Unique repression domains of Pumilio utilize deadenylation and decapping factors to accelerate destruction of target mRNAs

René M. Arvola\textsuperscript{1,2}, Chung-Te Chang\textsuperscript{3}, Joseph P. Buytendorp\textsuperscript{1}, Yevgen Levdansky\textsuperscript{3}, Eugene Valkov\textsuperscript{3}, Peter L. Freddolino\textsuperscript{2,4} and Aaron C. Goldstrohm\textsuperscript{1,*}

\textsuperscript{1}Department of Biochemistry, Molecular Biology and Biophysics, University of Minnesota, Minneapolis, MN 55455, USA, \textsuperscript{2}Department of Biological Chemistry, University of Michigan, Ann Arbor, MI 48109, USA, \textsuperscript{3}Department of Biochemistry, Max Planck Institute for Developmental Biology, Max-Planck-Ring 5, 72076 Tübingen, Germany and \textsuperscript{4}Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, MI 48109, USA

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ABSTRACT

Pumilio is an RNA-binding protein that represses a network of mRNAs to control embryogenesis, stem cell fate, fertility and neurological functions in \textit{Drosophila}. We sought to identify the mechanism of Pumilio-mediated repression and find that it accelerates degradation of target mRNAs, mediated by three N-terminal Repression Domains (RDs), which are unique to Pumilio orthologs. We show that the repressive activities of the Pumilio RDs depend on specific subunits of the Ccr4–Not (CNOT) deadenylase complex. Depletion of Pop2, Not1, Not2, or Not3 subunits alleviates Pumilio RD-mediated repression of protein expression and mRNA decay, whereas depletion of other CNOT components had little or no effect. Moreover, the catalytic activity of Pop2 deadenylase is important for Pumilio RD activity. Further, we show that the Pumilio RDs directly bind to the CNOT complex. We also report that the decapping enzyme, Dcp2, participates in repression by the N-terminus of Pumilio. These results support a model wherein Pumilio utilizes CNOT deadenylase and decapping complexes to accelerate destruction of target mRNAs. Because the N-terminal RDs are conserved in mammalian Pumilio orthologs, the results of this work broadly enhance our understanding of Pumilio function and roles in diseases including cancer, neurodegeneration and epilepsy.

INTRODUCTION

Proper control of gene expression is accomplished in part by RNA-binding proteins (RBPs) that affect processing, transport, translation, and degradation of messenger RNAs (mRNAs). \textit{Drosophila} Pumilio (Pum) is a quintessential sequence-specific RBP that regulates the fate of mRNAs in the cytoplasm. Pum is a member of the eukaryotic PUF family (named after Pum and \textit{C. elegans} fem 3-binding factor), which share a conserved Pum homology domain (Pum-HD) (1). Pum is essential for development and impacts a wide range of biological processes (2). Pum is broadly expressed and is abundant in embryos, the nervous system, and the female germline. During early embryogenesis, Pum represses expression of the morphogen Hunchback, a crucial factor in the establishment of polarity and body plan (3–9). In the germline, Pum regulates stem cell proliferation and differentiation (10–14). Moreover, Pum plays multiple roles in the nervous system, where it controls neuronal morphology, electrophysiology, motor function, and learning and memory formation (15–21).

Pum regulates specific mRNAs by binding to a short RNA sequence, 5′ UGUANAUA, termed the Pumilio Response Element (PRE), via its RNA-binding domain (RBD) that encompasses the Pum-HD and flanking residues (2,5,22–25). The RBD is comprised of eight repeats of a triple alpha-helical motif which form an arched molecule that recognizes single-stranded RNA (25,26). Each repeat presents three amino acids that specifically interact with a ribonucleotide base. Pum binds to an extensive network of mRNAs, the majority of which contain one or
more PREs located in the 3′ untranslated region (3′UTR) (2,5,27–29).

Notwithstanding substantial insights into Pum’s biological roles, structure, and RNA-binding activity (2), our understanding of the mechanisms by which it represses gene expression remains incomplete. An early model proposed that Pum recruits Nanos (Nos) and Brain tumor (Brat) to block translation of hunchback mRNA (30–32); however, recent developments have substantially revised that model. We now know that Pum, Nos, and Brat are each sequence specific RBPs that can combinatorially regulate a subset of mRNAs (2,25,28,33,34). Nos can bind in a cooperative manner with Pum to certain mRNAs that contain a Nos Binding Site (NBS) immediately upstream of a PRE, thereby strengthening Pum-mediated repression (25). Additionally, Brat was shown to bind specific mRNAs on its own and confers repressive activity independent of Nos or Pum (28,33,34). In the case of the hunchback mRNA in embryos, Brat, Pum and Nos collectively repress it by binding to two Nos Response Elements (NREs), each of which contain a Brat binding site, an NBS and a PRE (2,25,28,33–35). Importantly, Pum can repress PRE-containing mRNAs independent of Nos or Brat (36). For example, Pum potently represses PRE-bearing reporter mRNAs in cultured Drosophila d.mel2 cells that do not express detectable Nos. Moreover, depletion of Nos and/or Brat did not alter Pum’s ability to repress. Further, Pum can repress mRNAs that are not bound by Nos or Brat. In this study, we focus on determining the mechanism by which Pum represses mRNAs. The resulting knowledge will be essential to understand how Pum regulates its multitude of targets and how it collaborates with other RBPs, such as Nos and Brat, to regulate subsets of those mRNAs.

Multiple studies have provided insights into the mechanism of Pum-mediated repression. Early evidence correlated repression of hunchback mRNA by Pum—along with Nos and Brat—during embryogenesis with shortening of that transcript’s 3′ poly-adenosine (poly(A)) tail (i.e. deadenylation) (8,35). The poly(A) tail promotes translation and stability of mRNAs, and deadenylation reduces protein expression and initiates mRNA decay (37,38). Like all eukaryotes, Drosophila possesses multiple deadenylase enzymes (39–41). Pum was reported to interact with the Ccr4–Not (CNOT) complex (42–44), which contains both Pop2/Caf1 and Ccr4/twin deadenylases.

Pum also cooperates with Nos or Brat in other contexts, and again deadenylation is implicated. In the germline, Pum and Nos regulate cyclin B (cycB) in pole cells and mei-P26 mRNA in germline stem cells (GSCs) (42,43). In both cases, Pum and Nos are thought to utilize the CNOT deadenylase complex. Pum and Brat regulate targets in the cytoskeleton to attenuate the local effects of Dpp signaling, and this effect is thought to require CNOT, as the Pop2 deadenylase was necessary for Pum and Brat to repress a reporter bearing the mad 3′UTR (11). In terms of the Pum repression mechanism, a complication in interpreting these experiments is that Nos and Brat are also linked to CNOT and deadenylation (40,45,46). Thus, it was necessary to develop approaches that specifically dissect repression of mRNAs by Pum alone.

We previously used PRE-containing reporter genes to measure Pum repression activity in Drosophila cells and showed that it reduces both protein and mRNA levels (36). Four regions of Pum contribute to its repressive activity. The highly conserved RBD made a minor contribution, whereas the N-terminus of Pum contains the major repressive activity. Repression by the Pum RBD required a poly(A) tract in the target mRNA and the cytoplasmic poly(A) binding protein (pAbp) (44). The Pum RBD associates with pAbp and antagonizes its ability to promote translation. The Pum RBD also interacts with Pop2 and promotes deadenylation (42,44). While depletion of Pop2 and Ccr4 blocked mRNA decay induced by the Pum RBD, it did not prevent RBD-mediated translational repression, whereas pAbp depletion did. Thus, the Pum RBD appears to primarily act via inhibition of poly(A)-dependent translation.

The robust repressive activity of the Pum N-terminus is conferred by three repression domains (Figure 1A, RD1, RD2 and RD3) (36). These RDs are unique to Pum orthologs spanning from insects to vertebrates (47). They do not share homology with each other or previously characterized protein domains. Each is capable of repressing protein expression when directed to a reporter mRNA (36). The crucial remaining challenge is to determine how the Pum N-terminal RDs regulate target mRNAs. In this study, we characterize their regulatory activities and investigate the co-repressors necessary for repression.

An earlier model proposed that the 5′-7-methylguanosine cap of target mRNAs is important for Pum repression (48). The 5′ cap plays a key role in translation and mRNA stability, and its enzymatic removal (i.e. decapping) initiates 5′ mRNA decay (37,38). Analysis of the Xenopus Pum ortholog identified a 5′ cap-binding motif that contributes to cap-dependent translation inhibition in oocytes (48). Because this motif is conserved in Drosophila Pum (11,48), it was postulated to contribute to translational inhibition; however, conflicting data have been reported. First, deletion of the putative cap-binding region (PCMB) did not alleviate Pum repression, and the cap-binding region did not display repression activity when directly tethered to an mRNA (36). In contrast, another study reported that mutation of a conserved tryptophan in the cap-binding motif reduced Pum activity (11). In this study, we further scrutinize the potential contribution of this motif.

A previous study reported repression by Pum and Nos of reporter genes whose translation is driven by either 5′ cap-dependent or independent translation using a Drosophila eye phenotypic assay (49). There are several important caveats to that analysis. First, the internal ribosome entry site that was used remains poorly characterized (50,51). Second, the contribution of mRNA decay was not assessed. Third, the experimental system could not separately analyze contributions by Pum and Nos. Therefore, the potential relevance remains unknown.

In this report, we show that Pum accelerates mRNA degradation of PRE-bearing mRNAs, mediated by N-terminal repression domains. We find that the putative 5′ cap binding motif is not necessary for Pum repression. Instead, the Pum RDs directly bind to the CNOT deadeny-
Figure 1. Pumilio accelerates mRNA degradation. (A) Diagram of *Drosophila melanogaster* Pumilio (Pum) protein with N-terminal (N) Repression Domains (RD1, RD2 and RD3) and C-terminal RNA-Binding Domain (RBD). Amino acid boundaries are listed at the top. Amino acid substitutions used in this study, including the putative cap-binding amino acid (W783G) and the RNA-binding defective mutant repeat seven (mut R7), are annotated below the diagram. (B) Diagram of nano-luciferase (Nluc) reporter mRNAs containing three Pum Response Elements (3× PRE) sequences in 3′ UTR, along with 7-methyl guanosine cap (m7G) and poly(A) tail (pA). An equivalent Nluc reporter lacking the PRE sequences (∆PRE) was used as a control. Diagram is not to scale. (C) Transcription shut-off with Actinomycin D (ActD) was performed to compare the half-lives of the Nluc 3× PRE and Nluc ∆PRE reporter mRNAs in transiently transfected d.mel2 cells. A representative northern blot of Nluc reporters and the 18S ribosomal rRNA internal control is shown. Each lane of the gel contains 10 μg of total RNA. The measured half-life of each reporter mRNA is shown below the respective blots. Mean values from three experimental replicates are reported with 95% credible intervals. Data and statistics are reported in Supplementary Table S1. (D) The fraction of Nluc mRNA remaining, normalized to internal control 18S rRNA, is plotted relative to time (minutes) after inhibition of transcription. Datum points for each of three experimental replicate are plotted. First order exponential decay trend lines, calculated by non-linear regression analysis, are plotted for each experimental condition (red for Nluc ∆PRE and blue for 3× PRE). (E) RNAi-mediated depletion of Pum or Pop2 mRNAs after 4 days of dsRNA treatment was measured by RT-qPCR. The Pum or Pop2 mRNA level was normalized to internal control Rpl32 mRNA and fold change was calculated relative to the non-targeting control RNAi condition (NTC). The mean log2 fold change of the indicated mRNA level is plotted with 95% credible intervals based on three biological replicates with three technical replicates each. Data and statistics are reported in Supplementary Table S1. (F) Transcription shut-off with ActD was performed using the Nluc 3× PRE reporter mRNA to measure the effect of RNAi depletion of Pum or Pop2 relative to NTC. A representative Northern blot of Nluc 3× PRE reporter and 18S rRNA is shown. Each lane of the gel contains 10 μg of total RNA.
lase complex, and specific CNOT subunits are required to repress and degrade target mRNAs. We also detect that the Pum N-terminus has an additional repressive activity that circumvents the requirement for CNOT and the poly(A) tail and involves the mRNA decapping enzyme, Dcp2. We measured the contribution of multiple mechanisms to repression by Pum, emphasizing the importance of CNOT, Dcp2 and pAbp. Taken together, our data reveals that Pum utilizes deadenylation and decapping pathways to repress and degrade its target mRNAs.

MATERIALS AND METHODS

Plasmids and cloning

The plasmids used in this study are listed in Supplementary File 1 and were verified by DNA sequencing. The sequences of all oligonucleotide primers are listed in Supplementary File 1. The pIZ plasmid (Invitrogen) was used for effector expression and contains the OpIE2 promoter. Drosophila Kozak sequence, C-terminal V5 epitope and His6 tags, and the SV40 cleavage/poly-adenylation site. For experiments employing over-expression of Pumilio, the coding region of wild type (NP_001262403.1) or RNA-binding defective (mut R7) Pumilio was cloned into pIZ to create pIZ Pumilio V5H6 and pIZ Pumilio mut R7 V5H6 plasmids, as previously described (36). Amino acid residues S1342A, N1343A and E1346A of the seventh repeat of the Pum-HD are mutated in the RNA-binding defective Pum mut R7 (Figure 1A), as previously described (25,36).

For tethered function assays, the MS2 fusion effector-encoding plasmids pIZ MS2-PumN V5H6, pIZ MS2-RD1 V5H6, pIZ MS2-RD2 V5H6 and pIZ MS2-RD3 V5H6 plasmids were previously described (36). The negative control pIZ MS2-EGFP plasmid was created by inserting the coding sequence for enhanced green fluorescent protein (EGFP), amplified by PCR using oligos CW115 and CW116, into KpnI and XhoI sites in the pAc5.1 vector (36). Likewise, the Dcp1 coding sequence (NP_611842.1, amplified with oligos RA 102 and RA 103) was inserted into SpeI and NotI sites to create pIZ MS2-Dcp1.

For RNAi rescue experiments, cDNA clone pIZ myc-Pop2 was generated by insertion of the Pop2 coding sequence (NP_648538.1, amplified using oligo CW 033 and CW 034) with an N-terminal Myc tag into HindIII and XbaI sites of the pIZ plasmid. Pop2 mutations, D52A and E54A, were introduced into pIZ myc-Pop2 using quick-change site-directed mutagenesis (Agilent) with primers CW 161 and CW 162.

The expression plasmid vector pUBKz3 × Flag contains the Drosophila ubiquitin 6E3 promoter, Kozak sequence, N-terminal 3 × Flag, and SV40 cleavage and poly-adenylation site in a pUC19 backbone (provided by Dr Eric Wagner, University of Texas Medical Branch). For inhibition of decapping by over-expression, the plasmid pUBKz3 × Flag Dcp2 E361Q was created by inserting the Dcp2 coding sequence (NP_001246776.1, amplified with oligos RA 086 and RA 087) into SpeI and NotI sites in pUBKz3 × Flag vector followed by site directed mutagenesis to introduce the E361Q mutation (as used in (45,52,53)) using oligos RA 166 and RA 167.

Reporter genes are based on vector pAc5 (Invitrogen). The internal control plasmid, pAc5.1 FFluc min 3′ UTR, which expresses firefly luciferase, was described previously (36). The reporter plasmid pAc5.4 Nluc2 ΔPRE 3′ UTR was cloned by inserting the Nano-luciferase (Nluc) coding sequence with C-terminal PEST sequence, derived from pNL1.2 plasmid (Promega) and amplified using oligos CW 578 and RA 066, into the KpnI and XhoI sites of vector pAc5.4 (25). The tethered function reporter plasmid pAc5.4 Nluc2 × MS2 was cloned by inserting oligos AG 784 and AG 785, encoding two copies of the binding site for MS2 coat protein (MS2), into XhoI and NotI sites of pAc5.4 Nluc MCS. To create the Histone Stem Loop (HSL) reporter, pAc5.4 Nluc2 2 × MS2 HSL, inverse PCR with oligos RA 255 and RA 256 was performed using pAc5.4 Nluc2 2 × MS2 template, thereby replacing cleavage/polyadenylation element with the HSL and Histone Downstream Element (HDE) sequences. To create the Pum reporter plasmid, first a unique XhoI site was inserted into pAc5.1 Rnuc (36) using inverse PCR with oligos RA 214 and RA 215. Next, the Nluc2 coding sequence was inserted into KpnI and XhoI sites in the pAc5.1 vector to create pAc5.1 Nluc2 3×PRE.

For production of recombinant Pum constructs in Escherichia coli, cDNA sequences encoding Pum RD1 (aa 1–378), RD2 (aa 548–776), RD3 (aa 848–1090) or Pum RBD (aa 1091–1426) were inserted using the Gibson assembly method (54) into the pYC-pM plasmid vector (55) lin-
earized with NdeI. The resulting Pum RD1, RD2, RD3 and RBP fusion proteins have N-terminal MBP tags that are cleavable by the human rhinovirus 3C (HRV3C) protease, and a C-terminal StrepII tag.

Cell culture and transfection

D.mel-2 cells (Invitrogen) were cultured in Sf900III media (Thermo-Fisher, see Supplementary File 1 for reagents) at 25°C in 100 μg/ml penicillin and 100 μg/ml streptomycin. In our standard transient transfection procedure, 2 million d.mel-2 cells were plated in a six-well plate and transfected with 150 μl of transfection mix containing FuGene HD (Promega) and 3 μg effector DNA at a 4 μl Fugene HD:1 μg DNA ratio in Sf900III media. This transfection mix was incubated for up to 15 min at room temperature prior to application to cells. In experiments that measure regulation by endogenous Pum, the transfection mix contained a total of 1.5 μg of transfected DNA with FuGene HD. For dual luciferase assays and Northern blotting, 20 ng of pAc5.4 NLuc poly(A) or 100 ng Nluc HSL, along with 20 ng pAc5.1 FFluc, were included in the transfection mix.

For the Pop2 rescue experiments, 5 ng of pAc5.4 NLuc 2× MS2 poly(A) reporter and pAc5.1 FFluc internal control plasmid were included in the transfection mix. In the Pop2 RNAi rescue experiment (Figure 4F–H), cells were transfected with 2.25 μg of the indicated tethered effector and 750 ng of either pIZ EGFP V5, pIZ myc-Pop2 or pIZ myc-Pop2 D52A E54A. For analysis of Not1 RNAi rescue with exogenous Pop2 (Supplementary Figure S3), either 150 ng of pIZ EGFP V5 control, 100 ng myc-Pop2 (with 50 ng pIZ EGFP V5) or 150 ng myc-Pop2 were transfected into cells, along with 2.85 μg of the indicated MS2-tethered effectors.

For analysis of decapping, cells were transfected with 1.5 μg of the indicated MS2-tethered effectors and 1.5 μg of either pUbKz 3× Flag empty vector or pUbKz 3× Flag Dcp2 E361Q plasmid. In this approach, the cells were also treated with either non-targeting control (NTC) or Dcp2 double stranded RNA, as indicated in the figure. Importantly, RNAi of the Dcp2 mRNA targeted the 3' UTR and thus did not affect expression of Dcp2 E361Q. To measure regulation by endogenous Pum and co-repressors in Figure 11, the transfection mix contained a total of 1.5 μg of transfected DNA with FuGene HD, along with 20 ng of FFluc and 20 ng of the indicated Nluc reporter plasmids.

RNA interference

To induce RNAi, gene-specific double stranded RNA (dsRNA), ranging from 133–601 bp, were designed using the SnapDragon web-based tool provided by the Harvard Drosophila RNAi Screening Center (URL: http://www.flyrnai.org/cgi-bin/RNAi_find_primers.pl) to minimize potential off-target regions. Transcription templates were PCR-amplified with primers that add opposing T7 promoters to each DNA strand (see Supplementary File 1 for dsRNA template primer sequences with T7 RNA polymerase promoters). The dsRNAs were transcribed from these templates using HiScribe T7 high yield RNA synthesis kit (New England Biolabs). The dsRNAs were then treated with RQ1 RNase-free DNase (Promega) and purified using RNA Clean & Concentrator-25 (Zymo Research). Non-targeting control (NTC) dsRNA, corresponding to the E. coli LacZ gene, was described previously (36).

For all RNAi experiments measuring dual luciferase activity, d.mel-2 cells were plated in 6-well plates with 24 μg dsRNA per well. In the standard protocol, one million cells were plated with dsRNA, incubated 24 h, then reporters and effectors were transfected with FuGene HD as described above. Forty-eight hours after transfection, cells were harvested for luciferase assays, western blotting, and RNA isolation. For Figures 4F–H, 10 and Supplementary Figure S3, a half-million cells were plated and incubated with dsRNAs for 72 h, then reporters and effectors were transfected with FuGene HD as described above. Forty-eight hours after transfection, cells were harvested for luciferase assay, western blotting and RNA isolation.

Reporter gene assays

D.mel-2 cells were harvested from a transfected 6-well plate and 100 μl of cell culture (~0.5–6 × 10^5 cells, depending on experimental conditions) was aliquoted into a 96-well plate. Luciferase assays were then performed using Nano-Glo Dual-Luciferase Reporter Assay System (Promega) and a GloMax Discover luminometer (Promega) per manufacturer's instructions, using 10 μl/ml of Nluc substrate. To measure regulation by endogenous or over-expressed Pum, the Nluc 3× PRE reporter was used. To measure activity of tethered effectors, the Nluc 2× MS2 poly(A) or Nluc 2× MS2 HSL reporters were used. The internal control pAC5.1 FFluc 3′UTR poly(A) was used in all reporter experiments. In tethered assays, MS2-EGFP served as the negative control for effectors. In experiments analyzing repression by full-length wild type or mutant W783G Pum, the RNA-binding defective mutant Pum (mut R7) served as a negative control, as previously established (36).

The Nluc and FFluc reporter activities from each sample, measured in Relative Light Units (RLU), were used to calculate fold change values, as described below. First, Relative Response Ratios (RRR) for each sample were calculated by dividing the Nluc value by the FFluc value to normalize variation in transfection efficiency. Next, the log2 fold change in RRR for a given effector/condition was calculated relative to a negative control effector/condition. For tethered function assays, log-scaled fold change by an effector was determined relative to the mean RRR for the negative control effector, MS2-EGFP, unless otherwise noted. For tethered function assays utilizing RNAi of a putative co-repressor, fold change induced by an effector was measured relative to the negative control effector, MS2-EGFP, within the same RNAi condition.

For Pum-mediated repression of the PRE-containing reporter, Nluc 3× PRE, the log2 fold change induced by wild type or mutant Pum was determined relative to the mean RRR for the RNA-binding defective Pum mut R7 negative control. For RNAi experiments in Figure 11C, log2 fold change induced by depletion of a regulatory factor was determined relative to the mean RRR for the non-targeting control siRNA. The data was analyzed by determining the log2 RRR of Pum-repressed Nluc 3× PRE reporter activ-
ity, relative to the log$_2$ RRR of the non-Pum regulated Nluc ΔPRE reporter activity within the same RNAi condition. In doing so, the Pum specific effect is measured while normalizing for Pum/ΔPRE independent effects of the RNAi. From this data, the fold change in reporter expression induced by each RNAi condition was calculated relative to the NTC. The data were separately analyzed and reported in Supplementary Figure S4 in the same manner, except that the FFluc values were omitted from the calculations. The PRE-dependent effect of each RNAi condition on the Nluc 3× PRE reporter was normalized to the effect on the unregulated Nluc ΔPRE reporter. The log$_2$ fold change in PRE-mediated regulation within each RNAi condition was then calculated relative to the negative control NTC dsRNA. Further details of statistical analysis are described below.

**Pum antibody**

The anti-Pumilio rabbit polyclonal antibody was generated using the recombinant purified antigen containing Pum residues 1434–1533 fused to GST. Pum-specific antibodies were antigen-affinity purified from the resulting serum using a column containing immobilized, recombinant, purified Halotag-Pum aa1434–1533 immobilized to Halolink resin (Promega) (56). The Pum antigen sequence was: PIITVGTGAGGVPAAASSAAVSSGATSASVTACTSGS STTTTSTTNLSAPTICSVQENGSMVVEPSSPDAS ESSSSVSGAVNSSLPGLPVPTNGNVL.

**Western blotting**

Cell lysates were prepared by adding 100 μl of lysis buffer (50 mM Tris–HCl pH 8.0, 150 mM NaCl, 2× complete mini, ethylenediaminetetraacetic acid (EDTA)-free protease inhibitor cocktail (Roche), and 0.5% Non-Idet P40 (NP40)) to cell pellets containing 0.5–6 × 10$^6$ cells, dependent on experimental design. Cells were lysed for 10 seconds using a cell disruptor. Cell debris was removed by centrifugation at 21 100 × g for 10 minutes. Total protein in the resulting cell lysate was quantitated using Lowry DC assay (BioRad) with a bovine serum albumin (BSA) standard curve. Equal mass (10 μg, unless noted otherwise) of cell lysates were then analyzed on SDS-PAGE gels (4–20% Mini-PROTEAN TGX or Criterion TGX, BioRad) along with PageRuler Prestained Plus Molecular Weight Markers (Thermo Fisher). For detection of endogenous Pum protein, 30 μg of total cellular protein per lane was analyzed. Protein was then transferred onto Millipore Immobilon-P (or Immobilon-PSQ for detection of Pop2 and Ccr4) western blot membranes (50% slurry) of Strep-Tactin sepharose resin (IBA). After 1 h incubation, the beads were washed three times with lysis buffer and once with binding buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl, 0.3% (v/v) Tween-20, pH 7.4). The cleared lysates were incubated with 30 μl (50% slurry) of Strep-Tactin sepharose resin (IBA). After 1 h incubation, the beads were washed three times with lysis buffer and once with binding buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl). Purified human CCR4–NOT complex (50 μg) was then added to the beads. The reconstitution of the human CCR4–NOT complex (50 μg) was then added to the beads. The reconstitution of the human CCR4–NOT complex was described elsewhere (57) and includes the following eight components: CNOT1 (amino acids 1–2376), CNOT2 (1–540), CNOT3 (1–753), CNOT10 (1–299). After 1 h incubation, the beads were washed three times with lysis buffer and once with binding buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl). Purified human CCR4–NOT complex (50 μg) was then added to the beads. The reconstitution of the human CCR4–NOT complex was described elsewhere (57) and includes the following eight components: CNOT1 (amino acids 1–2376), CNOT2 (1–540), CNOT3 (1–753), CNOT10 (25–707), CNOT11 (257–498), CAF1 (1–285), CCR4a (1–558), CAF40 (1–299). After 1 h incubation, the beads were washed three times with binding buffer and the proteins were eluted with binding buffer supplemented with 2.5 mM D-dephtiothiotobiotin. The eluted proteins were analyzed by SDS-PAGE and western blot.

**In vitro pulldown assays**

The StrepII-tagged MBP and MBP-tagged Pum fragments (RD1, RD2, RD3 and RBD) were expressed in E. coli BL21 (DE3) Star cells (Thermo Fisher Scientific) grown in LB medium overnight at 37°C. Cells were lysed by a cell disruptor in Flag Buffer A (50 mM Tris–HCl pH 8.1, 200 mM NaCl, 1 mM EDTA, 0.2% Triton X100 and 2× complete protease inhibitor cocktail (Roche) and cell debris was removed by centrifugation at 21 100 × g for 10 minutes. Cell extracts were split into RNase treated (4 units RNaseOne, Promega) and untreated (120 units of RNasin ribonuclease inhibitor, Promega) samples and incubated with 10 μl bed volume of EZview Red anti-FLAG M2 Affinity Gel (Sigma) overnight at 4°C. Beads were washed three times with Flag Buffer A, and three times with Flag Buffer B (50 mM Tris–HCl pH 8.1, 200 mM NaCl and 1 mM EDTA) and then resuspended in 60 μl Flag Buffer B. Bound proteins were eluted by heating in 1× SDS-PAGE loading dye. Samples were then analyzed by SDS-PAGE and western blot.

**Immunoprecipitation**

D.mel-2 cells (2 million per sample) were transfected with the Flag-tagged bait protein and V5-tagged prey protein expression plasmids indicated in Figure 7 using FuGene HD as described above and incubated for 3 days at 25°C to allow protein expression. Cells were lysed with a cell disruptor in Flag Buffer A (50 mM Tris–HCl pH 8.1, 200 mM NaCl, 1 mM EDTA, 0.2% Triton X100 and 2× complete protease inhibitor cocktail (Roche) and cell debris was removed by centrifugation at 21 100 × g for 10 minutes. Cell extracts were split into RNase treated (4 units RNaseOne, Promega) and untreated (120 units of RNasin ribonuclease inhibitor, Promega) samples and incubated with 10 μl bed volume of EZview Red anti-FLAG M2 Affinity Gel (Sigma) overnight at 4°C. Beads were washed three times with Flag Buffer A, and three times with Flag Buffer B (50 mM Tris–HCl pH 8.1, 200 mM NaCl and 1 mM EDTA) and then resuspended in 60 μl Flag Buffer B. Bound proteins were eluted by heating in 1× SDS-PAGE loading dye. Samples were then analyzed by SDS-PAGE and western blot.

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The StrepII-tagged MBP and MBP-tagged Pum fragments (RD1, RD2, RD3 and RBD) were expressed in E. coli BL21 (DE3) Star cells (Thermo Fisher Scientific) grown in LB medium overnight at 37°C. Cells were lysed by a cell disruptor in Flag Buffer A (50 mM Tris–HCl pH 8.1, 200 mM NaCl and 1 mM EDTA). The StrepII-tagged MBP and MBP-tagged Pum fragments were purified from E. coli BL21 (DE3) Star cells (Thermo Fisher Scientific) grown in LB medium overnight at 37°C. Cells were lysed by a cell disruptor in Flag Buffer A (50 mM Tris–HCl pH 8.1, 200 mM NaCl and 1 mM EDTA) and then resuspended in 60 μl Flag Buffer B. Bound proteins were eluted by heating in 1× SDS-PAGE loading dye. Samples were then analyzed by SDS-PAGE and western blot.
Radioactive antisense RNA or DNA probes were used for Northern blot detection. Transcription templates for Nluc and FFLuc antisense RNA probes were PCR-amplified using DNA oligonucleotides with a T7 RNA polymerase promoter appended to the antisense strand, described in Supplementary File 1. Using these templates, in vitro transcription was performed for 10 min at 37°C with the T7 MAXIscript transcription kit (Thermo-Fisher) in the supplied 1× Transcription Buffer with 1 µg of DNA template, 0.4 mM final concentration of ATP, CTP, GTP and 8 µM UTP, 2 µl of 800 Ci/mmol 10 µCi/ml 12.5 µM UTP α-32P (1 µM, 10–20 µCi final) (PerkinElmer), and 30 units T7 RNA polymerase in a 25 µl reaction. Next, 1 µl Turbo DNase (2 U) (Thermo-Fisher) was added to the reaction for 10 min at 37°C and then 1 µl of 250 mM of EDTA and 250 mM EGTA was added to the reaction. The probes were purified using a G25 sephadex (GE Life Sciences) spin-column. To detect 18S rRNA, 1.7 µg of 18S rRNA deoxy-oligonucleotide antisense probe (see Supplementary File 1) was phosphorylated using 2 µl of 6000 Ci/mmol, 150 mCi/ml, 25 µM ATP γ-S32P (2.5 µM final, 25 to 100 µCi) (PerkinElmer) and 40 units of T4 Polynucleotide Kinase (New England Biolabs) in a 20 µl reaction incubated at 37°C for 40 min. The probe was then purified with G25 Sephadex column.

For anti-sense Nluc and FFLuc probes, 2.5–7.5 × 106 total cpm was added to the blot that had been pre-hybridized for 45 min at 68°C in 8 ml of ULTRAhyb hybridization buffer (Invitrogen). The blot was then incubated with probe at 68°C overnight, washed two times sequentially with 2 ml each of 2× SSC, 0.1% SDS, and then two more times with 0.1× SSC, 0.1% SDS at 68°C for 15 min each wash. For 18S rRNA probes, 5–6 × 106 cpm was added to the blot that had been pre-hybridized with 8 ml ULTRAhyb-Oligo hybridization buffer (Invitrogen) at 42°C. The blot was incubated with probe overnight and then washed twice with 25 ml 2× SSC containing 0.5% SDS for 30 min each wash at 42°C. Blots were then exposed to phosphor screens and visualized using a Typhoon FLA phosphorimager (GE Life Sciences) and analyzed using ImageQuant TL software (GE Life Sciences). Background signal was subtracted using the ‘Rolling Ball’ method in ImageQuant.

Nluc and FFLuc levels were measured in phosphorimager units (PIU). For analysis of steady state reporter mRNA levels, fold change was determined in the same manner as described for the reporter activity measurements, first normalizing Nluc signal to the corresponding FFLuc signal in that sample, and then calculating the log2 fold change relative to the negative control effector/condition. In tethered function assays, MS2-EGFP served as the negative control for normalization of the effectors. In the full-length Pum experiment, the RNA-binding defective mutant, Pum mut R7, served as the negative control effector.

For experiments measuring effect of endogenous Pum and corepressors on reporter mRNA levels, Northern blot data was analyzed in two ways. First, the values of the Pum regulated Nluc 3× PRE were divided by the Nluc ΔPRE reporter to normalize Pum specific activity to global effects on gene expression. From this data, the log2 fold change was calculated relative to non-targeting control (NTC) negative control RNAs. In the second approach, the RRR of
Nluc 3 × PRE reporter mRNA was normalized to the RRR of Nluc ΔPRE reporter mRNA within each RNAi condition. From these RRR values, the log₂ fold change in Nluc 3 × PRE mRNA was calculated relative to NTC.

To measure RNA decay rates, Nluc signal was normalized to stable 18S rRNA signal for each sample to adjust for potential variation in loading and transfer of RNA in each lane over the time courses. The fraction of reporter mRNA remaining at each time point was plotted relative to time in minutes after Actinomycin D addition. Half-lives and statistical parameters were calculated as described below. Mean mRNA half-lives and 95% credible intervals are reported for each experiment.

High resolution Northern blotting was performed to analyze Nluc 2 × MS2 pA and HSL reporter mRNAs. First, 3 μg of total RNA was heated at 70°C with 20 pmol of antisense Nluc cleavage oligo RA 296 (see Supplementary Table S1) in a 30 μl reaction containing 200 mM KCl and 1 mM EDTA. In control reactions that remove the poly(A) tail (the A₀ control) 1.5 μg of oligo deoxothyymidine (dT) was included. Reactions were then cooled at room temperature for 20 min. Next, 5 units of RNase H (New England Biolabs) in 20 mM Tris–HCl pH 8.0, 28 mM MgCl₂ and 48 units of RNasin (Promega) were added and reactions were incubated at 37°C for 1 h. Next, EDTA (final 30 mM) was added and reactions were incubated for 15 min at 37°C. The RNA was then purified with Clean and Concentrator–25 kit (Zymo). Next, 1.2 μg of purified RNA was combined in equal volume (15 μl) of RNA loading buffer (88% formamide, 0.025% Bromophenol blue, 0.025% xylene cyanol, 10 mM EDTA and 0.025% SDS), heated for 10 min at 75°C. Samples were then electrophoretically separated on a 5% poly-acrylamide, 1 × Tris–borate–EDTA (TBE), 8 M urea gel (BioRad) that had been pre-run at 20–25 mA, 200 V, in 1 × TBE buffer (89 mM Tris, 89 mM Boric acid and 2 mM EDTA). Next, the RNA was transferred onto Immobilon-Ny+ Membrane (Millipore) for 45 min in 0.5 × TBE buffer at 60 V at 4°C using a Trans-Blot Cell (BioRad). The blot was crosslinked with 120 J/cm² UV (λ = 254) and probed with a radioactive, antisense 2 × MS2 RNA probe (see Supplementary File 1 for primers).

Reverse transcription and quantitative polymerase chain reaction

All parameters for RT-qPCR, including primer sequences, amplification efficiencies, and amplicon sizes, are reported according to MIQE guidelines (59) in Supplementary File 2. Data and statistics for RT-qPCR are reported in Supplementary Table S1. Reverse transcription was performed using GoScript (Promega) following the manufacturer’s instructions. Purified RNA (1.6–4 μg) and 500 ng of random hexamers were combined and heated in 10 μl volume of RNase free water at 70°C for 5 min, followed by cooling on ice for 5 min. Next, GoScript Buffer (1 × final), dNTPs (0.5 mM final), MgCl₂ (2 mM final) and 20 units of RNase inhibitor, and 160 units GoScript reverse transcriptase were combined in a 20 μl reaction that was then incubated at room temperature for 5 min, 42°C for 45 min and 70°C for 15 min. As a negative control for each primer set, mock ‘no RT’ reactions were performed using identical conditions except that the reverse transcriptase was omitted. Next, qPCR was performed using GoTaq qPCR Master Mix (Promega) with the equivalent of either 50 or 100 ng (specified in Supplementary File 2) of input RNA and 0.1 μM of each primer per reaction in 50 μl final volume. In addition, no template reactions were also performed wherein cDNA was omitted, so as to assess potential false positive signal for each primer set.

The following cycling parameters were performed using a CFX96 Real-Time PCR System thermocycler (BioRad): (i) 95°C for 3 min, (ii) 95°C for 10 s, (iii) 65°C (for Not1, Not2, Not3, Caf40 Dep1 and Dep2 reactions) or 62°C (for Pop2, Ccr4, Not10 and Not11 reactions) for 30 s, (iv) 72°C for 40 s, (v) repeat steps 2–4 for 40 cycles. Melt curve was generated with range 65–95°C at increments of 0.5°C. Data and statistical analysis are described below.

Data and statistical analyses

To assess replication and reproducibility of measurements, we employed several types of replicates including experimental replicates (i.e. independent assays performed on separate days using the same approach), biological replicates (i.e. parallel measurements of distinct cell samples) and technical replicates (i.e. multiple measurements from the same sample). The number and type of replicates were dictated by the assay type, experimental design, and feasibility. All data, number and types of replicates, and statistics (e.g. credible intervals and posterior probabilities) are reported in Supplementary Table S1.

For each data set of interest, we fitted a hierarchical Bayesian model to account for the differences in variation between technical, biological, and experimental replicates and draw robust inferences on the biological parameters of interest. All models were fitted using STAN via the brms package (60–62), and assessed for convergence using both the Gelman–Rubin shrinkage statistic (63) and visual inspection of the posterior predictive distributions (as well as all built-in diagnostics in STAN/brms). As a general rule, we fitted data using at least four independent Monte Carlo chains for at least 2000 iterations, with the fits expanded when needed to ensure convergence. Unless otherwise noted, all models fitted were linear, using a Student’s t family error distribution, with group-level effects for all batch variables that could be identified (e.g. date of assay, plate identity, etc.), and population-level effects for the biological variables of interest (e.g. genotype and RNAi status), as well as relevant interaction terms between those biological variables. We used default priors from brms unless otherwise noted. The response variable for each fit was globally centered prior to fitting, as the differences between parameters were the topic of interest.

For the majority of our assays (exceptions are noted below), the key parameter of interest was the log₂ ratio of Nluc to FFLuc detected, either at the mRNA or protein level. The reported values are given as the difference between the fitted values for each combination of biological variables of interest, and the corresponding values for a negative control (as described above and in the Results and figure legends). The control samples for each experiment appear with a mean of 0, as they are similarly subjected to subtraction of their
posterior mean. Posterior means are shown as the plotted values, with error bars giving the 95% credible intervals. In addition, we flag results based on three types of significance criteria: * indicates a significant difference for which the 95% credible interval excludes zero (thus, we are reasonably certain that there is some nonzero effect in the indicated direction). ** indicates a difference for which there is a 95% posterior probability of a difference of at least 1.3-fold in the indicated direction (thus, we are reasonably certain that there is a substantial difference in the indicated direction). Finally, ‘X’ indicates a difference for which 95% of the posterior density indicates a less than 1.3-fold change in either direction (thus, we are reasonably certain that there is no substantial difference in either direction). We used similar approaches to analyze quantitative PCR data, except that in this case the observables were the amplification efficiency-weighted \( C_q \) values (calculated as in Equation 3 of Ganger et al. (64)). Our inferences are subsequently drawn from differences in efficiency-weighted \( C_q \) values of the transcripts of interest and corresponding reference transcript, Rpl32. Efficiencies of qPCR assays are reported in Supplementary File 2.

In the case of the mRNA decay data shown in Figures 1 and 6, we instead fitted exponential decay equations for the fraction of material of interest remaining at a given time \( t \), \( f(t) \), of the form:

\[
f(t) = (1 - \eta) + \eta \exp(-\lambda t) + \varepsilon
\]

With \((1 - \eta)\) representing a non-reacting fraction of the population (or one with a far longer half-life than the majority), \(\lambda\) representing a decay constant (fitted on a logarithmic scale) and \(\varepsilon\) representing Student’s \( t \) distributed errors. Both \(\eta\) and \(\lambda\) are taken to be a sum of population-level effects from the biological variables of interest and the specific biological effect being measured. Nonstandard priors were used in both cases; the \(\eta\) contributions from biological variables had a beta \((0.5,0.5)\) prior with an upper bound of 1 and lower bound of 0.5, and the replicate contributions had a normal \((0,0.1)\) prior. The biological contribution to log(\(\lambda\)) had a normal \((0,0.1)\) prior, and the replicate-level contribution had a normal \((0,0.1)\) prior. Other priors (e.g. for the Student’s \( t \) error distribution) used BRMS defaults. We then report numeric values for the effective half life (that is, the time required for only 50% of the initial signal to remain, assuming that decay follows the equation above), and plot the decay curves that would arise from the posterior mean values for the biological parameters (neglecting the replicate-level parameters). For the cases shown in Figure 6, we found that one biological replicate for WT cells (in the Not1 RNAi case) and one for Pum mutant cells (in the Pop2 RNAi case) were profound outliers; those two biological replicates were excluded from contributing to the overall analysis.

RESULTS

Pumilio accelerates mRNA degradation

To investigate the mechanism of Pum-mediated repression, we utilized luciferase reporter gene assays in the Drosophila cell line, d.mel-2. We previously demonstrated that d.mel-2 cells express a limiting amount of endogenous Pum, and RNAi-mediated depletion of Pum specifically increased protein expression from a luciferase reporter mRNA bearing three PRE sequences in a minimal 3′ UTR (25,36,44).

We also reported that over-expression of full-length Pum protein (Figure 1A) further repressed expression of the PRE-containing reporter in a dosage dependent manner. The effect is highly specific, as mutation of the PRE sequence or the RNA recognition amino acids of the 7th repeat in the Pum RBD (Figure 1A, mut R7) alleviated RNA-binding and repression (25,36,44).

Here we report that endogenous and over-expressed Pum specifically accelerate degradation of a PRE-containing Nano-luciferase reporter mRNA. First, we analyzed the decay rate of reporters with or without PRE sequences (Figure 1B, Nluc 3× PRE versus Nluc ΔPRE) using a transcription shut-off approach and Northern blot detection (58). The PREs substantially decreased the Nluc 3×PRE mRNA level and significantly reduced its half-life by 8.3-fold relative to Nluc ΔPRE, with a \( P(\text{sig}) > 0.999 \) (Figure 1C and D), indicating that Pum recognition of the mRNA caused its degradation. We note that the statistical analyses in this report utilized a Bayesian approach (described in Materials and Methods). Data reported in figures include mean values and 95% credible intervals, with posterior probabilities and number and type of replicates reported in the figure legends and Supplementary Table S1. Following previous work on Pum proteins (65), we assess differences based on 95% credible intervals and \( P(\text{sig}) \), the posterior probability of a difference of at least 1.3-fold in the direction indicated. High values of \( P(\text{sig}) \) indicate high confidence in a biologically meaningful effect being present. Values of \( P(\text{sig}) \geq 0.95 \) are indicated by a ** in the figures. Differences that are significant (posterior probability of \( \geq 0.95 \) that a difference in the indicated direction exists) but do not meet this stringent 1.3-fold threshold are marked with a * in the figures. We also use a converse measure, \( P(\text{insig}) \), which we define as the posterior probability that a change is no larger than 1.3-fold ((\( P(\text{insig}) \geq 0.95 \) are indicated by a ‘X’ in the figures). We note that a large value of \( P(\text{insig}) \) is far more informative than a large \( P \)-value would be in a frequentist statistical test; whereas the latter only represents a failure to reject the null hypothesis, and thus has limited inferential value, a large value of \( P(\text{insig}) \) represents true confidence that an effect is small.

We next determined that PRE-mediated mRNA decay is dependent on Pum. RNAi depletion of Pum, verified by RT-qPCR in Figure 1E, increased the half-life of the Nluc 3× PRE mRNA by 7.2-fold (\( P(\text{sig}) > 0.999 \)) relative to non-targeting control RNAi (Figure 1F and G). Because the Pop2 deadenylase subunit of the CNOT complex is implicated in Pum-mediated repression, we tested the effect of Pop2 depletion (itself, confirmed in Figure 1E), and observed a 3.9-fold increase in Nluc 3× PRE mRNA half-life (\( P(\text{sig}) = 0.99 \)) (Figure 1F and G). This result indicates that Pum-PRE-mediated mRNA decay occurs through the deadenylation-mediated pathway.

Next, we measured the effect of over-expressed Pum on mRNA decay. We previously showed that Pum expression specifically repressed PRE-containing reporter mRNAs, whereas an RNA-binding defective mutant, Pum mut R7, did not (25,36,44). Consistent with those observations,
expression of wild type Pum, but not mut R7, repressed protein expression from the Nluc 3× PRE mRNA by >4-fold (P(sig) > 0.999) in a dual luciferase reporter assay (Figure 1H and I) and accelerated mRNA decay (Figure 1J and K), reducing the Nluc 3× PRE mRNA half-life by 2.8-fold (P(sig) > 0.999). Expression of the V5 epitope-tagged Pum effector proteins was confirmed by western blot of equal mass of cell extracts (Figure 1I). Together, these results indicate that Pum-PRE mediated repression accelerates mRNA degradation and that Pop2 deadenylation is important for decay of PRE-bearing mRNAs.

N-terminal Pumilio repression domains cause mRNA decay

We previously showed that the N-terminus of Pum confers its major repressive activity, mediated by three Repression Domains (Figure 1A, RD1, RD2 and RD3) (36). These domains can repress in a tethered function assay, wherein they are fused to the RNA-binding domain of the MS2 phage coat protein and directed to the 3′ UTR of a reporter mRNA bearing tandem MS2 binding sites, Nluc 2×MS2 (Figure 2A) (36,66–68). These assays include a co-transfected control Firefly luciferase gene (FFLuc) for normalization of transfection efficiency in each sample. In agreement with our previous observations (36,44), tethering the Pum N-terminus or each RD reduced reporter protein expression (Figure 2B, Nluc 2×MS2), whereas the negative control MS2-EGFP fusion did not. Under these conditions, Pum N and RD1 repressed by 2-fold relative to MS2-EGFP, while RD2 and RD3 repressed by 1.4-fold and 1.5-fold, respectively (all Pum effectors had P(sig) > 0.999). As a positive control, the decapping enzyme subunit Dcp1, fused to MS2, elicited 3.9-fold repression (P(sig) > 0.999) (Figure 2B), consistent with the previously reported effect of tethering decapping factors (53,69–73). Importantly, the observed repression was dependent on binding of effector proteins to the reporter, because they had little or no effect on a reporter that lacks the MS2 binding sites, Nluc ΔMS2 (each Pum effector had P(insig) > 0.999) (Figure 2B). We also compared the repressive activity of each effector between the two reporters, demonstrating that Pum N repressed by 3.3-fold (P(sig) > 0.999), RD1 by 3.1-fold (P(sig) > 0.999), RD2 by 2-fold (P(sig) = 0.99), RD3 by 2.5-fold (P(sig) > 0.999), and the Dcp1 control repressed by 6.5-fold (P(sig) > 0.999) (Supplementary Figure S1A). The expression of the V5 epitope-tagged effector proteins was confirmed by western blotting of equal mass of cellular lysates (Figure 2C). These results support the independent repressive activity of each Pum RD.

To further characterize the activities of the N-terminal repression domains, we analyzed the relationship of effector protein dosage to repression of Nluc 2×MS2 reporter activity. To do so, the mass of transfected effector expression plasmid (MS2-Pum N, MS2-RD1, MS2-RD2, MS2-RD3 or MS2-EGFP) was titrated over a 30-fold range and repression was measured relative to cells transfected with empty expression vector, pIZ (Figure 2D and E). To maintain identical transfection conditions, the total mass of transfected DNA was balanced across samples with empty expression vector. Pum N and RDs exhibited repression activities proportional to the mass of expression plasmid, and substantially above the equivalent amount of negative control MS2-EGFP (Figure 2D).

We then examined the relationship of effector protein level to repression by performing quantitative western blotting on equal mass of cell extract from these samples (Figure 2E). We observed a log-linear relationship wherein increased Pum effector protein level caused a proportional decrease in reporter expression (Supplementary Figure S1B). In contrast, MS2-EGFP was far less effective. The only substantial deviation from a log-linear relationship of effector amount to repressive activity occurs in the case of MS2-RD1, for which higher effector protein levels appear to be somewhat disproportionately more effective (Supplementary Figure S1B). Importantly, repression activity by each Pum effector was observed to be consistent and proportional across a broad range of plasmid or protein levels; even a 30-fold reduction in effector did not eliminate repression by Pum effectors (Figure 2D and Supplementary Figure S1B).

To directly compare repression activities, the fold change relative to empty vector was determined for equivalent amounts of expressed effector proteins. This analysis shows that per unit of effector protein, the order of efficacy is: Pum N > RD3 = RD1 = RD2 > EGFP (Figure 2F). The posterior probability of the difference of Pum N is greater than the RDs is P(sig) > 0.999 and the activities among the three RDs are not distinguishable from each other (P(insig) ≥ 0.93), whereas their activities are consistently greater than the EGFP negative control (P(sig) ≥ 0.97) (Supplementary Table S1). We conclude that the tethered function assay provides a robust and specific means of assessing Pum RD activity. Minor fluctuations in effector level do not result in loss of activity, nor do they alter our qualitative conclusions regarding the effects of the various constructs.

To measure the impact of each effector on Nluc 2×MS2 mRNA, we performed Northern blot analysis. The Pum N-terminus and RDs reduced the Nluc reporter mRNA level with magnitudes corresponding to their effects on reporter protein level (P(sig) ≥ 0.97) (Figure 2G–I), whereas the internal control Firefly luciferase mRNA was not affected by the tethered effectors (Figure 2I). Northern blot of the 18S ribosomal RNA served as an internal control for gel loading and blotting. These results indicate that the N-terminal Pum RDs promote mRNA decay.

The putative Pumilio cap-binding motif is not required for repression

We interrogated a model wherein Pum was hypothesized to repress translation via a 5′ cap-binding motif located in its N-terminus (11,48). This motif does not correspond to the three RDs, but instead resides within a conserved region (previously designated PCMb) (36). First, we introduced a mutation in the Pum N-terminus, W783G (Figure 1A), which was reported to disrupt cap binding (48) and compared its repressive activity (Figure 3A) and expression (Figure 3B) to wild type in the tethered function assay. By direct comparison of the repressive activities of wild type Pum N to the mutant W783G, we see that there is no significant difference in their repressive activi-
Figure 2. The Pumilio N-terminal Repression Domains repress mRNA and protein expression. (A) Diagram of tethered function nano-luciferase reporter mRNA (Nluc 2×MS2) and co-transfected control, figure Firefly luciferase (FFluc). Nluc 2×MS2 bears two copies of the MS2 stem loop RNA structure.
ties (P(sign) > 0.999), for each transfected amount. We also examined the effect of the W783G mutation on repression of a PRE-containing reporter by over-expressed full length Pum (Figure 3C and D); no significant reduction in the ability of the mutant Pum to repress was observed. Based on this data, and our previous observation that PCMB was neither necessary nor sufficient for repression (36), we conclude that, at least in these experimental conditions, the proposed 5' cap binding motif does not contribute to Pum-mediated repression.

CNOT complex components are important for Pumilio repression domain activity

We sought to identify co-repressors necessary for repression by the Pum RDs. The CNOT deadenylase complex plays a crucial role in the initiation of mRNA decay (39) and is important for mRNA decay by Pum-PRE (Figure 1F and G), thus we evaluated its role in repression by Pum N-terminus and individual RDs. The Drosophila CNOT complex contains eight subunits and the Pop2 subunit is thought to be the major deadenylase (Figure 4A) (40); therefore, we first performed RNAi using two different double stranded RNAs (dsRNAs) to deplete Pop2. DsRNA1 targets the open reading frame of Pop2 mRNA, whereas the dsRNA2 targets its 5'UTR. Both dsRNAs depleted Pop2 from the d.mel-2 cells relative to non-targeting control dsRNA (NTC), as confirmed by RT-qPCR (Figure 4B) and western blotting (Figure 4C). Pop2 depletion was effective and reproducible in each condition using either dsRNA, though depletion by dsRNA1 was stronger (Figure 4B).

We then measured the effect of Pop2 depletion on repression by tethered Pum N-terminus and RDs. It is important to note that in all reporter gene assays that incorporate RNAi, the repressive activity of the effector was measured relative to the negative control effector, MS2-EGFP, within the same RNAi condition, as described in the Methods. In this manner, the specific effect of RNAi-mediated depletion on the Pum effector is determined. We observed that Pop2 depletion substantially reduced but did not eliminate repression by the Pum N-terminus, significantly decreasing repression activity by 1.3- to 1.5-fold relative to the NTC condition (Figure 4D). Pop2 depletion eliminated repressive activity of RD2 and RD3, and greatly reduced repression by RD1. As anticipated based on the ability of Dcp1 to interact with the mRNA decapping enzyme Dcp2, Pop2 depletion did not alleviate repression by tethered decapping enzyme subunit Dcp1, which maintained repressive activity in all conditions (P(sign) > 0.95 in all cases; Figure 4D). Effector expression was verified in each condition (Figure 4E). Based on this data, we conclude that Pop2 is important for repressive activity of the Pum N-terminus and RDs.

To investigate whether the deadenylase activity of Pop2 is involved in Pum RD mediated repression, we tested the ability of wild type or mutant Pop2 to rescue repression. To do so, cells were treated with Pop2 dsRNA2 and then were transfected with plasmid expressing RNAi resistant cDNAs encoding myc-tagged wild type or catalytically inactive Pop2 mutant (D53A and E55A) (40), or V5-tagged EGFP as a control. Depletion of Pop2 mRNA was verified by RT-qPCR (Figure 4F). As before, depletion of Pop2 reduced repression by the Pum effectors in the control EGFP expressing conditions (Figure 4F), compare Pop2 dsRNA2

in its 3'UTR, which is bound by the sequence-specific MS2 RNA-binding protein. By expressing Pum or other effectors as a fusion to MS2 RNA-binding protein, the impact of the effector on reporter protein and mRNA levels can be measured. Diagram is not to scale. (B) The repressive activity of Pum N-terminus and individual repression domains (RD1, RD2, RD3) was measured using the tethered fusion dual luciferase assay using the Nluc × MS2 reporter or an equivalent reporter wherein the MS2 binding sites are deleted, Nluc ΔMS2. Mean log2 fold change values in normalized reporter activity, measured relative to the negative control, MS2-EGFP, are plotted with 95% credible intervals, from three experiments with four technical replicates each. Tethered decapping enzyme subunit, Dcp1, serves as a positive control that strongly represses the reporter when tethered. Data and statistics are reported in Supplementary Table S1. Comparison of the activity of each Pum effector on the Nluc × MS2 reporter relative to Nluc ΔMS2 is shown in Supplementary Figure S1A. For significance calling, a * denotes a posterior probability > 0.95 that the difference relative to the negative control is in the indicated direction. The ** indicates a posterior probability of > 0.95 that the indicated difference is no more than 1.3-fold in either direction. An ‘x’ marks a posterior probability > 0.95 that the indicated difference is greater than 1.3-fold in one direction. (C) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in panel B from a representative experimental replicate. Equivalent mass of protein from each sample was probed with anti-V5 antibody, followed by the Integrator Subunit 1 protein (IntS1) to assess equal loading of lanes. (D) The relationship of repression activity to level of each MS2 tethered effector was measured using the tethered function assay by titrating the amount of transfected effector expression plasmid, as indicated at the top. Total mass of transfected plasmid was maintained across all conditions by supplementing with empty expression plasmid vector, pIZ. In this experiment, repression of reporter activity was calculated relative to the control condition containing only the empty expression vector. Data from three experiments with four technical replicates each, are plotted along with 95% credible intervals. The yellow dashed line marks the repression activity of the negative control effector, MS2-EGFP, for each amount of transfected effector plasmid (relative to the vector-only control). The relationship of effector protein level, measured by quantitative western blot, repression activity is reported in Supplementary Figure S1B. Data and statistics are reported in Supplementary Table S1. (E) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in panel D from a representative experimental replicate. Equivalent mass of protein from each sample was probed with anti-V5 antibody, followed by IntS1 as a loading control. (F) The repression activity for equivalent expression level of each MS2 tethered effector protein, determined by tethered function assays and quantitative western blotting as shown in Supplementary Figure S1B. Fold change was calculated relative to empty vector. Mean log2-fold change values are plotted with 95% credible intervals from three experiments with four technical replicates each. The yellow dashed line marks the repression activity of the negative control effector, MS2-EGFP. Data and statistics are reported in Supplementary Table S1. (G) The effect of each MS2 fusion effector protein on Nluc × MS2 reporter protein and mRNA level was determined by dual luciferase assay (reporter protein, dark gray bars) and Northern blotting (reporter mRNA, light gray bars). Log2 fold change of Nluc × MS2 levels, normalized to internal control FFluc, for each effector were calculated relative to negative control MS2-EGFP for three biological replicates. Mean log2 fold change and 95% credible intervals are reported in the graph. Data and statistics are reported in Supplementary Table S1. (H) Expression of V5-tagged MS2 fusion effector proteins from three biological replicates (panel G) was confirmed by western blotting of an equal mass of protein cell extract for each sample. (I) Northern blot detection of tethered function reporter Nluc × MS2, internal control FFluc, and loading control 18S rRNA for three biological replicate samples for each tethered effector protein. Each lane of the gel contains 5 μg of total RNA. This data was used to determine fold change in reporter mRNA level shown in panel G. The Mock sample contained total cellular RNA from untransfected cells and demonstrates specificity of the reporter probes.
Figure 3. The putative Pumilio cap-binding motif is not required for repression. (A) Repression activity was measured for three amounts of transiently transfected wild type or cap-binding mutant (W873G) Pum N-terminus via the tethered function dual luciferase assay using the Nluc 2× MS2 pA reporter. Repression activity was calculated relative to the MS2-EGFP negative control at the lowest transfected amount (100 ng). Empty expression vector, pIZ, was used to balance the total mass of transfected plasmids in samples with 100 and 500 ng of MS2 effector plasmid. Mean log2 fold change and 95% credible intervals for three experimental replicates with four technical replicates each are reported in the graph. Data and statistics are reported in Supplementary Table S1. For significance calling, a ‘*’ denotes a posterior probability >0.95 that the difference relative to the negative control is in the indicated direction. The ‘**’ indicates a posterior probability of >0.95 that the indicated difference is at least 1.3-fold. An ‘x’ marks a posterior probability >0.95 that the indicated difference is no more than 1.3-fold in either direction. (B) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in panel A from a representative experimental replicate. Equivalent mass of protein from each sample was probed with either anti-V5 antibody or anti-IntS1 as a loading control. (C) Repression activity was measured for three amounts of transfected wild type or cap-binding mutant (W873G) full length Pum via dual luciferase assay using the Nluc 3x PRE pA reporter. The fold change values were calculated relative to the equivalent amount of transfected RNA-binding defective mutant Pum (mut R7) negative control. V5-tagged EGFP plasmid served to balance the total mass of transfected plasmids in samples with 50 and 500 ng of Pum effector plasmid. Mean log2 fold change and 95% credible intervals for three experimental replicates with four technical replicates each are reported in the graph. Data and statistics are reported in Supplementary Table S1. (D) Western blot of the V5-epitope tagged Pum effector and EGFP balancer proteins used in panel C from a representative experimental replicate. Equivalent mass of protein from each sample was probed with anti-V5 antibody and, to assess equal loading of lanes, anti-tubulin antibody.

to NTC). Importantly, expression of wild type Pop2, but not the active site mutant Pop2, increased repression activity of RD2 and RD3 in cells treated with Pop2 dsRNA2 (Figure 4G). RD1 activity also increased, but the change did not meet our significance threshold. No effect of Pop2 expression on the activity of Pum N was evident. Protein expression of all effectors and the wild type and mutant Pop2 proteins was confirmed by western blot analysis of equivalent mass of cell extracts in Figure 4H. These observations indicate that the deadenylation activity of Pop2 is important for RD activity.

Ccr4 is the second deadenylase present in the CNOT complex (Figure 4A) (41, 74). We next evaluated the involvement of Ccr4 and observed that substantial depletion of its mRNA (Supplementary Figure S2A) did not affect repression by the Pum N-terminus or RDs (P(insig) ≥ 0.91) (Supplementary Figure S2B and C). This observation suggests that Ccr4 is not crucial, whereas Pop2 has an important
role in Pum repression. That said, the potential limitation of RNAi is notable, wherein residual low levels of Ccr4 might be sufficient to support Pum activity.

We also examined the role of non-catalytic CNOT subunits in Pum RD-mediated repression, starting with Not1, which is the central scaffold of the complex (Figure 4A) (74). RNAi depletion of Not1 was confirmed by RT-qPCR (Figure 5A) and western blotting (Figure 5B). We observed that knockdown of Not1 significantly abrogated repression by all three RDs (Figure 5C); a similar level of loss of repressive activity was observed for Pum N, although it did not meet our statistical significance thresholds.

We observed that RNAi depletion of Not1 also reduced the level of its protein partner Pop2 (Supplementary Figure S3A and B), consistent with a previous report (40). Therefore, the effect of RNAi of Not1 on Pum activity could be the result of diminished Pop2. To test this idea, we attempted to rescue the effect of Not1 depletion by expressing Pop2 from a transfected plasmid. First, we titrated Pop2 expression vector to approximate the level of endogenous Pop2 protein (Supplementary Figure S3B). We then performed tethered function assays. Again, depletion of Not1 reduced (for Pum N and RD1) or eliminated (for RD2 and RD3) repression activity (Supplementary Figure S3C). Expression of exogenous Pop2 did not rescue the loss of repression by the Pum N-terminus or RDs caused by Not1 depletion. Based on these observations, we conclude that Not1 is important for Pum RD activity, and its depletion mimics the effect of Pop2 depletion.

The other CNOT subunits form modules that interact with Not1, as depicted in Figure 4A (74). We observed that depletion of Not2 (Figure 5D–F) and Not3 (Figure 5G–I) also reduced Pum RD-mediated repression. Overall, depletion of Not2 or Not3 uniformly reduced the activity of Pum RDs (but not of Pum N) comparably to depletion of Not1, although in some instances the effect did not meet our statistical significance threshold (Figure 5F and I); relaxation of our criteria to require the 90% (instead of 95%) credible interval to omit zero would lead to significance calls in six of the nine RD/RNAi combinations (not shown). Pum effector expression in each RNAi condition was confirmed by western blotting (Figure 5J). We also tested the effects of depletion of Caf40, Not10 and Not11 (Supplementary Figure S2D–I), none of which alleviated the repressive activity of the Pum N-terminus or individual RDs (P(insig) > 0.88). We note that Caf40 knockdown did affect RD2, wherein its repressive activity was enhanced, an observation that is currently not understood. Overall, these results indicate that certain CNOT subunits are important for the repression activity of Pum RDs (i.e. Pop2, Not1, 2 and 3) whereas others appear to be dispensable (i.e. Ccr4, Caf40, Not10 and 11).

CNOT is required for efficient Pumilio-mediated mRNA decay

Given the importance of CNOT for Pum RD activity, we further examined its involvement in Pum-mediated mRNA degradation. To do so, we depleted Pop2 by RNAi and measured decay of the Nluc 3×PRE reporter in response to over-expressed Pum. In the control RNAi condition (NTC), over-expressed wild type Pum accelerated decay of the reporter mRNA, reducing its half-life by 3.2-fold relative to the negative control, RNA-binding defective mut R7 (Figure 6A and B). Depletion of Pop2 stabilized the reporter RNA in the presence of WT (by an estimated 17.8-fold change in half life, P(sig) > 0.999) or mut R7 (by an estimated 11.8-fold, P(sig) > 0.999) (Figure 6A and B). This result demonstrates that Pop2 is required for efficient Pum-mediated mRNA decay, consistent with the observation that Pop2 is important for PRE-mediated decay in Figure 1F and G. Using the same strategy, we analyzed the involvement of Not1 and observed that depletion of Not1 also led to impairment of Pum-mediated mRNA decay (Figure 6C and D), stabilizing the reporter mRNA by >11.9-fold (P(sig) > 0.999) in the wild type Pum condition and by more than an estimated 38-fold (P(sig) > 0.999) in the mut R7 condition. This data indicates that the CNOT complex plays a crucial role in the mechanism by which Pum accelerates mRNA decay.

Pumilio N-terminal Repression Domains bind to the CNOT complex

The observation that the repressive activity of Pum N-terminal RDs requires CNOT components suggested a model wherein they act to recruit the CNOT complex to target mRNAs. We used a co-immunoprecipitation assay to assess a possible physical interaction between flag-tagged Pum N-terminus or RBD with endogenous CNOT. Flag-tagged GST served as a negative control. Several positive controls were also utilized including the RNA-binding protein Nanos, which was previously shown to directly contact Not1 (45), and the stoichiometric CNOT complex subunits Not2 and Not3. We observed that the Pum N-terminus associates with Not1 (Figure 7A), similar to Nanos, and that this interaction is resistant to treatment with RNome (Figure 7A and B), indicating that the association is not bridged by RNA. As expected, Not1 robustly co-purified with Not2 and Not3. Interestingly, while the Pum RBD was previously shown to interact with Pop2 (42,44), we did not detect co-immunoprecipitation with Not1 under these conditions, perhaps reflecting differences in protein-protein contacts or affinities. Together, these observations indicate that the Pum N-terminus associates with CNOT.

We then investigated the potential for direct interaction between the Pum RDs and the CNOT complex by performing in vitro protein interaction assays using bead-bound purified Pum RD1, RD2, RD3 or RBD proteins, which were purified as fusions to maltose binding protein (MBP) and the StrepII affinity tag. Bead-bound MBP-StrepII served as a negative control. The eight subunit human CNOT complex was purified as described by Raisch et al. (Figure 7C and D, input) (57). We observed that each Pum RD interacted with the intact CNOT complex, including both deadenylases. The RBD similarly interacted with the CNOT complex, consistent with its reported interaction with Pop2 (42,44). This data indicates that each Pum repression domain can directly and individually bind to
Figure 4. The Pop2 deadenylase is required for Pum RD activity. (A) Diagram of the Drosophila melanogaster Ccr4–Not complex, containing eight subunits. Adapted from Temme et al. (74). (B) The efficiency of Pop2 mRNA depletion after 3 days of treatment with either of two double stranded RNAs (dsRNA1 and dsRNA2) was measured using RT-qPCR. The dsRNA1 targets the Pop2 coding sequence, whereas dsRNA2 targets the Pop2 mRNA 5′ UTR. Fold changes were calculated relative to non-targeting control RNAi (NTC) for the indicated experimental conditions. Mean log2 fold change and 95% credible intervals for a representative experimental replicate with three technical replicates for each measurement are reported in the graph. Data and statistics are reported in Supplementary Table S1. (C) Western blot confirming RNAi-mediated depletion of Pop2 deadenylase induced by treatment of d.mel-2 cells (three biological replicates) with two different double-stranded RNAs in comparison to non-targeting control dsRNA (NTC). Equivalent mass of cellular extract was loaded for each sample. Anti-tubulin western blot serves as a loading control. (D) The effect of Pop2 depletion on repression by Pum N-terminus and individual RDs was measured via tethered function dual luciferase assay. Repression by each effector was calculated relative to the corresponding negative control effector MS2-EGFP within each RNAi condition. Non-targeting control (NTC) RNAi serves as a negative control for RNAi. Tethered decapping enzyme subunit, Dcp1, serves as a positive control. Mean log2 fold change and 95% credible intervals for three experimental
the CNOT deadenylase complex. It is noteworthy that the CNOT complex is highly conserved throughout eukarya (39), as are Pumilio orthologs (1), and we previously reported that human Pumilio proteins are active in Drosophila cells (36,44,75). The observation that Drosophila Pum RDs bind to human CNOT accentuates the probable conservation of the repressive mechanism.

The poly(A) tail is necessary for maximal activity of the Pum N-terminus

As a complementary approach to analyze the role of deadenylation in the mechanism of repression by the Pum N-terminus, we tested whether the poly(A) tail is necessary. To do so, we compared Pum repression of the polyadenylated Nluc reporter to that of a non-adenylated reporter bearing a 3′ Histone Stem Loop (HSL) processing signal (Figure 8A) (76). First, we confirmed that each reporter generated the correct 3′ end product by cleaving them with RNase H and an antisense deoxyligonucleotide that is complementary to the Nluc coding region (Figure 8A). In addition, deadenylated 3′ end RNA fragments were generated for each reporter by adding oligo deoxynucleotide (oligo dT) to the indicated samples (Figure 8B). The resulting 3′ end fragments were detected by high resolution Northern blotting, thereby verifying that the Nluc × MS2 pA reporter produced a distribution of poly(A) lengths spanning up to 200 adenosomes appended to the 228 nucleotide 3′ end product, whereas the Nluc × MS2 HSL reporter mRNA produced the expected nonadenylated 210 nucleotide product (Figure 8B).

We then compared the ability of the Pum N-terminus to repress the adenylated and non-adenylated reporters. Repression by Pum N was significantly reduced by 2.5-fold on the HSL reporter relative to the adenylated reporter \((P\text{sig} > 0.999; \text{Figure } 8\text{C and D})\). Two conclusions can be drawn from this result. First, the reduction in activity emphasizes the importance of the poly(A) tail in maximal repression by the Pum N-terminus. Second, the residual repressive activity of Pum N on the non-adenylated mRNA indicates that it wields an additional poly(A) independent repression mechanism. We also analyzed the individual Pum RDs and observed that RD1 behaved similarly to the N-terminus, whereas RD2 and RD3 exhibited comparable repressive activity on the two reporters (Figure 8C and D), indicating that the RDs contribute to poly(A) independent repression. The tethered Dep1 control repressed both reporters (Figure 8C and D), consistent with its ability to promote 5′ decapping of the transcripts.

We postulated that the poly(A) independent repression activity of the N-terminus and RDs may be mediated via CNOT recruitment. This hypothesis is based on previous studies that reported deadenylation-independent repression by the CNOT complex (77,78). Moreover, the CNOT complex interacts with translational repressors and mRNA decay enzymes, including the decapping enzyme complex (74,77,79). We therefore analyzed the role of CNOT in poly(A) independent repression by the Pum N-terminus by measuring the impact of Not1 depletion. As in Figure 5, RNAi of Not1 significantly reduced repression by Pum N on the polyadenylated reporter mRNA, but no substantial effect on the poly(A)-independent repression of the HSL reporter was observed (Figure 8E and F). In contrast, when we examined the effect of Not1 knockdown on the individual RDs, we found that their ability to repress the HSL reporter was abrogated, and in all cases became indistinguishable from that of a corresponding EGFP control (Figure 8G and H). Our interpretation of these observations is that the Pum RDs rely on the CNOT complex to cause both poly(A) dependent and independent repression.

The Pum N-terminus utilizes an additional CNOT-independent repression activity

The residual repressive activity of the Pum N-terminus on the non-adenylated HSL reporter (Figure 8), and when CNOT components are depleted (Figures 4, 5 and 8), indicates that an additional mechanism contributes to repression. We further assessed the CNOT independent activity by simultaneously depleting both Not1 and Pop2. As before, depletion of each co-repressor individually decreased Pum N activity by 1.3- to 1.4-fold relative to NTC (Figure 9A). Simultaneous depletion of Not1 and Pop2 further significantly reduced Pum N-mediated repression by 1.6-fold relative to the NTC (Figure 9A). These results emphasize the importance of CNOT in poly(A) dependent repression, but also further argue for the presence of residual CNOT-independent repression activity.

To further understand CNOT-independent repression by the Pum N-terminus, we measured its effect on the levels of Nluc × MS2 pA mRNA when Pop2 or Not1 are depleted. As shown in Figure 9C–E, depletion of either CNOT component diminished Pum N mediated reduction of reporter mRNA levels, decreasing Pum N activity by 2.3-fold \((P\text{sig} > 0.99)\) or 3.1-fold \((P\text{sig} > 0.99)\) when Not1 or Pop2 were depleted, respectively, which replicates with four technical replicates each are reported in the graph. Data and statistics are reported in Supplementary Table S1. For significance calling, a ‘*’ denotes a posterior probability >0.95 that the difference relative to the negative control is in the indicated direction. The ‘**’ indicates a posterior probability of >0.95 that the indicated difference is at least 1.3-fold. An ‘x’ marks a posterior probability >0.95 that the indicated difference is no more than 1.3-fold in either direction. (E) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in panel D from a representative experimental replicate. Equivalent mass of protein from each sample was probed with anti-V5 antibody, followed by western blot of IntS1 as a loading control. (F) The efficiency of Pop2 mRNA depletion after 5 days of Pop2 dsRNA2 treatment was measured using RT-qPCR. Fold changes were calculated relative to non-targeting control (NTC). Mean log; fold change and 95% credible intervals for three biological replicates with three technical replicates each are reported in the graph. Data and statistics are reported in Supplementary Table S1. (G) The ability of wild type Pop2 (WT) or active site mutant Pop2 (mtt) to rescue repression by Pum N-terminus and RDs was measured via tethered function dual luciferase assay. Endogenous Pop2 was depleted by treating cells with dsRNA2. NTC dsRNA serve as a control. The effect of Pop2 expression was compared to EGFP control. Mean log; fold change and 95% credible intervals from three to six experimental replicates with four technical replicates each are reported in the graph. Data and statistics are reported in Supplementary Table S1. (H) Western blot of V5-tagged tethered effectors and myc-tagged Pop2, mutant Pop2, or negative control V5-tagged EGFP from a representative experimental replicate in panel G. Equivalent mass of cellular extract was loaded for each sample.
Figure 5. CNOT components are involved in Pum RD mediated repression. (A) The efficiency of RNAi-mediated depletion of Not1 mRNA after 3 days of dsRNA treatment was measured using RT-qPCR. Fold changes were calculated relative to non-targeting control (NTC). Mean log2 fold change and 95% credible intervals are reported in the graph for one representative experiment with three technical replicates of each measurement. Data and statistics are reported in Supplementary Table S1. (B) Western blot with anti-Not1 antibody confirms depletion of endogenous Not1 protein from a representative experiment. Equivalent mass of cellular extract was loaded for each sample. Anti-tubulin western blot serves as a loading control. (C) Tethered function dual luciferase assays measured the effect of Not1 depletion on the repression activity of Pum N-terminus and RDs. Non-targeting control (NTC) serves...
Pop2 were depleted, respectively, relative to NTC. Again, Pum N retained residual ability to reduce reporter mRNA levels. These observations are consistent with the results in Figures 1 and 6, wherein CNOT depletion stabilized the Pum-repressed mRNA, but did not completely eliminate decay.

Decapping enzyme participates in repression by the Pum N-terminus

We postulated that decapping may play a role in the residual repression activity of the Pum N-terminus. Removal of the 5′ cap of mRNAs plays a key role in the destruction of mRNAs through the 5′ decay pathway (38), and decapping is catalyzed by the enzyme Dcp2, which forms a complex with Dcp1 (79). We therefore inhibited decapping and examined its role in repression by the Pum N-terminus. To do so, Dcp2 was depleted via RNAi (as verified by RT-qPCR in Figure 10A) while simultaneously over-expressing an RNAi-resistant, catalytically inactive, dominant negative Dcp2 mutant (E361Q). As previously established (45,52,53), this combined approach was necessary to effectively abrogate decapping-mediated mRNA decay.

Inhibition of decapping significantly diminished repression of reporter mRNA levels by 2.2-fold and protein levels by 1.3-fold from the Pum N-terminus, relative to NTC (Figure 10B and C), indicating that decapping participates in the repression mechanism of Pum N-terminus. Supporting the efficacy of the approach, inhibition of decapping stabilized the Nluc reporter mRNA and Firefly luciferase internal control mRNA (Figure 10C). Further corroborating is provided by the observation that mRNA degradation by tethered Dcp1 was prevented by inhibition of Dcp2 (Figure 10B and C). Interestingly, tethered Dcp1 retained the ability to repress reporter translation upon Dcp2 knockdown, which likely reflects its association with translational inhibitory factors (Figure 10B) (79). Based on this collective data, we conclude that decapping contributes to the repression activity of the Pum N-terminus.

Multiple mechanisms contribute to Pumilio-mediated repression

Having identified multiple co-repressors – Not1, Pop2, Dcp2 (this study) and pAbp (44) – that are important for the individual activities of multiple Pum repression domains (N-terminal RDS and the C-terminal RBD), we investigated their contributions to regulation by full-length, endogenous Pum. To do so, Pum repression was measured using the Nluc 3×PRE reporter and compared to the unregulated reporter Nluc ΔPRE, which lacks PRE sequences. As before, FFLuc mRNA served as an internal control. Not1, Pop2, Dcp2 or pAbp were each depleted by RNAi, as confirmed by western blot (Figure 11A) or RT-qPCR (Figure 11B), and then reporter protein (Figure 11C) and mRNA levels (Figure 11D) were measured by dual luciferase assay and Northern blot, respectively. As before, the RNAi depletion of Dcp2 was accompanied by over-expression of mutant Dcp2 E361Q.

We analyzed the resulting data in two ways, focusing on the relative response ratio (RRR) either on the relative response ratio (RRR) between the Nluc reporter signal and the FFLuc internal control (Figure 11C), or the Nluc signal alone (Supplementary Figure S4). In either case, we considered log ratios relative to the corresponding non-targeting control (NTC) RNAi condition. The RRR-based approach normalizes potential sample-to-sample variation in transfection efficiency, and also responds to global effects on gene expression. To specifically measure PRE-dependent regulation in each RNAi condition, the RRR value of the Nluc 3× PRE reporter was divided by that of the RRR of the unregulated Nluc ΔPRE reporter, followed by calculation of the fold change relative to NTC. The results measure the PRE specific effect of depletion of the Pum co-repressors on reporter protein and mRNA levels. RNAi depletion of Pum served as a positive control (Figure 11A) and alleviated PRE-dependent repression, resulting in increased reporter protein (2.4-fold, \(P_{\text{sig}} > 0.99\)) and mRNA levels (2.8-fold, \(P_{\text{sig}} > 0.99\)) (Figure 11C). Depletion of Not1 or Pop2 alleviated PRE-dependent repression (2.1-fold, \(P_{\text{sig}} > 0.99\)) as a negative control for comparison. Activity of each effector was determined relative to the corresponding negative control effector MS2-EGFP within each RNAi condition. Mean log fold change and 95% credible intervals are reported in the graph for three experimental replicates with four technical replicates each. Data and statistics are reported in Supplementary Table S1. (E) Western blot confirms RNAi-mediated depletion of endogenous Not2 from a representative experiment using an anti-Not2 antibody. Equivalent mass of cellular extract was loaded for each sample. Anti-actin western blot serves as a loading control. The * designates a protein that cross-reacts with the Not2 antibody. (F) Tethered function dual luciferase assays measured the effect of Not2 depletion on the repression activity of Pum N-terminus and RDS. Non-targeting control (NTC) serves as negative control for comparison. Activity of each effector was determined relative to the corresponding negative control effector MS2-EGFP within each RNAi condition. Mean log fold change and 95% credible intervals are reported in the graph for three experimental replicates with four technical replicates each. Data and statistics are reported in Supplementary Table S1. (G) The efficiency of Not3 mRNA depletion after 3 days of dsRNA treatment was measured using RT-qPCR. Fold changes were calculated relative to non-targeting control (NTC). Mean log fold change and 95% credible intervals are reported in the graph for one representative experiment with three technical replicates of each measurement. Data and statistics are reported in Supplementary Table S1. (H) Western blot confirms RNAi depletion of endogenous Not3 protein from a representative experiment using an anti-Not3 antibody. Equivalent mass of cellular extract was loaded for each sample. Anti-tubulin western blot serves as a loading control. (I) Tethered function assays measured the effect of Not3 depletion on the repression activity of Pum N-terminus and RDS. Non-targeting control (NTC) serves as negative control for comparison. Activity of each effector was determined relative to the corresponding negative control effector MS2-EGFP within each RNAi condition. Mean log fold change and 95% credible intervals are reported in the graph. Data and statistics are reported in Supplementary Table S1. (J) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in panels C, F and I from a representative experiment. Equivalent mass of protein from each sample was probed with anti-V5 antibody and IntS1 loading control.
Figure 6. Pum-mediated mRNA decay requires Not1 and Pop2. (A) The effect of RNAi-mediated depletion of Pop2 by Pop2 dsRNA1 on the mRNA decay rate of Nluc 3× PRE reporter mRNA was measured in response to over-expressed wild type Pum (Pum WT) or the RNA-binding defective mutant Pum (Pum mut R7) following inhibition of transcription with ActD. Cells treated with NTC dsRNA served as negative control. The Nluc 3× PRE was detected by Northern blot along with 18S ribosomal RNA (rRNA), as a loading control. Each lane of the gel contains 10 μg of total RNA. The mRNA half-lives and 95% credible intervals measured in each condition are shown below the respective blots, and were calculated from three experimental replicates. Data and statistics are reported in Supplementary Table S1. (B) The fraction of Nluc 3× PRE mRNA remaining, normalized to 18S rRNA, is plotted relative to time (minutes) after inhibition of transcription. Datum points for each of three experimental replicates are plotted. First order exponential decay trend lines, calculated using non-linear regression analysis, are plotted for each effector (Pum WT in blue, and mut R7 in red) in each RNAi condition (NTC, solid lines, and Pop2, dashed lines). (C) The effect of RNAi-mediated depletion of Not1 on the mRNA decay rate of Nluc 3× PRE reporter mRNA was measured in response to over-expressed wild type Pum (Pum WT) or the RNA-binding defective mutant Pum (Pum mut R7) following inhibition of transcription with ActD. Cells treated with NTC dsRNA served as negative control. The Nluc 3× PRE was detected by Northern blot along with 18S ribosomal RNA (rRNA), as a loading control. Each lane of the gel contains 10 μg of total RNA. The mRNA half-lives and 95% credible intervals measured in each condition are shown below the respective blots, and were calculated from three experimental replicates. Data and statistics are reported in Supplementary Table S1. (D) The fraction of Nluc 3× PRE mRNA remaining, normalized to 18S rRNA, is plotted relative to time (minutes) after inhibition of transcription. Datum points for each of 3 experimental replicates are plotted. First order exponential decay trend lines, calculated using non-linear regression analysis, are plotted for each effector (Pum WT in blue and mut R7 in red) in each RNAi condition (NTC, solid lines and Not1, dashed lines).

and 2.0-fold, \( P(\text{sig}) > 0.99 \), respectively) and stabilized the reporter mRNA (1.6-fold, \( P(\text{sig}) = 0.98 \) and 1.9-fold, \( P(\text{sig}) = 0.86 \), respectively) (Figure 11C). Inhibition of Dcp2 more modestly increased reporter protein expression (1.4-fold, \( P(\text{sig}) = 0.99 \)) and stabilized the PRE-containing reporter mRNA (1.5-fold, \( P(\text{sig}) = 0.83 \)) (Figure 11C), as did depletion of pAbp (1.4-fold increase in PRE reporter protein, \( P(\text{sig}) = 0.99 \); 1.3-fold increase in PRE reporter mRNA, \( P(\text{sig}) = 0.44 \)), reflecting their respective contributions to repression by the Pum N-terminus and RBD.

Consistent with their global roles in basal mRNA decay, depletion of CNOT, and to a lesser degree Dcp2, affected FFluc levels, as observed by Northern blot (Figure 11D). Hence, we also analyzed PRE-dependent regulation by omitting the internal control FFluc from the calculations. In this case, to specifically measure PRE-dependent regulation, the level of the Nluc 3× PRE reporter was divided by the unregulated Nluc ΔPRE reporter for each RNAi condition, and then fold change values were calculated relative to the negative control RNAi condition. The resulting PRE-dependent fold change values are reported in Supplementary Figure S4. Again, we observed a substantial loss in PRE-dependent repression upon depletion of Pum (2.2-fold increase, \( P(\text{sig}) > 0.99 \)), Not1 (2.0-fold increase, \( P(\text{sig}) > 0.99 \)), or Pop2 (2.1-fold increase, \( P(\text{sig}) > 0.99 \)), demonstrating that the observed changes are not simply artifacts due to the global effects of CNOT or Dcp2 depletion. Depletion of pAbp caused a 1.4-fold increase (\( P(\text{sig}) = 0.91 \)) and Dcp2 inhibition had a minor effect (1.1–1.2-fold increase, though without normalization to transfection efficiency, the variation limited our ability to draw a firm conclusion). Taken together, our results demonstrate the multiple co-repressors contribute to Pum-mediated repression of target mRNA and protein levels in Drosophila cells.

DISCUSSION

The results of this study provide new insights into the molecular mechanism by which Pum represses gene expression. Previous work correlated Pum repression with reduction...
in mRNA levels, and we now show that Pum uses multiple repression domains to accelerate mRNA decay via the CNOT deadenylase and decapping complexes. This information enhances our understanding of the biological roles and impact of Pum on the transcriptome.

The model that emerges from our findings is that Pum utilizes four domains that interact with the CNOT deadenylase complex to repress target mRNAs (Figure 11E). Multiple lines of evidence directly link CNOT to Pum-mediated repression. First, Pum requires CNOT subunits to repress protein expression and accelerate RNA decay, specifically the Not1, Not2, Not3 and Pop2 subunits (Figures 4 and 5). Second, the catalytic activity of Pop2, the major deadenylase enzyme in Drosophila, is necessary to rescue Pum RD-mediated repression in the Pop2 RNAi condition (Figure 4). Third, the poly(A) tail is necessary for maximal repression by Pum (Figure 8). Fourth, CNOT co-immunoprecipitates with the Pum N-terminus in an RNase resistant manner (Figure 7). Fifth, the Pum RDs and RBD directly bind to the CNOT complex (Figure 7). In addition, the C-terminal RBD of Pum was previously shown to interact with Pop2 and promote deadenylation (42,44). Moreover, Pop2 has been linked to deadenylation of target mRNAs during Drosophila development (8,43). It is noteworthy that we were not able to measure Pum-enhanced deadenylation in cultured cells due to technical limitations. At the earliest time points at which we could detect the PRE-containing reporter mRNA, the poly(A) tail was already predominantly short (Supplementary Figure S5A–C). Depletion of Pum or deletion of the PRE increased the reporter mRNA level, as expected, but did not alter the poly(A) length distribution, which again was predomi-
Figure 8. The poly(A) tail is necessary for maximal activity of the Pum N-terminus. (A) Diagram of the Nluc 2×MS2 reporters with either 3′ poly(A) tail or the Histone Stem Loop (HSL) used for tethered function assays. The reporters are identical except that the cleavage/poly-adenylation element was
nantly short for both the reporter with or without PREs. We did observe that depletion of Pop2 increased poly(A) length and stabilized the PRE-containing mRNA, consistent with its importance in catalyzing the reporter’s deadenylation.

Why does Pum use multiple domains to recruit CNOT? Perhaps their activities combine to increase the avidity of Pum for CNOT, mediated by multiple contacts between Pum domains and CNOT. Indeed, the four repressive domains each directly interact with the CNOT complex (Figure 7). Mapping the precise protein-protein interactions necessary for CNOT recruitment by Pum will require detailed biochemical and structural analysis, to be pursued in future studies.

Recruitment of CNOT by RNA-binding factors has emerged as an important mechanism of post-transcriptional regulation (39,45,80–83). Utilization of CNOT by Pum orthologs has been reported in Saccharomyces cerevisiae, Schizosaccharomyces pombe, Caenorhabditis elegans, Drosophila melanogaster and mammals (11,43,44,75,84–86), and thus represents an evolutionarily conserved mechanism. Like Drosophila Pum, the highly conserved RBD of Pum orthologs spanning from yeast to human universally interact with Pop2 orthologs. In contrast, the N-terminal RDs are a more recent evolutionary addition, being found in Pum orthologs spanning insects through mammals (36). Based on the results of this study, we speculate that analogous regions of those Pum orthologs repress by recruiting the CNOT complex. Consistent with this idea, experimental evidence showed that the N-termini of human Pum orthologs, PUM1 and PUM2, exhibit repressive activity when directed to an mRNA (36). Future research should dissect the repressive mechanism of mammalian Pum N-termini.

It is interesting to speculate that Pum might recruit a subcomplex of CNOT, based on the observation that Pop2, Not1, Not2 and Not3 were important for Pum RD activity whereas depletion of others had little to no effect (i.e. Ccr4, Not10, Not11, Caf40) (Supplementary Figure S2). Germaine to this idea, evidence in yeast and human cells revealed heterogeneity in size and composition of CNOT complexes (87–90). The requirement of specific CNOT subunits for repression by Pum orthologs was also observed in S. cerevisiae (85,91). Still, interpretation of the negative results in our experiments is limited by the effectiveness of RNAi-mediated depletion, and we acknowledge that residual levels of a CNOT component may be sufficient to support activity. Moreover, our biochemical analysis indicates that Pum repression domains can bind the intact CNOT complex (Figure 7). We also found that decapping contributes to repression by Pum, supported by several lines of evidence. First, the N-terminus retains partial repressive activity when the target mRNA lacks a poly(A) tail and therefore is not subject to deadenylation (Figure 8). In addition, the N-terminus retains repression and RNA decay activities when Not1 and/or Pop2 are depleted (Figures 1, 6 and 9). Moreover, inhibition of the decapping enzyme Dcp2 reduced repression and RNA decay activity of the Pum N terminus (Figures 10 and 11). Future analyses will be necessary to delineate the specific region(s) of Pum that modulate decapping and how it associates with decapping enzyme—either via direct protein contacts or via bridging proteins that interface with the decapping machinery (79). Interestingly, decapping appears to be a conserved mechanism of Pum repression, supported by the observations that yeast PUF proteins associate with decapping factors and promote decapping of target mRNAs (85,92,93).

How does Pum affect protein synthesis? Because the 5’ cap is crucial for translation of most mRNAs, and the poly(A) tail and pAbp promote translation, Pum-mediated deadenylation and decapping can contribute to both repression of protein synthesis and mRNA destruction. Indeed, Pum-mediating repression of protein level corresponds in magnitude to the reduction in mRNA level. Based on con-

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replaced with a Histone Stem Loop and Histone Downstream Element (HDE) in the Nluc 2× MS2 HSL reporter, which produces a non-adenylated 3’ end. The location of the probe used for Northern blotting (green) and the DNA oligonucleotide (red) used for RNase H cleavage of the mRNAs for high resolution Northern blotting are indicated. Diagram is not drawn to scale. (B) High resolution Northern blot of the Nluc 2× MS2 pA and HSL reporter mRNAs expressed in d melan2 cells confirms proper poly-adenylation of the pA reporter and lack of poly-adenylation of the HSL reporter. Where indicated (+), RNA was treated with DNA oligonucleotide of 15 thymidines (dT) and RNase H to degrade the poly(A) tail. RNA size markers are indicated on the left. The lengths of the 3’ end reporter fragments are also indicated. (C) The repression activity of the Pum N-terminus and RDs was measured via tethered function dual luciferase assay, using the Nluc 2× MS2 poly(A) and HSL reporters. The activity of each effector was determined relative to the tethered EGFP negative control effector on the same reporter. Mean log₂ fold change and 95% credible intervals are reported in the graph for three experimental replicates with four technical replicates each. Data and statistics are reported in Supplementary Table S1. For significance calling, a ** denotes a posterior probability >0.95 that the difference relative to the negative control is in the indicated direction. The *** indicates a posterior probability of >0.95 that the indicated difference is at least 1.3-fold. An ‘x’ marks a posterior probability >0.95 that the indicated difference is no more than 1.3-fold in either direction. (D) Western blot detection of the V5-epitope tagged MS2 fusion effector proteins used in panel C from a representative experimental replicate. Equivalent mass of protein from each sample was probed with anti-V5 antibody. (E) The effect of RNAi-mediated depletion of Not1 on repression activity of the Pum N-terminus was measured via tethered function using the Nluc 2× MS2 poly(A) and HSL reporters. The non-targeting control (NTC) served as negative control for RNAi. The repression activity of each effector was determined relative to tethered EGFP negative control in the same RNAi condition. The mean log₂ fold change and 95% credible intervals are graphed for three experimental replicates with four technical replicates each. Data and statistics are reported in Supplementary Table S1. (F) Western blot of the V5-epitope tagged MS2 fusion effector proteins from a representative experimental replicate from panel E. Equivalent mass of protein from each sample was probed with anti-V5 antibody or anti-IntS1 to assess equal loading of lanes. The depletion of Not1 protein was assessed using an anti-Not1 antibody, with anti-tubulin western blot serving as a loading control. (G) The effect of RNAi depletion on the repression activity of the Pum RDs was measured via tethered function dual luciferase assay, using the Nluc 2× MS2 HSL reporter. Non-targeting control (NTC) serves as negative control for comparison. Activity of each effector was determined relative to tethered EGFP negative control in the same RNAi condition. The mean log₂ fold change and 95% credible intervals are reported in the graph for three experimental replicates with four technical replicates each. Data and statistics are reported in Supplementary Table S1. (H) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in from a representative experimental replicate from panel G. An equivalent mass of protein from each sample was probed with anti-V5 antibody, and then with anti-IntS1 to assess equal loading of lanes. Depletion of the Not1 protein was assessed using an anti-Not1 antibody, with anti-tubulin western blot serving as a loading control.
Figure 9. The Pum N-terminus utilizes an additional deadenylation-independent repression activity. (A) The effect of RNAi-mediated depletion of Not1, Pop2 or both simultaneously on the repression activity of the Pum N-terminus was measured in the tethered function reporter assay. The activity of each effector protein was determined relative to the negative control MS2-EGFP within each RNAi condition. Tethered Dcp1 served as a control. The mean log₂ fold change and 95% credible intervals are graphed for three experimental replicates with four technical replicates each. Data and statistics are reported in Supplementary Table S1. For significance calling, a ‘*’ denotes a posterior probability >0.95 that the difference relative to the negative control is in the indicated direction. The ‘**’ indicates a posterior probability of >0.95 that the indicated difference is at least 1.3-fold. An ‘x’ marks a posterior probability >0.95 that the indicated difference is no more than 1.3-fold in either direction. (B) Western blot of the V5-epitope tagged MS2 fusion effector proteins used in panel A from a representative experimental replicate. Equivalent mass of protein from each sample was probed with anti-V5 antibody, and then with anti-tubulin to assess equal loading of lanes (Asterisks indicate residual anti-V5 signal on the blot). Depletion of Not1 and Pop2 proteins was assessed by Western blot detection of the endogenous proteins. Anti-tubulin Western blot served as a loading control. Note that Not1 depletion also reduces Pop2 protein level, as discussed in the text. (C) The effect of depletion of Pop2 or Not1 on repression activity of Pum N-terminus was measured using the tethered function dual luciferase assay and Northern blotting. Repression of reporter protein and mRNA levels by tethered Pum N-terminus was calculated relative to the negative control MS2-EGFP within the same RNAi condition. Tethered Dcp1 served as a positive control. Mean log₂ fold change and 95% credible intervals from three biological replicates are reported in the graph. Data and statistics are reported in Supplementary Table S1. (D) Northern blot detection of Nluc 2× MS2 reporter and internal control FFluc mRNAs in the three biological replicates for each effector and RNAi condition. Each lane of the gel contains 5 μg of total RNA. Quantitation of this blot is represented in panel C. Ethidium Bromide detection of rRNA was used to assess integrity and equivalent loading of the RNA samples. (E) Western blot of the V5-tagged MS2 fusion effector proteins used in panel C from three biological replicates. Equivalent mass of protein from each sample was probed with anti-V5 antibody.
Figure 10. Decapping enzyme participates in repression by the Pum N-terminus. (A) RNAi-mediated depletion of the Dcp2 mRNA after 5 days of dsRNA treatment was measured using RT-qPCR. Fold changes were calculated relative to non-targeting control (NTC). Mean log2 fold change and 95% credible intervals for three biological replicates with three technical replicates each are reported in the graph. Data and statistics are reported in Supplementary Table S1. (B) The effect of inhibition of decapping on repression of Niuc 2 x MS2 reporter protein and mRNA levels by the Pum N-terminus was measured using the tethered function dual luciferase assay. Decapping was inhibited by RNAi mediated depletion of Dcp2 and over-expression of the dominant negative mutant Dcp2 E361Q. Repression activity was calculated relative to tethered EGFP negative control within the same RNAi condition. Tethered Dcp1 served as a positive control. Mean log2 fold change and 95% credible intervals are graphed from three biological replicates. Data and statistics are reported in Supplementary Table S1. For significance calling, a ‘*’ denotes a posterior probability > 0.95 that the difference relative to the negative control is in the indicated direction. The ‘**’ indicates a posterior probability of >0.95 that the indicated difference is at least 1.3-fold. An ‘x’ marks a posterior probability >0.95 that the indicated difference is no more than 1.3-fold in either direction. (C) Northern blot detection of Niuc 2 x MS2 reporter and FFluc internal control in three biological replicate samples for tethered effectors analyzed in panel A. Each lane of the gel contains 5 μg of total RNA. Ethidium Bromide detection of rRNA was used to assess integrity and equivalent loading of the RNA samples. (D) Western blot of the V5-epitope tagged MS2 fusion effector proteins and Dcp2 E361Q used in panel B from three biological replicates. Equivalent mass of protein from each sample was probed with anti-V5 antibody.

servation of a cap-binding motif that contains a key tryptophan residue (W783) (48), Pum was proposed to directly inhibit translation by binding the 5’ cap of target mRNAs, and mutation of W783 moderately reduced Pum’s ability to repress a GFP reporter bearing the mad 3’UTR in S2 cells (11). However, we did not observe an effect of this mutation on repression by the Pum N-terminus in the tethered function assay or by full length Pum using the PRE reporter (Figure 3). Likewise, our previous analysis showed that the PCMbdomain encompassing the putative cap binding motif was neither necessary nor sufficient for repression (36). Thus, the proposed mechanism of cap-dependent inhibition does not appear to make an essential contribution to the Pum activity measured here. We cannot rule out that it might have a potential role in repression in other cellular or developmental contexts, or within the context of certain mRNAs (10,11).

The Pum RBD contributes to translational repression by associating with and antagonizing the activity of pAbp (44). Consistent with this role, we observed that depletion of pAbp diminished Pum/PRE-mediated repression of protein and mRNA levels (Figure 11). These observations lead us to speculate about the potential functional interplay between Pum, pAbp, and deadenylation. Binding of pAbp to poly(A) was originally thought to interfere with deadenylation (94); however, recent studies indicate that the relationship is more complex (95,96), wherein Ccr4 and Pop2 deadenylase activities were shown to be differentially affected by pAbp orthologs. Analysis of human PABPC1 and S. pombe Pab1 indicate that they are required for Ccr4 deadenylase activity, whereas they inhibit activity of Pop2 orthologs. Further evidence in vitro suggests that S. pombe Ccr4 can displace Pab1 from the poly(A) tail, whereas Pop2 cannot. Whether these properties carry over to Drosophila remains to be determined. In principle, Pum could cause displacement of pAbp from the mRNA, thereby bypassing the role of Ccr4, resulting in accelerated Pop2-catalyzed deadenylation. Contradicting this hypothesis, however, is the observation based on RNA immunoprecipitation of pAbp, that the Pum RBD did not dislodge pAbp from mRNA (44).
Figure 11. Multiple mechanisms and co-repressors contribute to Pumilio-mediated repression. (A) RNAi-mediated depletion of endogenous Pum, Not1, and Pop2 proteins was assessed by Western blot of three biological replicate samples each from d.mel2 cells that were treated with the indicated dsRNA for three days. Note that Pum and Not1 antibodies each recognize two isoforms of their respective proteins. Equivalent mass of protein was analyzed for each sample, and anti-tubulin western blots were performed as a loading control. (B) RNAi-mediated depletion of pAbp mRNA (left) and Dcp2 mRNA (right) were measured using RT-qPCR. Fold changes were calculated relative to non-targeting control (NTC). Mean log2 fold change and 95% credible intervals for three biological replicates are reported in the graph. Data and statistics are reported in Supplementary Table S1. (C) The effect of RNAi depletion of Pum co-repressors Not1, Pop2, Dcp2 and pAbp on repression of Nluc 3× PRE reporter protein and mRNA expression levels by endogenous Pum was measured in d.mel-2 cells. Data was analyzed by calculating the Relative Response Ratio for each sample by dividing the Nluc signal by corresponding FFluc signal, thereby normalizing variation in transfection efficiency. Next, the PRE-dependent effect of each RNAi condition on the Pum repressed, PRE containing reporter was normalized to the effect on the unregulated Nluc ΔPRE reporter, which contains a minimal 3′ UTR that lacks Pum binding sites. The fold change in PRE-mediated regulation within each RNAi condition was then calculated relative to the negative control (NTC) dsRNA. RNAi of Pum served as a positive control. Mean log2 fold change and 95% credible intervals for three biological replicates are reported in the graphs. Data and statistics are reported in Supplementary Table S1. For significance calling, a † denotes a posterior probability >0.95 that the difference relative to the negative control is in the indicated direction. The †‡ indicates a posterior probability of >0.95 that the indicated difference is at least 1.3-fold. An † marks a posterior probability >0.95 that the indicated difference is no more than 1.3-fold in either direction. (D) Northern blot detection of Pum-regulated Nluc 3× PRE, unregulated Nluc ΔPRE reporter mRNA and FFluc internal control mRNAs in three biological replicate samples for each RNAi condition analyzed in panels C and D. Each lane of the gel contains 5 μg of total RNA. Ethidium bromide detection of rRNA was used to assess integrity and equivalent loading of the RNA samples. (E) Model of Pum-mediated repression. The RNA-binding domain (RBD) of Pum binds to mRNAs that contain a Pum Response Element (PRE). Multiple domains of Pum contribute to repression activity including the N-terminal repression domains (RD1, RD2, and RD3) and C-terminal RBD. Pum represses the target mRNA by multiple mechanisms including acceleration of mRNA decay via recruitment of Ccr4–Pop2–Not (CNOT) deadenylase complex, leading to deadenylation of the 3′ poly-adenosine tail, and via decapping enzyme (Dcp2) mediated removal of the 5′ 7-methyl guanosine cap (7mGppp). Pum RBD also antagonizes the translational activity of poly-adenosine binding protein (pAbp). CNOT subunits that are important for Pum RD-mediated repression are shaded in red. Red arrows indicate enzyme-catalyzed hydrolysis of the RNA. Arrows with gray-black gradient indicate Pum-CNOT interactions, as described in the Discussion. The means by which Dcp2 is modulated by Pum N-terminus remains to be determined.
Alternatively, the Pum RBD may interfere with the ability of pAbp to promote translation initiation. This idea is supported by functional assays, wherein we showed the Pum RBD inhibits the ability of pAbp to promote translation when bound to an internal poly(A) tract within an mRNA engineered without a 3′ poly(A) tail (44). Future work will include determining the specific interactions between Pum and pAbp, and how this might influence translation efficiency and/or deadenylation.

Our results have important implications for understanding Pum-mediated repression in embryos, the germline and neurons. Pum repression of hunchback mRNA in early embryogenesis was linked to poly(A) tail shortening (8). Pum repression was also correlated with mRNA decay during the maternal-to-zygotic transition (MZT), wherein many PRE-containing, maternally provided mRNAs are coordinate degradation (28, 29, 97). Pum-mediated repression was also linked to deadenylation by CNOT in primordial germ cells, where Pum contributes to repression of cyclin B mRNA (42), and in GSCs, where Pum participates in repression of mRNAs such as mei-P26 (43). The mechanisms of Pum-mediated deadenylation and decapping likely contribute to mRNA degradation observed in these contexts. Pum also represses specific mRNAs in neurons, such as paralytic, which encodes a voltage-gated sodium channel (15, 16, 98); however, the impact on mRNA decay remains to be examined in this context. Future analysis should investigate the contributions of Pum RDs to these and other processes.

Regulation of a growing number of Pum target mRNAs involves collaboration with other RBPs, with Nanos and Brat being the best documented examples. How can the mechanisms of Pum repression integrate with the activities of these RBPs? In the case of Nanos, it binds cooperatively with Pum to certain target mRNAs that possess a Nanos binding site preceding a PRE motif (25, 99). Nanos also confers its own repressive activity that accelerates deadenylation (45). Nanos may synergize with Pum in the recruitment of CNOT by contributing additional contacts with the Not1 and Not3 subunits. Nanos was also reported to interact with Not4 (42), though the potential role of Not4 in deadenylation and its involvement in Nanos activity are unclear. In fact, Drosophila Not4 does not appear to be a stable, stoichiometric CNOT component (40). Nanos also promotes decapping, but like Pum, how it does so is presently not well understood (45). Thus, combinatorial control by Pum and Nanos together accelerate the same key steps contributing to silencing of translation and mRNA destruction (2).

Pum can also collaborate with Brat to repress certain mRNAs that contain both a PRE and Brat binding site (10, 11, 28, 33, 34). The mechanism of Brat-mediated repression in embryos was reported to involve recruitment of the translational repressor eIF4E homologous protein (4EHP) (32). Brat may also affect mRNA decay, supported by the observation that it co-purifies with the CNOT complex (40). Depletion of Pop2 reduced the combined repressive activity of Pum and Brat (11). Future analyses will be necessary to interrogate the role of CNOT in Brat-mediated repression and whether Pum and Brat can synergistically recruit CNOT to their mutual target mRNAs. Pum, Nanos and Brat all collaborate to bind and repress the maternal hunchback mRNA during embryogenesis through a combination of cooperative RNA-binding, translational repression, deadenylation, and mRNA decay (reviewed in (2)). It is noteworthy that the assembly of this triumvirate of repressors on one mRNA is likely a rare scenario, as few mRNAs are predicted to contain the requisite cluster of binding sites for these three RBPs (2).

The mechanistic insights into Pum-mediated repression also have important implications for Pum orthologs in other species. Drosophila Pum serves as an archetype, and the conservation of its repressive domains, including those in its N-terminus and RBD, in animals ranging from insects to vertebrates indicates that the mechanisms and co-repressors described here will be relevant (47). Indeed, as noted above, accumulating evidence supports the role of deadenylation, decapping and translational inhibition in Pum repression in multiple model organisms. Mammalian Pum orthologs have crucial, diverse roles in growth and development, gametogenesis, hematopoiesis, neurogenesis, behavior, motor function and memory formation (100–109). Their dysfunction has now been linked to cancer, neurodegeneration, epilepsy, memory impairment, reduced fertility, and developmental defects in mammals (100, 101, 103, 106, 110–114). Thus, we anticipate that further elucidation of Pum function will facilitate better understanding of its role in development and disease, and perhaps inform future therapeutic interventions.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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