNA (data not shown). This result suggests that several RNA isoforms could originate from the circMblV5 minigene. Therefore, we visualized the RNA by northern blot (Figure S1C). Indeed, we found that circMbl and circMblV5 minigenes generate linear concatemers that can also potentially produce the observed peptides (Figures 2C, S1D, and S1E). The circMbl minigene produces more protein than the circMblV5 minigene (see “Input” in Figure S1B) despite much lower levels of concatemers, suggesting that the circRNA is the main source of the detected protein. The circPde8V5 minigene construct also produces substantial amounts of linear concatemers (Figures S1F and S1G). Interestingly, other minigenes (i.e., circCamK1V5 and circCdiV5; Figures S1G–S1I) produce lower levels or no concatemers.

Figure 1. A Subset of CircRNAs Is Associated with Translating Ribosomes

(A) Approach used to detect circRNA reads in RFP data. (B) Datasets utilized and ribo-circRNAs detected. (C) circRNA expression was plotted against their ranking in these datasets. Red dots indicate ribo-circRNAs. (D and E) Integrated genome browser (IGV) snapshots of small and large RFP datasets for tai (D) and ct (E) show that more RFP reads are in the circularized exons, mainly in the lighter polysomes (red).

See Table S1.

Björklund, 2015) and searched for reads across the circRNA-specific junctions. Using a database of backsplice and canonical junctions (Figure 1A), we identified 37 circRNAs (referred hereafter as ribo-circRNAs) with at least one specific RFP read (Figure 1B; Table S2). The association of these ribo-circRNAs with the ribosomes seems to be more than a chance event because: (1) ribo-circRNAs contain circRNAs expressed at various levels (Figure 1C); and (2) despite the low RFP counts, some ribo-circRNAs are detected on the same scale as their hosting mRNA (Figure S1A; Table S2), suggesting comparable translational rates. Moreover, for a few circRNAs, we found more RFP reads covering the exons containing the circRNAs than other exons (Figures 1D and 1E). In this experiment, the polysomes were separated according to size before the footprinting assay. Indeed, the enrichment of RFP reads in specific circRNA-containing exons was prominent only in RNAs associated with light polysomal fractions. This was highly significant for circTai (7.2-fold enrichment, p < 1 × 10⁻⁵⁴; Figure 1D) and circCt (2.6-fold enrichment, p < 1 × 10⁻³¹; Figure 1E).

Minigenes of Ribo-CircRNAs Produce Proteins

We then generated intron-exon-intron minigenes for expression of specific circRNAs in *Drosophila* S2 cells. We focused on the ribo-circRNAs circMbl, circCdi, and circPde8 and used as negative controls circRNAs for which we did not observe RFP reads in this cell type: circHaspin and circCamK1. We engineered the minigenes to express a V5-tagged protein in case of a circularization event (Figure 2A). Cells transfected with circMblV5, circCdiV5, or circPde8V5 minigenes produced V5-tagged proteins, whereas those transfected with circHaspinV5 or circCamK1V5 minigenes failed to express a V5-tagged protein (Figure 2B). The two V5-tagged protein products in the cells transfected with the circPde8V5 minigene are due to the translation of two proteins generated from the circRNA with or without an internal intron. We detected a non-specific band above the circCdi protein (Figure 2B).
protein construct contains the CamKI flanking intronic sequences that drive very efficient RNA circularization, with no detectable concatemers (Figures 2D and S1G–S1I). Cells transfected with the split Cherry constructs display strong expression of CHERRY protein (Figure 2D). We then performed sucrose density gradient centrifugation. Both GFP and an endogenous mRNA (rp49) were strongly associated with monosomes and polysomes (Figures S2A and S2B). Most circCamKI RNAs co-migrate with the lighter fractions, whereas most of the circMbl, circCdi, and circCherry RNA co-migrates with the monosome and polysomal fractions (Figure S2A and S2B). We then treated the RNA from the different polysome gradient fractions with RNaseR. This assay confirmed that the RNAs derived from the circMblV5, circCherry minigenes, and to a lesser extent, the circCdiV5 minigene are indeed circular (Figures S2C–S2E).

Importantly, the expression of the V5-tagged protein from the circMblV5 minigene depends on the start codon in the circMbl exon (Figure 2E). In addition, mutation of the splice site resulted in the disappearance of anti-V5 immunoreactivity and in the generation of a slightly longer protein because of a stop codon in the downstream intron (Figure 2E). Indeed, mutation of the 5’ splice site (5’ss) leads to the use of a cryptic 5’ss splice site and the generation of a new type of circRNA molecule (circMblV5’ss; Figures S1D and S1E), which is likely the template for the longer protein product observed in Figure 2E. From these experiments we conclude that a substantial fraction of the V5-tagged proteins detected in Figure 2B originates from circRNA molecules. We also concluded that the intron-exon-intron minigenes should be used with caution.

To extend these findings to an in vivo system, we generated transgenic flies expressing a circMbl minigene. To minimize formation of concatemers, we did not tag the circMbl-encoded protein. We expressed this circMbl minigene with the actin-gal4 driver and analyzed muscleblind RNA and protein isoforms from fly heads. Indeed, expression of the minigene resulted in a 4-fold increase in the levels of circMbl (Figures 2F and 2G). In these flies, we did not observe linear concatemers, although we observed small amounts of a larger RNaseR-resistant RNA, likely a larger circMbl molecule (see asterisk in Figure 2F). Importantly, in these flies we consistently detected MBL-immunoreactive bands of
Figure 3. Genome-wide Analysis of Translation in Fly Heads by RFP
(A) Meta-analysis of the RFP reads.
(B) “Pile ups” of the RFP reads at two different resolutions in the Arr2 locus.
(C) Meta-analysis of the RFP reads in the proximity of the start and stop codons of all annotated genes. The total frequency of RFP read 5’ end positions plotted against the relative distance (in nucleotides) to the start/stop codon (left). The same information, but subdivided into reads of different length, is shown, with frequency represented by color (middle). Relative enrichment of RFP reads around real start/stop codons over the background frequency for reads of such length in the vicinity of start/stop codons, expressed as log ratio of these frequencies (right). Positive scores (red) indicate consistency with the signature of start/stop codons, and negative scores (blue) consistency with the uniform background distribution.
(D) Main over-represented GO terms for the mRNAs significantly higher in the normal detergent library. The detergent libraries were sequenced 3.4 times deeper.
approximately 10 kDa (Figure 2H), strongly suggesting that circMbl can also produce a protein in adult flies.

**RFP from Fly Heads**

To determine whether circRNAs are translated in a more relevant tissue, we set up the RFP methodology in fly heads. For doing so, we used two different extraction buffers, which contain none or standard detergent concentrations. As expected, we obtained strong coverage of the known ORFs and their 5’ UTRs (Figure 3A).

RFP fragment sequencing experiments preferentially fall into particular positions with respect to the ORF. As expected, we observed phasing of the reads (Figure 3B), which indicates that particular positions with respect to the ORF. Despite these limitations, we observed a high stop codon score for the most abundant ribo-circRNA, circMbl (score = 37; p < 0.007; Figure 3G, bottom panels). Thus, this result supports translation of circMbl in fly heads. Interestingly, the high score of this stop codon disappears when all the RFP reads (not only the ones containing the backspliced reads) are considered (Figure 3G, top panels). Therefore, this stop codon is used only in the context of the circRNA. In summary, the RFP pattern around the stop codon of circMbl strongly supports endogenous translation of this circRNA.

It could be that the RFP backspliced reads arise from a small, undetected fraction of linear, trans-spliced products rather than circRNA molecules. To test this possibility, we utilized a fly strain carrying a transgene for expressing EGFP-tagged ribosomes (Huang et al., 2013). We immunopurified the tagged ribosomes and analyzed the RNA bound to them. We measured the relative immunoprecipitation (IP) enrichment by comparing the enrichment (immunoprecipitate/input ratio) between the assayed target and an abundant pre-mRNA (pre-mbl mRNA), which is not expected to be associated with ribosomes. Indeed, we found a strong enrichment for four ribo-circRNAs assayed, demonstrating their association with ribosomes (Figure 4A). To distinguish between associations of ribosomes with linear and circRNA molecules, we treated ribosome-bound RNAs with RNaseR. For doing so, we set up an approach to determine RNaseR sensitivity from low RNA amounts (see STAR Methods). Indeed, we found that the four assayed ribo-circRNAs are strongly resistant to RNaseR, demonstrating that they are bound to the ribosome as circRNA molecules (Figure 4B). Importantly, the RNaseR treatment resulted in complete degradation of a linear RNA in the same sample (rp49 mRNA; Figure 4C). In addition, we found that the ribosome-bound circSh fraction is strongly resistant to RNaseR treatment (Figure 4C, red bars), even when this circRNA displays sensitivity to RNaseR.
Figure 4. A Subset of CircRNAs Is Associated with Translating Ribosomes

(A) Results of the translating ribosome affinity purification (TRAP) assay. Ribo-GFP strains: Actin-Gal4; UAS-EGFPL10a. Control strain: UAS-EGFPL10a. The number indicates the difference in the normalized ratio between the Ribo-GFP and control strains. n = 4.

(B) RNaseR validation of the ribosome-bound circRNAs. RNA from the input and IP TRAP samples was subjected to RNaseR treatment. Mouse RNA was added and quantified as spike-in for the normalization. The RNaseR resistance was assessed and normalized to the one observed in the input sample. n = 3.

(C) RNaseR validation of the ribosome-bound RNAs. Normalization was performed using the internal spike-in. n = 3.

(D) RNA expression in the mbl locus. Red track represents the expression of the different exons in fly heads. Hemi-circles indicate the most abundant circRNAs. Top inset: comparison of the expression levels in the mock and RNase-treated samples corroborates the high expression of circMbl3 and circMbl4. Right boxes indicate the size of the proteins produced by the annotated mbl mRNAs or circRNAs.

(F) circMbl3 Northern Blot

(legend continued on next page)
A CircRNA Generated from the mbl Locus Produces a Detectable Protein

The mbl locus produces several circRNAs, some of which are highly abundant (see Figure 4D; Ashwal-Fluss et al., 2014; Westholm et al., 2014). Most of those circRNAs include the second exon, which contains the start codon for the putative circRNA-encoded ORF and would also include the same 5’ exon, which contains the start codon for the putative circRNA-encoded ORF and would also include the same 5’ exon, which contains the start codon for the putative circRNA-encoded ORF and would also include the same 5’ UTTr regulated sequences. To determine the presence of any of these proteins in fly head extracts, we performed targeted mass spectrometry from a fly head MBL immunoprecipitate. We focused on the C terminus of the putative proteins, which is encoded across the circRNA junction. We utilized synthetic peptides to determine their characteristic spectra for which we then searched in the fly head immunoprecipitate. Unfortunately, the ionization capacity of the peptide designed to identify the circMbl1-encoded peptide was extremely low, precluding the efficient identification of our best candidate. However, we found a consistent and very high confidence hit for a peptide that can only be produced by circMbl3 (Figures 4D, 4E, and S4F). This circRNA encodes a 37.04 kDa protein and might either include or skip an alternative internal exon (exon 5 of mbl). To verify the existence of this circRNA in fly heads, we performed a Northern blot analysis using a specific circMbl3 probe. We detected two RNaseR-resistant bands of the predicted size due to alternative splicing of the fifth mbl exon in the assayed fly strains (yw and CantonS [CS]; Figure 4E). Importantly, circMbl3 was identified as resistant to RNaseR treatment in our published dataset (Ashwal-Fluss et al., 2014). Because the identified peptide could only have originated from the circMbl3 molecule and not from any of the other known MBL isoforms (Figure 4E), these data constitute very strong evidence for the translation of this circRNA.

Analysis of CircRNA Features

We utilized 151 ribo-circRNAs (Table S2) for further analysis. We compared these ribo-circRNAs with the set of control exons utilized for cORF identification, as well as the ORF contained within circRNAs for which we did not find evidence of translation (untranslated cORFs, or utcORFs). Interestingly, we found that the ribo-circRNAs have a strong bias toward 5’ UTRs (p < 0.0055; Figure 5A). In 40% of these circRNAs, translation is predicted to share the start codon with the host gene (Figure 5B). We observed a similar bias for both parameters for the utcORFs, suggesting the presence of false negatives (Figures 5A and 5B). Interestingly, ribo-circRNAs are significantly longer than the control groups (Figure 5A), suggesting that they can carry out functions. We compared the protein domains present in the predicted cORFs with those present in random exons from the same hosting genes. Interestingly, we found a subset of domains, which were significantly enriched in the cORF group (Figure SS5E). Half of those domains (three of six) are among those encoded by exons contained in ribo-circRNAs. We did not observe RFP reads for exons containing any of the domains enriched in our random dataset, strongly suggesting that ribo-circRNAs contain specific types of domains.

Ribo-CircRNAs in Mouse and Rat

To extend our observation to a mammalian system, we sought relevant RFP datasets (Ribo-seq) from mammalian cells and treatment in total RNA and in the input fraction (Figures 4C and S3E). This suggests the existence of two populations of transcripts carrying the backsplice junction (one linear and one circular). The circular transcript is bound to the translating ribosomes. If the translation of a subset of circRNAs is functionally important, it is reasonable to expect that their translation potential should be evolutionary conserved. Indeed, we observed strong conservation of the coding potential and levels of circMbl, the most abundant Drosophila circRNA. We based our conclusion on the results of multiple species alignment, as well as on the validation and quantification of backsplice junctions by Sanger sequencing for five different Drosophila species (Figures 5E, 5F, and S5B).

To investigate whether cORFs are under negative selection, we built multiple species alignments of each circRNA candidate from up to 24 insect species (see STAR Methods; unpublished data). Figure 5E shows excerpts of such an alignment for circMbl (see Table S3 for the scores of all cORFs). Indeed, we observed that the cORFs show higher conservation of the stop codon compared with randomly selected stop codons in the same 5’ UTR (Figure 5G). These results are encouraging but should be taken with caution because of two limitations of the conservation assessments. First, cORF stop codons tend to fall close to the head-to-tail junction of the circRNA, and hence near the splice sites (the distribution of distances from the splice site is indeed to a good approximation geometric; see Figure SS5C). However, splice sites are under additional evolutionary constraints because of the requirement for proper splice-site recognition. Second, the cORF stop codon could also be the stop of an upstream ORF (uORF), potentially starting in a preceding 5’ UTR exon. When accounting for these factors, we observe higher conservation of stop codons that are linked to a potential uORF in both circRNAs and controls (Figure 5G; Figure SS5D). Thus, despite the higher degree of evolutionary conservation in comparison with the hosting 5’ UTR sequences, it is impossible to determine whether the observed conservation is due to the cORF, the uORF, or both.

As stated earlier, a large fraction of the ribo-circRNAs are in frame and share the start codon with the ORF of the hosting gene. Interestingly, we observed that many (31 out of 132) proteins encoded in ribo-circRNAs contain at least one identifierizable protein domain (see Table S4), suggesting that they can carry out functions. We compared the protein domains present in the predicted cORFs with those present in random exons from the same hosting genes. Interestingly, we found a subset of domains, which were significantly enriched in the cORF group (Figure SS5E). Half of those domains (three of six) are among those encoded by exons contained in ribo-circRNAs. We did not observe RFP reads for exons containing any of the domains enriched in our random dataset, strongly suggesting that ribo-circRNAs contain specific types of domains.
Figure 5. Features of Fly Ribo-CircRNAs

(A) Ribo-circRNAs are strongly enriched for 5′ UTR and CDS overlap (p < 0.0055, Fisher’s exact test).

(B) Percentage of control, utc-circRNAs, and ribo-circRNAs ORFs, which share their putative start codon with their hosting linear mRNA. The differences between ribo-circRNAs and control or utc-circRNAs are statistically significant (p < 2 × 10^{-10} and p < 0.05, respectively, Fisher’s exact test).

(C) Percentage of control, cORFs, and ribo-circRNAs that contain cUTR. The differences between ribo-circRNAs and control or utc-circRNAs are significant (p < 0.002 and p < 0.05, respectively, Fisher’s exact test).

(D) Average length of cUTR for control groups as well as for ribo-circRNAs (p < 0.019, double-sided Mann-Whitney U test).

(E) Excerpt of a multiple species alignment of circMbl. First codon position is in uppercase, and other codon positions are in lowercase. STP, stop codon. Arrows indicate the beginning (green) and end (red) of the exon.

(F) Ratio between the linear and circular forms of the second exon of mbl in different Drosophila species. n = 2.
tissues (in which also most circRNAs have cORFs; Figure S6A). We utilized a Ribo-seq dataset from rat brain and liver, and one from mouse C2C12 cells (de Klerk et al., 2015; Ori et al., 2015). We found unequivocal RFP reads for 34 and 158 circRNAs in the rat and mouse samples, respectively (Tables S5 and S6). As in Drosophila, these ribo-circRNAs are not necessarily among the most abundant circRNAs in the assayed tissue (Figures S6B and S6C). However, the numbers of backsplice junction mapping RFP reads were too low to assess phasing or make predictions on the rate of translation. This finding agrees with the number of spliced reads spanning normal mRNA junction, suggesting that levels of circRNA translation may be comparable with those of the host mRNA (Figures S6D and S6E). As observed in Drosophila, we found that the ribo-circRNAs are strongly enriched for 5' UTR overlap and tend to share the start codon with linear RNAs (Figures S6F–S6H).

Ribo-CircRNAs Contain Sequences that Can Promote Cap-Independent Translation

Because circRNAs do not contain a 5' cap, their translation should be dependent on the presence of a translation regulatory sequence. Cap-dependent translation can be efficiently inhibited by overexpression of the 4E-BP protein (Marr et al., 2007; Olson et al., 2013). We examined whether overexpression of Drosophila 4E-BP inhibits translation of the V5-tagged circRNA minigenes. Although the levels of GFP expressed from a linear mRNA molecule were severely diminished by co-expression of 4E-BP, expression of the circMbl or circCdi V5-tagged proteins was not affected (Figure 6A). Because the main protein product of the Pde8 minigene co-migrates with the 4E-BP protein (which is also V5 tagged), it was not possible to distinguish between these proteins. However, we found that the smaller (14.5 kDa) isoform was insensitive to 4E-BP overexpression, suggesting that sequences in the cUTR of this circRNA can drive cap-independent translation (Figure S7A). We observed a similar phenomenon with our circCherry artificial minigene (Figures 2F and 6B).

To validate these results, we cloned the circMbl, circCdi, circPde8, and circTai CUTFs in the straight or inverted orientation into a bicistronic reporter (Figure 6C, top; Olson et al., 2013). Remarkably, we found that the circMbl, circCdi, circTai, and circPde8 CUTFs can drive higher expression of the luciferase ORF from this bicistronic construct (Figures 6C and S7B). Interestingly, we noticed that the putative regulatory elements seem to act at least partially by increasing RNA stability (Figure S7C). We then measured firefly and Renilla luciferase levels in the presence of 4E-BP. As previously reported, overexpression of 4E-BP resulted in a decrease in the firefly/Renilla ratio of the Hepatitis C virus (HCV) reporter by approximately 40% (Figure 6D; Olson et al., 2013). We observed a more than 4-fold increase in the firefly luciferase/Renilla luciferase ratio for all the cUTR reporters after co-transfection with a 4E-BP expression plasmid (Figure 6D). The constructs containing the inverted cUTR sequences displayed much lower resistance to 4E-BP overexpression with the exception of the reverse Pde8 sequence, which displays luciferase levels barely above background (Figures 6C, 6D, and S7B).

CircMbl Can Be Translated In Vitro in a Cap-Independent Way

To study the mbl 5' UTR in a more controlled system, we performed an in vitro translation assay. We generated RNA reporters that contain a Renilla luciferase ORF downstream of the mbl 5' UTR and used them in a Drosophila cell-free translation system (Castagnetti et al., 2000; Chekulaeva et al., 2006). We first compared translation of two reporters: one bearing m'Gppp-cap (cap-circMbl-RLuc) and another containing an Appp-cap analog (circMbl-RLuc). Substitution of m'Gppp-cap with Appp-cap analog reduced mbl reporter translation 8-fold (Figure 6E). Similarly, bicistronic reporter with mbl 5' UTR was translated 10-fold less efficiently than m'Gppp-capped monocistronic mbl reporter (cap-Fluc-circMbl-RLuc; Figure 6E). These results show that in this system, the linear version of the mbl mRNA is translated mainly via cap-dependent initiation. However, because circRNAs represent a unique case of RNAs in which there is no cap, we reasoned that even 10% activity might still be relevant. Therefore, we generated a circRNA reporter that contains the circMbl cUTR (circMblcUTR; Figures S6D and S6E). Additionally, we generated circRNA reporters with cUTR in a reverse orientation (RevUTR) and a reporter from which most of the cUTR was deleted (ΔcUTR). In the context of the circRNA, circMbl cUTR was able to drive Renilla translation two times less efficiently than CrPV IRES (Figure 6F). Deletion of most of the cUTR abolished this activity, and the cUTR activity was reduced 2-fold when this element was inserted in a reverse orientation. Addition of a cap analog in trans was used as a further control to demonstrate that the CrPV IRES and circMbl reporters do not carry m'Gppp-cap and, therefore, cannot be translated in a cap-dependent manner (Figure 6F). In summary, our experiments demonstrate that the circMbl cUTR sequences can drive translation from a circRNA in a cap-independent manner, although this mechanism is not prevailing for a linear mbl reporter, at least in vitro. Interestingly, the inverted circMbl sequence can also promote circRNA translation, perhaps suggesting that the functional element is of a structural nature.

A Putative CircMbl1-Encoded Protein Is Enriched in Synaptosomes and Modulated by Starvation and FOXO

Using the available antibody against MBL, we could not always detect an endogenous band for putative circMbl-encoded proteins in fly head extracts. Therefore, we determined whether we could detect bands of the expected size in synaptosomal preparations from fly heads (Depner et al., 2014). Indeed, following subcellular fractionation, we could observe a band of
the expected size (slightly lighter than 10 kDa for circMbl1) from the fractions P2 and LP1, which are strongly enriched for synaptosomes. We also observed a band of the expected size (37 kDa) for the protein product produced from circMbl3 in the cytoplasmic fractions. However, we cannot rule out that this band is not the endogenous MBL isoform of 32 kDa. In addition, circMbl1 and circMbl3 RNA were also present in the same synaptosomal preparation (Figure 7B).

To extend this observation to a more relevant system, we used Drosophila S2 cells line carrying a plasmid expressing 4E-BP. Proteins expressed by the circMbl and circCdi minigenes are resistant to the activity of 4E-BP. (B) Quantification of the fluorescent signal from cells transfected with the Cherry minigene and GFP in the presence or absence of 4E-BP, detected by high-throughput microscopy. n = 3. In each experiment more than 3,000 cells were measured. (C) Relative firefly luciferase levels of the indicated bicistronic reporters. To normalize between experiments, we divided the luciferase reads to the ones obtained with the HCV IRES construct. n = 6. Error bars represent SEM.

Figure 6. Characterization of CircRNAs Putative IRESs

(A) Proteins expressed by the circMbl and circCdi minigenes are resistant to the activity of 4E-BP.

(B) Relative firefly luciferase levels of the indicated bicistronic reporters. To normalize between experiments, we divided the luciferase reads to the ones obtained with the HCV IRES construct. n = 6. Error bars represent SEM.

(D) Drosophila S2 cells were transfected with a bicistronic IRES reporter and with or without a plasmid expressing 4E-BP. The results show the ratio between the firefly/Renilla coefficients in presence or absence of 4E-BP. Error bars indicate SEM. n = 6.

(E) Expression of the G-capped RLuc reporter bearing mbl 5’ UTR (m7Gppp-circMbl-RLuc) was compared with the analogous A-capped reporter (Appp-circMbl-RLuc) and bicistronic reporter (m7Gppp-FLuc-circMbl-RLuc) in a cell-free Drosophila translation system. RLuc activity was normalized to FLuc activity; expression of m7Gppp-circMbl-RLuc was taken for 100%, and the rest was expressed accordingly.

(F) Linear reporter containing the A-capped CrPV IRES (CrPV-RL) was used as a positive control, and circular reporters with mbl 5’ UTR in a reverse orientation (circMblRevcUTR-RLuc) or deleted mbl 5’ UTR (circMblcUTR-RLuc) were used as negative controls in a cell-free Drosophila translation system. Each reaction was assembled in the presence or absence of m7GpppG.

See Figure S7.

mRNA or pre-mRNA, suggesting that it is due to stimulation of their specific protein synthesis (Figure S7F). Those isoforms are not likely originated from circRNAs, because the levels of circMbl isoforms are very low in Drosophila S2 cells.

To extend this observation to a more relevant system, we determined the effect of starvation, which activates cap-independent translation (Bar-Peled and Sabatini, 2014), on the levels of the ~10 kDa MBL-immunoreactive band observed in Figure 7A. As a positive control, we utilized a fly extract of the fly strain overexpressing circMbl (see earlier). Overexpression of circMbl leads to the presence of at least one clear ~10 kDa band (which in some cases was accompanied by a slightly heavier MBL-immunoreactive band; see Figure 7A). However, following 12 hr starvation, we observed a clear band of this size also in wild-type flies (Figure 7D), indicating that this protein is either produced and/or stabilized following starvation. Moreover, inducible expression of FOXO in the fly brain for 16 hr (by the use of the CNS-specific GeneSwitch transgene) resulted in the presence of the ~10 kDa band (Figure 7F). Importantly, neither starvation nor FOXO overexpression resulted in any changes in the levels of circMbl1 or the tested mbl mRNA
isoforms (Figure 7E), demonstrating that the effect is post-transcriptional.

**DISCUSSION**

In this manuscript, we present multiple lines of evidence that strongly support circRNAs translation: (1) specific association of circRNAs with translating ribosomes; (2) capacity to generate proteins from circRNA minigenes; (3) finding of RFP reads supporting a stop codon in circMbl; (4) presence of sequences able to promote cap-independent translation on several circRNAs; (5) finding of a peptide matching circMbl3, but no known linear isoform; (6) detection of putative circMbl-encoded proteins by western blot; and (7) determination that physiological factors can regulate the levels of these proteins.

Analyzing circRNA data is difficult because their identification requires exclusive focus on backspliced reads, which cover only a small part of the circRNA molecule. Even for mRNA, the low number of spliced RFP reads prohibits isoform-specific analyses of phasing. The same difficulty arises in identifying circRNA-produced peptides by mass spectrometry. Similarly, evolutionary analysis of cORFs needs to focus on the stop codon, which even if it resides in 5' UTR sequence may well be shared with a putative uORF. Thus, none of the sequence or sequencing-based analyses alone can without ambiguity support circRNA translation. However, taken together and viewed in light of the additional experimental validation presented here, we provide strong evidence of endogenous circRNAs translation. In this context, although it is impossible to quantitatively estimate translation efficiencies of circRNAs from the ratio of backspliced and linear reads, the situation is comparable with mRNA isoforms when restricted to a single, diagnostic junction. Thus, it appears that, once initialized, translation of circRNAs is not much different from translation of mRNAs.

It was recently reported that artificially, strongly overexpressed circRNAs can be translated independently of IRES sequences (Abe et al., 2015). However, our results suggest the existence of a specific sequence to enable translation initiation in a natural, endogenous context. Moreover, the fact that we detected circRNAs with very different abundance associated to ribosomes and that some of our minigenes did not generate proteins strongly suggest that translation of a subset of circRNAs is not by chance, but a specific effect. Importantly, the results obtained using the bicistronic constructs, as well as the in vitro assay, do not necessary mean that the assayed cUTRs have conventional IRES activity. This is because the results obtained using the bicistronic constructs could be caused by the presence of cryptic promoters and/or cleavage and polyadenylation signals, and because circRNAs in the in vitro assay could potentially be cleaved in the extract. However, the fact that in both assays the luciferase signal obtained from constructs carrying the cUTRs is resistant to inhibition of cap-dependent translation demonstrates the capacity of these sequences to promote translation in a cap-independent way, and hence from circular
molecules. Further research is necessary to uncover how these sequences promote translation.

Many of our observations suggest that circRNA translation might be important in the brain. First, circMbl1 and the putative circMbl1-encoded peptide are present in synaptosome fractions. Moreover, ribo-circRNAs are predominantly bound to membrane-associated ribosomes. Importantly, the proteins encoded by the mbl1 and circMbl molecules do not contain identifiable peptide signal sequences, raising the possibility that circMbl and maybe some mbl1 mRNA isoforms are translated in synapses. Indeed, a recent report demonstrated that acute fast- ing regulates synaptic function by a mechanism dependent on 4E-BP and FOXO, both of which can regulate translation from the circMbl cUTR.

In summary, here we provide evidence for translation of circRNAs, an abundant and widespread type of RNA that is distinct from, but shares sequence with, mRNA. Our findings thus indicate the presence of a repertoire of protein isoforms and likely uncharacterized modes of regulation of gene and protein expression.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENTS AND RESOURCE SHARING
- METHOD DETAILS
  - Fly strains and Reagents
  - Molecular Biology Methods
  - Mass spectrometry
  - Computational Methods
- QUANTIFICATION AND STATISTICAL ANALYSIS
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures and seven tables and can be found with this article online at http://dx.doi.org/10.1016/j.molcel.2017.02.021.

AUTHOR CONTRIBUTIONS

N.R.P. and O.B. performed most of the experiments. M.J., R.A.-F., S.S., and M.S. performed the computational analysis. C.S. performed the northern blots. L.R. and M.C. performed the in vitro translation. M.H. performed the synapto- somal purifications. E.W. and M.L. set up the RFP assays. D.P.-H., E.R., and G.D. performed the mass spectrometry. N.R. supervised most of the computational experiments. S.K. wrote the manuscript and designed and supervised the project.

ACKNOWLEDGMENTS

We thank M. Marr for reagents and Y. Arava for help. S.K. is funded by the ERC Consolidator Grant (ERC Grant 647989). M.C., N.R., and L.R. are supported by the Einstein Foundation grant "Single molecule RNA network." E.R. was supported by the Berlin School of Integrative Oncology (grant GSC 1091). M.J. received a fellowship from DFG (GRK 1772). C.S. is funded by BIF. N.R.P. is funded by a JBC fellowship.

REFERENCES


## STAR METHODS

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CONTACT FOR REAGENTS AND RESOURCE SHARING

Further information and request of reagents should be addressed to the Lead Contact, Dr. Sebastian Kadener (skadener@mail.huji.ac.il).

METHOD DETAILS

Fly strains and Reagents

Plasmids

The minigenes constructs for *Drosophila* S2 cells were generated in pMT-V5 (Invitrogen). HCV IRES, MARL bicistronic reporter and 4E-BP plasmids were kindly provided by Mike Marr and previously described (Olson et al., 2013). The pAc-GFP plasmid was previously described (Weiss et al., 2014). Custom DNA fragments synthesis (gBlocks®, Integrated DNA Technologies) was used to generate split cherry exon plasmid construct with circCamk1 flanking introns. The primers used for cloning are described in Table S7.

Generation of UAS-circMbl flies

To generate circMbl overexpression flies we utilized a variant of the pUAS-attB plasmid (Addgene) in which we eliminated the SV40 small intron and replaced the SV40 3' UTR with the one present in the pMT plasmid (Invitrogen). We followed by cloning the circMbl minigene (Ashwal-Fluss et al., 2014) into this plasmid by standard cloning techniques. This plasmid was sent to injection to BestGene Inc (CA, USA).

Fly Strains

WT flies used in this study are w118, yw and CantonS (CS, CSst3H) strain (Bloomington Stock Centre, Indiana). *Drosophila yakuba, Drosophila simulans, Drosophila virilis, and Drosophila pseudoobscura* were obtained from the *Drosophila* Species Stock Center (DSSC) at the University of California, San Diego. Actin-gal4 and UAS-EGFPL10a (III) strains were previously described (Huang et al., 2013; Ito et al., 1997). The UAS-foxo (stock# 9575; (Puig et al., 2003)) was obtained from Bloomington stock center. Elav-Gal GeneSwitch (GS) were generously provided by Dr. Minoru Saitoe, Tokyo Metropolitan Institute of Medical Science, Japan.

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Software and Algorithms

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**Starvation Experiment**
Flies were starved for 12 hr on 2% agar vials, flash frozen in liquid nitrogen and heads were separated using a sieve after mechanical decapitation.

**Gene switch experiment**
6 days old flies were starved for 12 hr and transferred to 5% sugar containing 400μg/ml RU486 food vials to induce the Gal4 expression for 24 hr, flash froze and separated the heads performed western blotting as described below.

**Molecular Biology Methods**

**Cell culture and transfections**

*Drosophila* S2 cells were maintained in 10% fetal bovine serum (Invitrogen) insect tissue culture medium (HyClone). Cells were seeded in a six-well/ twelve-well plate. Transfection was performed at 60%–80% confluence according to company recommendations: 6μl of TransIT 2020 transfection reagent (Mirus Bio, MIR 5400A) and 2μg of total DNA. In the minigene experiments 1 μg of the minigene-expressing vector were used. 0.4μg of 4E-BP was used in all co-transfection experiment. When copper (Cu) induction was performed, 500μM of copper was added to the media and the cells were collected 48h after the induction. FOXO stable cell line was previously described (Puig et al., 2003).

**Western blotting**
Fly heads (20 heads per sample) were collected on dry ice. Heads were homogenized in RIPA lysis buffer (50 mM Tris-HCl at pH 7.4, 150 mM NaCl, 1 mM EDTA, 1% NP-40 0.5% Sodium deoxycholate, and 0.1% sodium dodecyl sulfate (SDS), 1 mM DTT, supplemented by protease inhibitor cocktail and phosphatase inhibitors) using motorized pestle. Head lysates were then centrifuged at max speed for 10 min and the supernatant was saved. Cell samples were collected by centrifugation (500 g 5min), washed twice with PBS and homogenized in RIPA lysis buffer. All lysates were boiled with protein sample buffer (Bio-Rad) and resolved by Criterion XT Bis-Tris gels (Bio-Rad). Antibodies used in this study: Anti-MBL antibody was kindly provided by Prof. Darren Monckton (School of Life Sciences, University of Glasgow). Anti-V5 (Mouse monoclonal V8012, Sigma Aldrich) was used for Immunoprecipitation and immunoblotting. Rabbit polyclonal anti-GFP (ab290, Abcam) and anti α-tubulin (mouse monoclonal DM1A, Sigma, Aldrich) were used for western blotting. All western blot analysis represents at least three independent biological repeats done.

**Northern Blotting**
RNA was treated with RNaseR (3U/μg, 15 min 37˚C) or mock treated and run on denaturing agarose gels (1% formaldehyde) at 90 V for 2:30 hr. The gels were stained with SYBR Gold to ensure equal loading. The RNA was blotted onto a Hybond N+ membrane (GE Healthcare) at 15 V for 1 hr, the membrane was dried and crosslinked (0.12 J/cm²), equilibrated and hybridized (68˚C, o/n) in NorthernMax Hybridization Buffer (Ambion) with 0.1 nM DIG-labeled RNA-probe (max. 150 nt, DIG labeling mix - Roche). The Northern blots were washed stringently (twice 2XSSC, 0.1% SDS and twice 0.2XSSC, 0.1% SDS at 68˚C) developed using an anti-Digoxigenin antibody (Roche), CDP star (Roche) and a LAS4000.

**Polysome sucrose gradient**
S2 cells were co-transfected with the circRNA minigenes. 12 hr later the cells were stimulated with copper and collected for 36 hr. Polysome profiling and RNA isolation from each fraction were done as previously described (Aspden et al., 2014). RNA from the each fraction was mock/ RNaseR treated as described below. In cases in which RNA amounts were lower than 2 μg, an in vitro transcribed Luciferase transcript (see utilized oligonucleotides on Table S6) was added to the RNaseR/ mock reaction.

**RNaseR treatment of total RNA**
Total RNA was RNaseR treated as previously described (Memczak et al., 2013; Suzuki et al., 2006). 2 μg of total RNA were treated with 3U/μg RNaseR (Epicenter) for 15 min at 37˚C or mock treated. The RNA was immediately transferred to ice, spiked with 10% mouse RNA, extracted with TRI Reagent (Sigma, Aldrich) according to the manufacturer’s protocol. The RNA concentration of the mock-treated sample was determined and 5μl of the mock treated RNA was used for reverse transcription (RT, using iScript and random primers, Bio-Rad), the same volume of the RNaseR treated RNA was used for reverse transcription. cDNA was used for qPCR quantification, whereas the spike-in was used for normalization.

**Analysis of gene expression by quantitative real-time PCR**
Total RNA was prepared from adult fly heads and *Drosophila* S2 cells using TRI Reagent (Sigma, Aldrich)) according to the manufacturer’s protocol. RNA was DNase treated (DNasel, NEB) and cDNA derived from this RNA (using iScript and random primers, Bio-Rad) was utilized as a template for quantitative real-time PCR performed with the C1000 Thermal Cycler Bio-Rad. The PCR mixture contained Taq polymerase (SYBR green Bio-Rad). Cycling parameters were 95˚C for 3 min, followed by 40 cycles of 95˚C for 10 s, 55˚C for 10 s, and 72˚C for 30 s. fluorescence intensities were plotted versus the number of cycles by using an algorithm provided by the manufacturer. Primer efficiency was determined for all primers described in this study and incorporated into the relative expression calculation. All the primers used in this assay are detailed in Table S7.

**Protein Immunoprecipitation (IP)**
*Drosophila* S2 Cells (one 10cm plate per sample) were washed twice with cold PBS and lysed in 500μl of lysis buffer (30 mM HEPES KOH at pH 7.4, 100 mM KAcetate, 2 mM MgAcetate, 5% glycerol, 0.1% Triton X-100, 1 mM EGTA, 5 mM DTT, protease inhibitors [Complete mini, Roche]). The lysates were incubated for 10 min on ice and centrifuged at 14,000 rpm for 12 min at 4˚C. 1/10 of the supernatant was used as input. 9/10 of the supernatant was incubated with either anti-v5 or anti-MBL antibodies at 4˚C with rotation, 2 hr later, 50μl of protein G-plus-Agarose beads (Santa Cruz, sc-2002; previously washed with lysis buffer) were added and incubated
one additional hour. After incubation, beads were washed five times for 5 min at 4°C in lysis buffer, before the addition of protein sample buffer and heating of the sample at 95°C for 5 min. As a negative control for the antibody specificity we included in each IP experiment a sample incubated with a normal IgG corresponding to the host species of the primary antibodies used (sheep IgG for α-MBL, sc-2717 and mouse IgG for α-V5, Jackson ImmunoResearch).

**RFP**

For “no detergent” samples, 200 frozen (in liquid N\(_2\)) heads fly heads (head1 and head2) were homogenized in homogenization buffer (20mM HEPES at pH 7.3, 150mM KCl, 5mM MgCl\(_2\), protease inhibitors (Complete mini, Roche), 0.5mM DTT, RNasin 10ul, 100μg/ml CHX). Heads were homogenized immediately using a motorized pestle and centrifuged at 20,000 g for 2 min at 4°C. The supernatant was carefully transferred to a new 1.5 mL tube and stored in −80°C until further use. For the “detergent samples” (Heads 3 and Head4), 200 heads frozen heads and lysed in 500ul of lysis buffer (30 mM HEPES KOH at pH 7.4, 100 mM KAcetate, 2 mM MgAcetate, 5% glycerol, 0.1% Triton X-100, 1 mM EGTA, 5 mM DTT, 0.4 U/μL murine RNase inhibitor (NEB), protease inhibitors (Complete mini, Roche), and 100μg/ml CHX). The lysates were incubated for 10 min on ice and centrifuged at 14,000 rpm for 12 min at 4°C. The supernatant was carefully transferred to new 1.5ml eppendorf tube, flash frozen in liquid nitrogen and stored in −80°C. For isolation of ribosome-protected RNA fragments we followed (Calviello et al., 2016). Briefly, 120ul of the lysate was digested with 3 μl of RNase I (Life Technologies, AM2294) for 45 min at room temperature with rotation. Digestion was stopped by the addition of 4 μl of Superase-In (Life Technologies, AM2694). Meanwhile, MicroSpin S-400 HR columns (GE Healthcare, 27-5140-01) were equilibrated with 3ml of mammalian polysome buffer by gravity flow and emptied by centrifugation at 600 g. We then immediately loaded 100 μl of the digested lysate on the column and eluted by centrifugation at 600g for 2 min. RNA was extracted from the flow-through (approximately 125μl) using Trizol LS (Life Technologies, 10296-010). Ribosomal RNA fragments were removed using the RiboZero Kit (Illuminara, MRZH11124) and separated on a 17% denaturing urea-PAGE gel (National Diagnostics, EC-829). The 27-33nt RNA fragments were cut eluted from the gel and used to generate small RNA libraries. The oligonucleotides utilized are listed on Table S7.

**Ribosome IP (TRAP)**

Ribopo IP (TRAP) was performed as previously described (Huang et al., 2013). The RNaseR treatment was performed in the following way: 100ng of RNA from the immunoprecipitated or input fraction was combined with 1.9 μg of in vitro transcribed 70nt RNAs from a luciferase template using the Megascript T7 Kit (ambion AM1334) according to the manufacturers protocol (see utilized oligonucleotides on Table S7). After performing the mock or RNaseR treatment, we added 100ng of mouse RNA which was used for normalization purposes and proceeded to RT-PCR as described above.

**Fluorescence reporter expression analysis in s2 cells**

24hs after transfection, cells were transferred to optic-suitable 384-well plates and visualized the fluorescence using Scan®R high-throughput fluorescence microscope (Olympus). The intensity from the transfection control reporter (Cherry for circMbl IRES and GFP for the cap-dependent activity experiments), cell size and roundness were evaluated. Cherry or/and GFP levels were assessed only for cell populations that met intensity and cell morphology criteria previously described (Lerner et al., 2015).

**Dual luciferase assay**

Thirty hours of post transfection, cells were assayed using the Dual Luciferase Assay kit (Promega) following the manufacturer’s instructions.

**In vitro translation**

All capped mRNA templates for in vitro translation were generated as previously described (Chekulaeva et al., 2006). For the circMbl-RLuc RNA production, we fused either the complete circMbl UTR region in the reverse / straight orientation / only the 20 most distal nucleotides with the Renilla luciferase or coding sequence by PCR. The resulting PCR fragment was used as a template for in vitro transcription. Transcription, circularization and purification of the circularized RNA by TBE-Urea PAAG electrophoresis were performed according to (Chen and Sarnow, 1995). Preparation of Drosophila embryo extract and in vitro translation assays was done as previously described (Castagnetti et al., 2000; Gebauer et al., 1999). Translation reactions contained 1.5 nM exogenous mRNA or circRNA.

**Synaptoneurosome preparation**

Synaptosome extraction was carried out as previously described (Depner et al., 2014). Briefly, 5ml of fly heads were first crushed into a fine powder with a pre-chilled mortar and ground. The powder was re-suspended in ice-cold homogenization buffer (320 mM sucrose and 4 mM HEPES, pH 7.4) and homogenized with a Polytron 3000 Kinematica homogenizer in 10 pulses. This homogenate was centrifuged (2,700 rpm, 10 min in a SS34 rotor). The supernatant was further centrifuged (11,000 rpm, 15 min in a SS34 rotor). The pellet contains a soft synaptosome layer, which was collected, and a hard mitochondria core. Samples were taken from all steps for RNA purification.

**Mass spectrometry**

**Collection of the reference spectra**

The synthetic heavy labeled RAADTTDMPLM (SpikeTides, JPT Inc.) was resuspended in 50% acetonitrile, 0.1% formic acid. The spectra were recorded using a Q-Exactive plus mass spectrometer (Thermo Scientific) using higher energy collision dissociation method (HCD) with a mass resolution of 70000 for the MS and 35000 for the MS/MS scans. The recorded spectra were analyzed using the MaxQuant software package (version 1.5.8) using a custom made database, with carbamidomethylation of cysteins as a fixed and oxidation of methionines as a variable modification. For peptides and proteins a FDR of 1% was applied.
SRM-Mass Spectrometry

Based on the fragmentation pattern of the RAADTTDMFPLIM peptide in the Q-Exactive plus mass spectrometer a SRM method for a Q-TRAP 6500 was developed monitoring the b8, b9 and b11 fragment ions.

Proteins bound on the antibody resin were released from the beads by a treatment with 1% formic acid (Merck) for 15 min at RT. The extract was then neutralized at pH = 8.0 with 1M ammonium bicarbonate (Sigma, Aldrich). Disulfide bonds were first reduced with 10 mM TCEP (tris(2-carboxyethyl)phosphine) (Sigma, Aldrich) for 20 min and then cysteine groups were alkylated with 55 mM 2-chloroacetamide (Sigma, Aldrich) for 30 min on an automated setup (Karashova et al., 2015). The proteins were digested with endopeptidase Lys-C (Wako). Peptides were extracted, desalted and stored on reversed-phase (C18) StageTips (Rappsilber et al., 2007). After elution, the peptides were lyophilized and resuspended in 3% TFA and 5% acetonitrile in preparation for LC-MS analysis. The heavy-labeled RAADTTDMFPLIM internal standard peptide (SpikeTides, JPT Inc.) was spiked into the solution at a concentration of 200 fmol per microliter of sample. Peptides were then separated on an in-house packed 20 cm reversed-phase column (75 μm inner diameter, 3 μm Reprosil C18-beads, Dr. Maisch) on a gradient from 3% to 32% acetonitrile in 90 min, and detected by a Q-Trap 6500 (AB Sciex). SRM signals were analyzed using the Analyst 1.6 software package (AB Sciex).

List of the transitions monitored for the identification of the peptide RAADTTDMFPLIM

<table>
<thead>
<tr>
<th>PRECURSOR MASS</th>
<th>TRANSITION MASS</th>
<th>DWELL TIME</th>
<th>FRAGMENTATION ID</th>
<th>COLLISION ENERGY</th>
<th>RETENTION TIME</th>
</tr>
</thead>
<tbody>
<tr>
<td>744.85</td>
<td>1219.57</td>
<td>30</td>
<td>b11_heavy</td>
<td>37.736</td>
<td>49 min</td>
</tr>
<tr>
<td>1009.43</td>
<td>862.36</td>
<td>30</td>
<td>b9_heavy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>741.35</td>
<td>1219.57</td>
<td>200</td>
<td>b11</td>
<td>37.604</td>
<td>49 min</td>
</tr>
<tr>
<td>1009.43</td>
<td>862.36</td>
<td>200</td>
<td>b9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>862.36</td>
<td></td>
<td></td>
<td>b8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Computational Methods

ORFs Prediction

To predict the cORFs we utilized circRNAs dataset generated from flies (Ashwal-Fluss et al., 2014), mouse and human (obtained from circBase.org; Glazar et al., 2014). Analysis was performed with only annotated circRNA (in which both junctions of the circRNA are matched to known annotated exons). Each circRNA sequence, excluding intronic sequences, was multiplied four times, and the longest ORF spanning the circRNA junction was selected for each one of the three frames (minimum cORF length threshold was 20aa). In the ribo-circRNA analysis we used the longest non-infinity (“moebius”) cORF detected in any frame. We generated the control dataset by randomly selecting junctions from genes not hosting circRNAs and correcting to obtain a similar 5' bias to the circRNA data.

cORF selection

We scored all possible ORFs in a circRNA by the following function:

\[ S = 100 \times KNOWN\_START + 1000 \times HEAD\_TO\_TAIL + \text{len(aa)} / 10 + 10,000 \times MOEBIUS\_ORF \]

KNOWN_START is 1 if the start codon coincides with the annotated start of the mRNA ORF

HEAD_TO_TAIL is 1 if the cORF spans the head-to-tail junction

MOEBIUS_ORF is 1 if the cORF has no stop codon and thus theoretically never terminates

The expression of circRNA in total-RNA seq data

In order to assess the ribo-circRNA expression levels in total RNA-seq data we applied the find_circ pipeline (Memczak et al., 2013) on the relevant RNA-seq datasets (for C2C12 and Kc167 cells, see STAR Methods table) or used an available circRNA expression data. For fly and S2 cells we used our recently published data (Ashwal-Fluss et al., 2014). For embryo we used data published in (Westholm et al., 2014).

Start/Stop scoring

From this, for each experiment, we compute a scoring-matrix for the vicinity of start and stop codons as follows:

\[ Q_{rl} = \log(f_{rl}) - \log(1 / R \Sigma_{r} f_{rl}) \]

with r being the relative position, I the length of the read, and R the number of positions taken into account. Thus we compare the specific occurrence frequency at each position to a mean, uniform background frequency, for each read length.

An observed pattern of reads is then scored as follows:

\[ S_{start/stop} = \sum Q_{rl} + O_{rl} \]
where $O_{rl}$ is the observation matrix, containing the number of observed reads of length $l$ at relative position $r$ to the start or stop codon.

**ORF-scoring**

The ORF score is slightly more complicated, because each read can not only support translation in one or more frames, but can also reduce support for translation in other frames. As a simple example, most RFP experiments show a clear preference of 28nt reads to occur in frame 1 relative to the start codon. Therefore, if we observed many such reads in frame 2 of a candidate region, this would lend strong support that the scored region is actually translated in frame 2 and weaken any support for frames 1 and 3. Thus, we need to properly sum up all evidence contributed by each read, even if occurring in an adjacent frame. First, the scoring matrix is computed completely analogous to $Q_{rl}$ with the frame replacing the relative position (also denoted by $r$). The same is true for the observation matrix $O_{rl}$, where we take the modulo of each RFP 5’ position in the candidate region with three to derive the frame $r$. Then, we compute one aggregate score for each frame of the candidate region:

\[
\begin{align*}
Q_1 &= \Sigma_l (O_{1, l} + Q_{1, l} + O_{2, l} + Q_{2, l} + O_{3, l} + Q_{3, l}) \\
Q_2 &= \Sigma_l (O_{2, l} + Q_{1, l} + O_{3, l} + Q_{2, l} + O_{1, l} + Q_{3, l}) \\
Q_3 &= \Sigma_l (O_{3, l} + Q_{1, l} + O_{2, l} + Q_{2, l} + O_{1, l} + Q_{3, l})
\end{align*}
\]

(the ORF-score matrix plots show everything on the right before being summed over $l$, i.e. the contribution of each length to translation in frame 1, 2, 3).

The highest score determines the most likely frame in which the region is translated:

\[
f_0 = \arg\max \{Q_1, Q_2, Q_3\}.
\]

We asserted that the vast majority of known ORFs with at least 10 RFP reads is assigned the correct frame 1 relative to the CDS (not shown).

**Negative controls and significance estimate**

We implemented two different negative controls. First, we randomly permuted the scoring matrix for each ORF that was scanned. This strongly abrogated the scores assigned to known Start/Stop codons or ORFs. However, since the scoring matrix still contains the same values, rare, large scores can be observed. Second, we randomly assign reads to a frame, ignoring its actual position relative to the candidate region, but keeping the length frequency distribution and the scoring-matrix intact.

This shuffling is very fast and can be used to compute an empirical $P$ value as the fraction of shuffled scores that reaches or exceeds the observed, non-shuffled score.

**Backspliced reads**

When scoring a putative circRNA start/stop codon or ORF, we extract the RFP alignments to the back-splice junction of the circRNA in question and convert the coordinates of each reads 5’ end to relative coordinates in the ORF. Then, scoring proceeds as explained above. Optionally, scoring can be performed exclusively for back-spliced alignments, excluding any possible interference with mRNA-derived RFP reads.

**Multiple species alignments**

We downloaded the 24way *Drosophila* alignment and 60 way mouse MAF-blocks from UCSC (Kent WJ, Sugnet CW, Furey TS, Roskin KM, Pringle TH, Zahler AM, Haussler D). The human genome browser at UCSC (Blanchette et al., 2004; Kent et al., 2002). Next we obtained the full genomic sequences of all used insect genomes. We then used a combination of indexing and custom scripts (to be published elsewhere) to extract and splice together the orthologous sequences to the exons in a dm6 *Drosophila melanogaster* or mm10 *Mus musculus* transcript model. The set of all obtained sequences was run through MUSCLE (Edgar, 2004) to rebuild a full multiple sequence alignment.

**Stop codon scoring**

We used our multiple-species alignment code to extract specifically the stop codon sequences, aligned to all known stop codons in the ENSEMBL gene annotation for dm6, and, separately mm10, downloaded from UCSC (Karolchik et al., 2004). We repeated this analysis for all stop codon sequences occurring in internal (not first) 5’UTR exons from the same transcript catalog, recording their distance from the 5’ splice site. From this data we computed for each aligned species the frequency with which the given stop codon in the dm6 reference (one of UAA, UAG, or UGA) is replaced by another, or a non-stop triplet. Thus, the 5’UTR stops yield position-dependent background models for the evolution of 5’UTR exonic sequence that is intrinsically tuned to evolutionary branch length and species-specific changes in sequence composition. Comparing these background frequencies to the observed substitution matrices for annotated stop codons yields a scoring matrix (the log-ratios), analogous to our approach for scoring RFP read patterns. We build separate matrices for splice-site distances up to 10nt. After this distance we accumulated all data to build a model for stop codons 11nt or further away from the 5’ splice site. Next, we selected from our circRNA sites only those with cORFs composed of coding and 5’UTR exonic sequence, where the stop codon resides in 5’UTR sequence, such that our model would be applicable. We then ran a script that uses these scoring matrices to score all stop codons of cORFs and the various controls we generated. For each species, the observed substitution or conservation is scored with the log-ratio matrix. Positive log-ratios indicate that the sequence fits the evolution of annotated stop codons, negative scores indicate that the sequence fits better to the 5’UTR background model. Scores are added for all species for which there is an alignable orthologous sequence.
**Domain Analysis**

We used ScanProsite tool (http://prosite.expasy.org/scanprosite) to find domains in the predicted cORF sequences. We used only cORF that contains STOP codon. As a control set we used ORF predicted from randomly selected exons obtained from the same hosting genes. We count the number of occurrence of each domain in the two datasets and select the ones enriched in the cORF dataset comparing to the random control (found at least 4 times more in the cORF).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

The statistical analysis as well as the number of repeats are stated in the figure legends as well as in the STAR method sections. For the meta-analysis described in this paper (in Figures 3, 5, S3, and S5) we used Fisher’s exact test or Mann-Whitney U test as indicated in the figure legends.

**DATA AND SOFTWARE AVAILABILITY**

All raw sequencing data are available at NCBI GEO: GSE79626. The cORF prediction algorithm has been deposited in https://github.com/kadenerlab/cORF_pipeline. The SRCP algorithm to detect circRNAs from RFP reads can be provided upon request.
Supplemental Information

Translation of CircRNAs

Figure S1 related to Figure 2

A. Minigene

B. Input

C. Control

D. 

E. 

F. 

G. 

H. 

I. 

Figure S6

A. 

B. 

C. 

D. 

E. 

F. 

G. 

H. 

I.
Figure S2 related to Figure 2
Figure S3 related to Figure 3

A. No Detergent RFP

B. Normal Detergent RFP

C. ORF Score

D. START and STOP Scores

E. No Detergent

F. Normal Detergent
Figure S4 related to Figure 4

A. circRNA RFP reads in Normal Detergent Samples

B. Fly Heads

C. RFP reads in circRNA/linear junctions

D. Venn diagram of Fly heads

E. RNaseR+/Mock Expression

F. Fly Head MBL immunoprecipitate
Figure S5 related to Figure 5

A. Observed stop distance [codons] for 5' UTR exons

B. CircRNA junction

C. STOP distance from splice site is geometrically distributed

D. Cumulative frequency

E. # Found in cORF

- ZF C3H1
- CNMP binding
- LAM G DOMAIN
- HMG BOX 2
- MHD1
Table 1: Summary of circRNA reads and footprints

<table>
<thead>
<tr>
<th>Species</th>
<th>Circ/Linear Reads</th>
<th>No cORF (%)</th>
<th>One (%)</th>
<th>Two (%)</th>
<th>Three (%)</th>
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</thead>
<tbody>
<tr>
<td>Rat</td>
<td>0.4</td>
<td>8.3%</td>
<td>62.3%</td>
<td>1.1%</td>
<td>30%</td>
</tr>
<tr>
<td>Mouse</td>
<td>0.2</td>
<td>5.2%</td>
<td>60.6%</td>
<td>1.2%</td>
<td>30%</td>
</tr>
</tbody>
</table>

Figure 7

A. Rat

- One 62.3%
- Two 31.3%
- No cORF 5.2%
- Three 1.2%

B. Rat

- circRNAs with footprinting Reads
- Rank Expression
- Log # Reads

C. Mouse

- circRNAs with footprinting Reads
- Rank Expression
- Log # Reads

D. Rat

- Circ/Linear Reads vs Number circRNA reads

E. Mouse

- Circ/Linear Reads vs Number circRNA reads

F. Rat

- % in Group
- Control
- ribo-circRNAs

G. Mouse

- % in Group
- 5'UTR
- 5'UTR-CDS
- CDS
- 3'UTR-CDS
- 3'UTR

H. Mouse

- % cORFs that share AUG with hosting mRNA

- Control
- ribo-circRNAs

Rat

- % cORFs that share AUG with hosting mRNA

- Control
- ribo-circRNAs
Figure S7 related to Figure 6

A. 

B. 

C. 

D. 

E. 

F.
SUPPLEMENTAL FIGURE AND TABLE LEGENDS:

Legends for Figure S1-S7:

Figure S1 related to Figure 2:
A. Non-ambiguous circRNAs foot-printing reads are in the same scale as the ones in junctions of the same exons. B. Both anti-V5 and anti-MBL antibodies recognize and precipitate (IP) the protein produced from circMblV5. C. Scheme of the probe design for the detection of circRNA and linear molecules by Northern Blot. D-F. Northern blot assay for detecting circMbl, circMbl 5'ss and circPde8 in transfected Drosophila S2 cells. RNA was treated with RNaseR or mock treated before running the gels. Blotting was done using probes for the relevant circRNA junctions or exons (which detect both the linear and circRNA molecules). Concatamer products migrate higher due to their size and are sensitive to RNaseR treatment. G. RT-PCR analysis for RNA extracted from Drosophila S2 cells, transfected with different minigenes, +/- RnaseR treatment. Gene expression was normalized to an endogenous circRNA. Data is presented as mean ±SD (n=3). H-J. Northern blot assay for detecting circCamKI in transfected Drosophila S2 cells. In H and I, three circRNA products originate from the minigene due to alternative splicing of the internal introns.

Figure S2 related to Figure 2:
Polysome profile analysis of Drosophila S2 cells transfected with plasmids expressing the circMblV5, circCdIv5 and circCamKIV5 minigenes, a plasmid driving expression of GFP and a plasmid driving expression of a circRNA in which a split Cherry protein is under the control of the putative circMbl IRES (see Figure 1F). A. RT-PCR analysis of the different fractions of a polysome gradient. The results are the average of three biological replicates and are expressed as fraction of the RNA target over the gradient. gadph mRNA amplified from a mouse RNA spike-in was used for normalizing between samples. Error bars indicate standard error of the mean (SEM). B. RT-PCR analysis of the different fractions for the indicated targets. The results are the average of three biological replicates and are expressed as fraction of the RNA target over the whole gradient. We utilized mouse gadph mRNA amplified from a mouse RNA spike-in for normalizing between samples across the gradient. Error bars indicate standard error of the mean (SEM). C-F. RT-PCR results showing the RNaseR sensitivity/resistance of the indicated RNA targets across the polysome gradients. The RNaseR sensitivity was obtained by calculating the RNaseR/Mock ratio for each target in each fraction. The data was normalized to mouse gadph mRNA; we added a mouse RNA spike to each fraction. To correct for the different efficiency of the RNaseR treatment across samples and fractions we used the GFP mRNA. Outliers were excluded for normalization and all the values are the average of at least 2 biological replicates.

Figure S3 related to Figure 3:
A. Top: Metagene profile of RFP read phasing analysis for no detergent samples. First (left) the overall frequency with which a read 5' end falls into a particular frame, relative to the annotated start codons. Next (middle), these frequencies subdivided by read length, represented by color in a heatmap plot. Last (right), the log-ratio scores of the position and length-dependent frequencies to a background model with uniform distribution across frames. Red indicates overrepresented, blue underrepresented frame/length combinations. Bottom: The highly expressed Adh gene as an example. Last panel contains the aggregate support each length-group of reads gives to the three possible frames relative to the annotated start. The correct frame1 receives very strong support. B. Same as A but for detergent containing lysis buffer conditions. C. Boxplots showing the distribution of aggregate log-ratio scores for translation (ORF-score) of known ORFs (ORF CDS). Negative controls are scores of the same ORFs when randomly permuting the scoring matrix for each ORF (ORF perm), or randomly shuffling the position of each read (ORF shuffle). Red line indicates the median and blue dot the mean. Box shows interquartile range. Whiskers show 5th to 95th percentile range. Outliers are omitted. Significantly different medians are indicated by stars *: P < 0.05; **: P < 0.01; ***: P < 0.001; ****: P < 0.0001; two-sided Mann-Whitney-U test. D. Box plots indicating the distribution of aggregate log-ratio score for start or stop codons. E. Meta-analysis of the RFP reads in the proximity of the start (top) and stop (bottom) codons of all annotated genes. The no detergent samples were used for these graphs. From left to right: the total frequency of RFP read 5' end positions plotted against the relative distance (in nucleotides) to the start/stop codon (Left). The same information, but subdivided into reads of different length, with frequency represented by color (Middle). Relative enrichment of RFP reads around real start/stop codons over the background frequency for reads of such length in the vicinity of start/stop codons, expressed as log-ratio of these frequencies (Right). Positive scores (red) indicate consistency with the signature of start/stop codons, negative scores (blue) with the uniform background distribution. E. Scatter plot showing the reproducibility of ORF scoring between biological replicates.
The ORF log-ratio scores for the coding sequence of each annotated gene are highly correlated between replicates (rho=Spearman rank correlation, R=Pearson R).

Figure S4 related to Figure 4:
A subset of circRNAs is associated with translating ribosomes. A. Comparison of RFP backsplicing reads between the FRP libraries prepared from with and without detergent samples. B. Backsplice RFP reads originate from circRNAs expressed at different levels of expression. Data from fly heads. C. Graph comparing the number of RFP reads on circRNA junctions with the number of observed reads in the linear junctions of the hosting mRNA. In the y-axis we plotted for each ribo-circRNA the ratio between the number of backspliced RFP reads and the highest number of flanking junction reads of the linear mRNA. D. Overlap of ribo-circRNAs between the RFP fly head and cell lines datasets. E. RT-PCR results showing RNaseR sensitivity (expressed as the ratio between the expression values in the RNaseR and mock-treated samples) for the candidate ribo-circRNAs. Values were normalized to mouse gapdh mRNA. Equal amounts of mouse spike-in RNA were added to the sample before the RT reaction. Previously validated circRNAs (left) were used as negative (resistant) controls and rp49 mRNA as a positive (sensitive) control. F. Detection of the RAADTDMFPLIM peptide by selected reaction monitoring. The uppertrace shows the elution profile for the three monitored transitions on a Q-TRAP 6500 mass spectrometer (b8-red, b9-green and b11-blue). Bottom: chromatographic profile of the native RAADTDMFPLIM peptide from the MBL IP experiment. Concentration of the peptide is 80 pmol.

Figure S5 related to Figure 5:
Computational analysis of ribo-circRNAs. A. ribo-circRNAs are longer than utc-circRNAs as well as control and all circRNAs. Boxplots showing the distribution circRNA spliced lengths, assuming all introns are spliced out. Negative controls (“ctrl”) are consecutive non-circRNA exons that are selected to match the spliced-length distribution of all circRNAs (“all”). “ribo”/“utc” are circRNAs with a cORF and with/without ribo-seq back-spliced read support. Red line indicates the median and blue dot the mean. Box shows interquartile range. Whiskers show 5th to 95th percentile range. Outliers are omitted. Significantly different medians are indicated by stars: *: P < 0.05; **: P < 0.01; ***: P < 0.001; ****: P < 0.0001; two-sided Mann-Whitney-U test. B. Validation of the circMbI expression in D. simulans, D. Yaacuba, D. pseudooscura, D. virilis using Sanger sequencing of the head to tail junction. Black arrow indicates the junction. The stop codon is highlighted in a pink box. C. The distribution of distances between 5′ splice site and the closest stop codon is well approximated by a geometric distribution. Black dots: observed stop distances for all internal (not first) Drosophila melanogaster 5UTR exons. Red line: geometric distribution with same mean. Insert: zoom into the first 5 nucleotides shows frame-specific deviation from the geometric distribution due to overlap with the exonic splice site motif. D. Distribution of stop codon conservation scores for different sets of cORF stop codons and controls shown as cumulative relative frequency plots. Prefix “rand” means randomly selected from the 5′UTR sequences also contained in the corresponding cORFs. Prefix “closest” means, control stop codons, selected for minimum distance to cORF stop, ignoring frame. Suffix “uORF” indicates that stop codon also terminates a possible uORF. “Known” are stop codons from the annotated mRNA ORFs corresponding to the circRNAs. “n.s.” for no significant difference of the medians by double-sided Mann-Whitney U test at P=5% cutoff. E, Proteins domains enriched in cORFs and present in ribo-circRNAs. We used ScanProsite to predict the domains from cORF predicted proteins. The same analysis was applied on exons selected randomly from the same hosting genes and was used as a control for filtering out randomly found domains. The most frequent domains found in cORF predicted proteins after filtering is presented. The y-axis indicates the number of those domains found in the cORF group.

Figure S6 related to Figure 5:
Analysis of cORFs in mammals. A. Top: Number of predicted cORFs per circRNA in rat circRNAs. Bottom: Number of predicted cORFs per circRNA in mouse circRNAs. B, C. Backsplice RFP reads originate from circRNAs expressed at different levels of expression in rat brain (A) and C2C12 cells (B). D. Non-ambiguous circRNAs foot-printing reads from rat brain are in the same scale than the ones in junctions of the same exons. E. Non-ambiguous circRNAs foot-printing reads in C2C12 cells are in the same scale than the ones in junctions of the same exons. F, G. Rat and Mouse ribo-circRNAs are strongly enriched for 5′ UTR and CDS overlap compared to control exons. H. Rat and Mouse ribo-circRNAs are more likely to start their cORF in the AUG of the hosting/linear mRNA than control.

Figure S7 related to Figure 6:
Ribo-circRNAs have IRES sequences. A. Western blot analysis for Drosophila S2 cells, transfected with pAcGFP and pMT-circPde8V5 with or without the 4E-BPV5 expressing plasmid. The V5 blot recognizes both the protein originated from the circPde8V5 minigene, which shows two bands (expected sizes: 14.5 and 17KD) and the 4E-BP (17 KD) that masks the higher Pde8 band. B. Firefly luciferase levels of the bicistronic carrying the putative circPde8 IRES in the straight or inverted orientation. The levels of S2 cells are presented as background luminescence. C. RT-PCR for firefly luciferase in cells transfected with the different bicistronic constructs. N=3. Error represents standard error of the mean (SEM). Data was normalized to the endogenous rp49 mRNA. D. Schematic representation of circMbl reporter used in cell-free translation. The UTR region of the circMBL was fused with the Renilla coding sequence and circularized. E. RT-PCR analysis examining circular and linear mbl transcript levels in response to starvation in circMbl OE flies. Data is presented as mean ±SD (N=2). Data were normalized to rp49 and 28S rRNA. F. Levels of foxo mRNA, mbl pre and mRNA and circMbl from Drosophila S2 cells at different times following foxo expression. The measurements were performed by RT-PCR at the indicated times after foxo induction by addition of copper (MT-FOXO stably transfected cell line). N=3. Error represents standard error of the mean (SEM).

Legends for Tables S1-S7

Table S1 related to Figure 1. List of cORFs in Drosophila melanogaster.
Table S2 related to Figure 3. List of Drosophila ribo-circRNAs.
Table S3 related to Figure 5. Evolutionary Conservation scores for Drosophila cORFs and ribo-circRNAs.
Table S4 related to Figure 5. Proteins Domains found on ribo-circRNAs.
Table S5 related to Figure 5. Summary of back-spliced reads detected in C2C12 ribosomal footprinting data (PRJEB7207).
Table S6 related to Figure 5. Summary of back-spliced reads detected in rat samples ribosomal footprinting data (GSE66715).
Table S7 related to Figures 1-7. List of oligonucleotides utilized in this study.